Department of Biosciences Faculty of Biological and Environmental Sciences University of Helsinki Finland

Computational genomics of lactobacilli

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ACADEMIC DISSERTATION

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"If you wish to make an apple pie from scratch, you must first invent the universe." - Carl Sagan

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Abstract

Lactobacilli are gram-positive lactic acid bacteria (LAB) and have important implications for food production and preservation as well as human health and wellbeing. These bacteria occupy various niches in and on the human body, such as the gastrointestinal, respiratory, and urogenital tracts, and have been used for centuries in the fermentation of dairy products, the pickling of vegetables, baking, and curing fish, meats, and sausages. Recently, the use of lactobacilli as biotherapeutic agents has attracted interest. However, the molecular basis of host-microbe interactions, food production abilities and beneficial effects on health of lactobacilli are not well understood and deserve more research. In this thesis research, bioinformatics approaches were developed for genome-scale protein function classification, and the genetic composition of two also human-associated *Lactobacillus* species was determined by means of genome sequencing and computational genomics. Taken together, the results of these analyses illustrated that genome sequencing and computational genomics represent valuable approaches to the study of lactobacilli and to understanding their physiology. Furthermore, these methods provide effective means of identifying lactobacillar components that are involved in host-interactions.

Protein function prediction is one of the most crucial tasks of any genome sequencing project. In this thesis, two bioinformatics software tools were developed for the systematic analysis of protein function that are more advanced than current methods in several respects. The first automatic function prediction method, called LOCP, was developed to fulfil the need for rapid and accurate genome-wide identification of putative pilus operons in gram-positive bacteria. The computational resource was designed to support for both nucleotide sequence input or annotated bacterial protein sequence data and introduced a novel approach that combines similarity searches and statistical detection of sortase- and pilin-motif enriched regions for the prediction of putative pilus operons in gram-positive genomes. Markedly, the tool identified all genuine pilus gene clusters from the test genome sequences and made in the benchmarking test no false predictions. The second bioinformatics tool disclosed an improved homology-based function prediction solution and was created to offer an effective approach to the large-scale computational annotation of uncharacterised bacterial protein sequences. Compared to existing solutions for homology-based function prediction, BLANNOTATOR groups sequences that are found using sequence similarity searches into subsets according to their biological function and uses a set of matches with consistent functional information as the basis for annotation transfer to the query sequence instead of relying on a single match or all matches as many competing tools do. This procedure improved the functional classification substantially, producing consistent results and facilitating comparisons among various organisms. Overall, the two tools developed in this thesis are important additions to the current repertoire of function classification systems that are applicable to bacterial proteins and provided a novel means to classify bacterial proteins at the genome level. Most importantly, annotation accuracy was high, and both tools provided information that otherwise might have been ignored or considered too labour intensive to find.

Lactobacillus rhamnosus GG is a probiotic bacterium that has a long history of safe use in foods and that has a well-documented beneficial effect on human health. To gain insight into its physiology and to elucidate the lactobacillar components that are involved in interactions with the host, the genomes of *L. rhamnosus* GG and its closely related dairy isolate *L. rhamnosus* LC705 were sequenced and analysed. Although the two genomes were shown to exhibit a high degree of synteny, altogether, nine regions of diversity punctuated the colinearity between the two genomes. The five GG-specific diversity regions included a number of genes encoding bacteriophage components and other genes that are implicated in sugar transport, metabolism, and exopolysaccharide (EPS) biosynthesis. In addition, genes for three pilus subunits and a pilin-dedicated sortase were identified in one of the diversity regions. Importantly, the presence of the pili on the cell surface of *L. rhamnosus* GG was confirmed and one of the GG-specific pilins was shown to be instrumental for the mucus interaction of *L. rhamnosus* GG. The diversity regions in LC705 strain encoded rhamnose, ribose, and maltose transporters and fermentative capacities that are missing from GG, thereby extending its metabolic versatility beyond that of GG and enabling LC705 to survive in a range of environments.

The genetic makeup of Lactobacillus crispatus was in turn explored by sequencing and analysing the genome of L. crispatus ST1 and performing a comparative genomic analysis of the chicken isolate ST1 and nine other L. crispatus genomes. The analyses revealed a rather compact genome and indicated that the genetic diversity present within L. crispatus has not yet been captured exhaustively. Specifically, the genomes of ST1 and nine vaginal strains were predicted to contain a pan-genome of 3,929 gene families, of which 1,224 families made up the L. crispatus conserved core. Mathematical extrapolations of these data to an infinite number of strains suggested that the core genome reaches a plateau of approximately 1,116 gene families and that the pan-genome doubles in size with the addition of the next 107 genomes, illustrating the value of sequencing many isolates. Interestingly, the comparison of protein-coding gene (CDS) contents revealed differences that are potentially relevant for the genetic adaptation of L. crispatus to different habitats, such as the existence of different EPS biosynthesis gene clusters in different strains and the Type II CRISPR (clustered regularly interspaced palindromic repeats)-Cas (CRISPRassociated) invader defence system that is specific to vaginal strains. In contrast, adhesin genes that are potentially involved in the exclusion and displacement of the bacterial vaginosis-associated species Gardnerella vaginalis were predicted to be present in the L. crispatus core genome, suggesting that all L. crispatus strains might have the potential to prevent bacterial vaginosis.

List of original publications

This thesis is based on the following publications:

- I **Plyusnin I, Holm L, Kankainen M. (2009)** LOCP--locating pilus operons in gram-positive bacteria. *Bioinformatics* **25**:1187-1188.
- II Kankainen M, Ojala T, Holm L. (2012) BLANNOTATOR: enhanced homology-based function prediction of bacterial proteins. *BMC bioinformatics* 13:33
- III Kankainen M, Paulin L, Tynkkynen S, von Ossowski I, Reunanen J, Partanen P, Satokari R, Vesterlund S, Hendrickx APA, Lebeer S, De Keersmaecker SC, Vanderleyden J, Hämäläinen T, Laukkanen S, Salovuori N, Ritari J, Alatalo E, Korpela R, Mattila-Sandholm T, Lassig A, Hatakka K, Kinnunen KT., Karjalainen H, Saxelin M, Laakso K, Surakka A, Palva A, Salusjärvi T, Auvinen P, de Vos WM. (2009) Comparative Genomic Analysis of *Lactobacillus rhamnosus* GG Reveals Pili Containing a Human Mucus-Binding Protein. *Proc Natl Acad Sci U S A* 106:17193-17198.
- IV Ojala T, Kuparinen V, Koskinen JP, Alatalo E, Holm L, Auvinen P, Edelman S, Westerlund-Wikström B, Korhonen TK, Paulin L, Kankainen M. (2010) Genome sequence of *Lactobacillus crispatus* ST1. J Bacteriol 192:3547-3548.
- V Ojala T, Kankainen M, Castro J, Cerca N, Edelman S, Westerlund-Wikström B, Paulin L, Holm L, Auvinen P. (2014) Comparative genomics of *Lactobacillus crispatus* suggests novel mechanisms for the competitive exclusion of *Gardnerella vaginalis*. *BMC Genomics* 15:1070.

The publications are referred to in the text by their roman numerals.

Abbreviations

Cas CDS	CRISPR-associated protein coding DNA sequence
COG	clusters of orthologous groups
Contig	contiguous sequenced region
CRISPR	clustered regularly interspaced palindromic repeat
DAG	directed acyclic graph
DE	description line
EC	enzyme commission
emPCR	emulsion polymerase chain reaction
EPS	exopolysaccharide
GIT	gastrointestinal tract
GO	gene ontology
GRAS	generally regarded as safe
HMM	hidden Markov model
HMP	human microbiome project
LAB	lactic acid bacteria
MGE	mobile genetic element
ncRNA	noncoding RNA
NGS	next generation sequencing
OLC	overlap-layout-consensus
ORF	open-reading frame
PCR	polymerase chain reaction
PTS	phosphotransferase system
RBH	reciprocal best hit
RBS	ribosomal binding site
RSD	reciprocal smallest distance
S-layer	surface layer
Sn	sensitivity
Sp	specificity
SVM	support vector machine
TC	transporter classification system
WGS	whole-genome shotgun

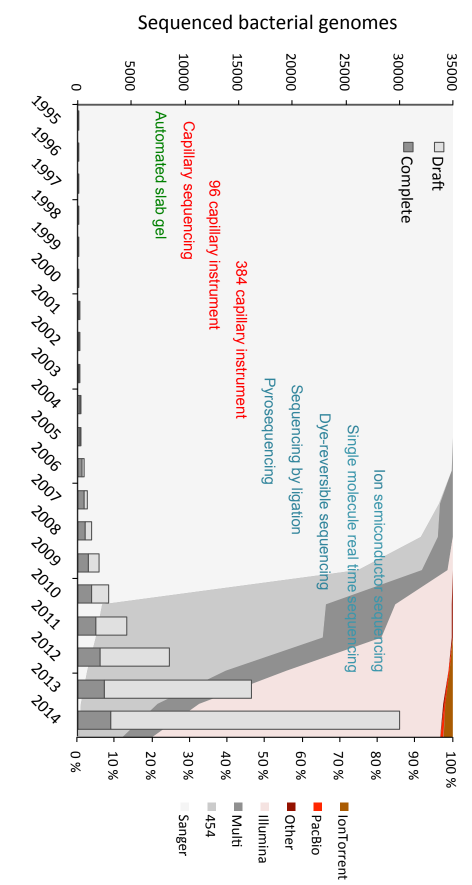
1 Review of the literature

This chapter introduces the objectives and scope of this dissertation and includes three sections: 1.1) a methodological review of DNA sequencing, sequence assembly and scaffolding, genomic feature calling, and functional classification techniques that are suitable for bacterial whole-genome sequencing; 1.2) an introduction to the general characteristics, phylogeny, taxonomy, and ecological distribution of lactobacilli; and 1.3) an overview of the existing literature on lactobacilli genomics.

1.1 Bacterial whole-genome sequencing

Whole-genome sequencing is the process of determining the order of nucleotides in the DNA molecules of an organism. Beginning with the first experimental determinations of DNA sequence using a location-specific primer extension strategy (Wu & Kaiser, 1968; Wu & Taylor, 1971) and following the introduction of the more practical 'plus and minus' (Sanger & Coulson, 1975), chemical degradation (Maxam & Gilbert, 1977), and chain-termination (Sanger *et al.*, 1977) sequencing strategies in the late 1970s, DNA sequencing has advanced to the point at which genomes can be sequenced rapidly and affordably (Mardis, 2008). This astounding growth in DNA sequencing capacity and speed has laid the foundation for determining the genomes of tens of thousands of bacterial (Figure 1) and thousands of other organisms in less than two decades and has permanently altered our understanding of microbial life.

Currently, the whole-genome shotgun (WGS) approach is the most widely used method for determining genome sequences (Anderson, 1981). When applied to microbial organisms (Figure 2), the initial standard approach is to grow the microbe from a single colony and then isolate a sufficient quantity of DNA for library construction. Depending on the protocol adopted, the amount used ranges from a few nanograms to tens of micrograms of DNA (Loman et al., 2012). In the second step, the DNA is fragmented into random overlapping DNA sequences that are used as templates for amplification and are sequenced from one (single-end reads) or both (paired reads) ends. For many years, the fragments were inserted into plasmids for cloning and then sequenced using the Sanger method. However, massively parallel sequencing methods are the current standard approach (Mardis, 2008). The third step of the process involves identifying overlaps between the reads and deriving contiguous consensus sequences from the reads, termed contigs. In most cases, however, full genome sequences cannot be built from a single shotgun read library. To improve the contiguity, paired reads from multiple libraries of different insert sizes are utilised or WGS reads are combined with mate-pair reads resulting from a process in which the ends of long (3-, 6-, or 8-kb) DNA fragments are brought together by circularisation and then sequenced across the ligation regions (Collins & Weissman, 1984). In addition, dedicated gap-closing techniques can be used to improve assembly contiguity (Figure 2). Previously, the closure of genomes was deemed



involving gel-based systems (green), capillary sequencing (red), and massively parallel DNA sequencing (blue). genome assemblies, respectively. Background surfaces indicate the fraction of genomes that have been sequenced using a particular sequencing technique. Others consist of genomes sequenced using the SOLiD or Complete Genomics platform. Labels indicate advances Figure 1. The cumulative number of bacterial genome entries in NCBI. Dark and light bars indicate the number of complete and draft necessary by the scientific community; however, this finishing phase is no longer routine (Figure 1) because of the prohibitive cost and labour required by the gap-closing phase. In future, the emergence of sequencing techniques that produce reads over 10 kb in length might provide a novel means to close genomes. The final step of the standard protocol is genome annotation, followed by analysis of the resulting information.

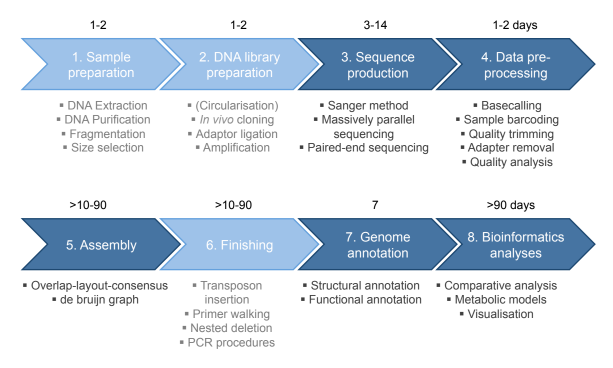


Figure 2. The principal steps involved in obtaining a bacterial genome sequence. In Sanger sequencing, genomic DNA is extracted from a single colony and then fragmented, ligated into a plasmid vector, and used to transform *Escherichia coli*. For each sequencing reaction, a single *E. coli* colony is selected, and the plasmid DNA is isolated. In massively parallel sequencing, common adaptors are ligated to fragmented genomic DNA, which is then subjected to PCR-based amplification and massively parallel sequencing. Following read preprocessing and assembly, unrevealed genome regions can be fixed using a variety of methods, wherein adjacent contigs are detected, and the gap between them is sequenced. Fixing gaps produces a finished genome. Genome annotation includes the identification of sequence features and the subsequent association of biological information with these features. These analyses are often followed by other computational analyses, such as whole-genome comparison, phylogenetic analysis, and metabolic network reconstruction.

1.1.1 DNA sequencing technologies

Six DNA sequencing technologies are currently used (Table 1). The Sanger method is the oldest of these and has been the workhorse of DNA sequencing for over 30 years (Sanger *et al.*, 1977; Mardis, 2008). In essence, the Sanger method uses mixtures of deoxy-

	Life technologies / Sanger 3730xl	Roche / 454's GS FLX+	lllumina / Hiseq2500	Life technologies / SOLiD 5500xl wildfire	Pacific Biosciences / PacBio RS II	Life technologies / lonTorrent PII
Released	1996	2005	2012	2006	2010	2014
Template preparation	PCR	Emulsion PCR on bead surface	BridgePCR on glass surface	Emulsion / Wildfire PCR	Single molecule	Emulsion PCR on bead surface
Sequencing chemistry	Dideoxy chain termination	Pyrosequencing	Reversible dye terminators	Sequencing by ligation and two- base coding	Sequencing by synthesis	lon semiconductor sequencing
Machine cost (\$)	95000	500000	740000	665000	700000	149000
Max read length (bases)	900	1000	125+125	50+50	11500	100
Fragments per run (M)	0.000096		4000	2400	0.04	320
Run time (h)		23	144	240	ω	4
Sequence yield (Gb/run)	0.0000864	0.7	1000	240	0.375	32
Sequence yield (Mb/h)	0.09	30	6944	1000	125	8000
Accuracy (%)	99.999	99.997	98.000	98.000	87.140	98.290
Operating cost (\$/Mb)	1111	8.86	0.03	0.02	1.07	0.01
Paired-end / Mate pairs / Multiplex	Yes / Yes / No	No / Yes / Yes	Yes / Yes / Yes	Yes / Yes / Yes	No / No / Yes	No / Yes / Yes
Pros	Read length, accuracy	Read length, rate of substitution errors	Throughput, operating costs	Accuracy, operating costs	Read length	Machine cost, no use of optics or fluorescence
Cons	Operating costs, throughput	Homopolymer errors, operating costs	Substitution errors	Color space data, read length	Operating costs, throughput, error rate	Homopolymer errors, throughput
Genome price (\$)	133333.3	1062.9	4.6	2.5	208.7	47.4
Genome coverage (×) / Number of genomes	0 / 0	233 / 5	333333 / 8333	80000 / 2000	125 / 3	10667 / 266

Table 1. Characteristics of widely-used DNA sequencing instruments (Liu *et al.*, 2012; Quail *et al.*, 2012). Genome price, genome coverage and number of genomes were computed assuming a bacterial genome of 3 Mb and a sequencing depth of ×40.

nucleotides and chain-terminating dideoxy-nucleotides to generate copies from the template that differ in length from each other by one nucleotide. In the process, the DNA sample is divided into four DNA sequencing reactions, which contain all four normal nucleotides, DNA polymerase, and one of four chain-terminating dideoxy-nucleotides in low amounts. As DNA synthesis progresses, DNA polymerase adds nucleotides to the chain. However, the occasional incorporation of a chain-terminating nucleotide into the strand causes DNA polymerase to cease DNA extension, resulting in fragments of different lengths (Sanger *et al.*, 1977). These DNA fragments are then denatured and separated according to mass using gel- or capillary electrophoresis, and the species of terminal base present is identified by exciting the fluorophore attached to the primer (Smith *et al.*, 1986) or chain-terminating base (Prober *et al.*, 1987) using a laser. Notably, incremental improvements to the Sanger method have rendered the technique advantageous for a number of applications, including the sequencing of long polymerase chain reaction (PCR) products and closing genomes by PCR.

Recently, a family of novel DNA sequencing technologies has supplanted the Sanger method (Mardis, 2008). These next-generation sequencing (NGS) platforms can process millions of DNA molecules in parallel and enable inexpensive and rapid sequencing, although at the expense of lower read length and accuracy. During template preparation, millions of template clusters, each comprising a large number of copies of a given template DNA molecule, are created using emulsion PCR (emPCR, Dressman et al., 2003) or bridge amplification (Adessi et al., 2000). Template aggregations are then sequenced in parallel using pyrosequencing (Margulies et al., 2005), sequencing by ligation (Shendure et al., 2005), sequencing by synthesis (Bentley, 2006; Bentley et al., 2008), or ion semiconductor sequencing (Rothberg et al., 2011). A notable exception is the single-molecule real-time sequencing system, which was developed by Pacific Bioscience (Eid et al., 2009) and involves sequencing single-template DNA molecules using DNA polymerases that are immobilised onto a zero-mode waveguide array. The key differences between the six sequencing technologies relate to the number and length of the reads produced (Table 1). In general, these two characteristics are negatively correlated, and technologies that provide long read lengths produce fewer data at higher cost than short-read sequencing instruments. However, the total sequence output from even the lowest capacity NGS instruments is far greater than the amount of sequence data needed to disclose a single bacterial genome; thus, these methods are an appealing choice for projects involving few isolates and benefiting from long contigs. In contrast, the Illumina and SOLiD platforms yield very large volumes of sequence data (Table 1) and are an affordable choice for large whole-genome projects that accommodate fragmented genome representations and for re-sequencing projects.

In terms of accuracy and error profiles, small but significant differences exist among the platforms (Mardis, 2008; Liu *et al.*, 2012; Quail *et al.*, 2012). The 454 and Ion Torrent platforms produce more nucleotide over- and under-calls than the other NGS platforms. These errors emerge from the methodology, which introduces all subsequent bases of one species at once, and from the difficulty in resolving the number of incorporated bases based on signal intensities. Especially troublesome are homopolymers of four or more bases (Voelkerding *et al.*, 2009). In contrast, the Illumina platform suffers from phasing,

fading, and crosstalk-generated noise (Erlich & Mitra, 2008). Phasing noise results from incorporation of none or more than one nucleotide during sequencing cycles and introduces lagging and leading nascent strands transmitting a mixture of signals; in contrast, fading noise arises from an exponential decay of the fluorescent signal intensity as a function of cycle number. The third noise factor arises from fluorophore crosstalk (Sheikh & Erlich, 2012). Regarding whole-genome sequencing, errors attributed to the Ion Torrent and 454 platforms are more fatal because the handling of insertion and deletion errors during genome assembly requires the use of computationally intensive gapalignments. Furthermore, insertion and deletion errors in the final contigs hamper gene calling and can cause fragmentation of predicted open reading frames (ORFs). The SOLiD platform, however, is considered accurate (Li *et al.*, 2012). Indeed, the investigation of each dinucleotide by two ligation reactions and the requirement that adjacent colour-calls must agree guarantees high accuracy. Nonetheless, the use of a colour-space coding scheme complicates data analysis, *de novo* assembly, and integration of the SOLiD data into other genomic resources (Li *et al.*, 2012).

1.1.2 Preprocessing of sequencing data

Modern DNA sequencing instruments generate large amounts of data that have to be preprocessed in several steps to convert them to a usable form (Figure 2). The recovery of human-readable read sequences from sequencing instrument data is referred to as base calling and is the typical first step toward usable data. It includes the transformation of intensity signals to nucleotide calls and the assignment of quality scores (indicating the reliability of the call) to each base (Sheikh & Erlich, 2012). In most cases, quality scores are reported in terms of logarithmically linked error probabilities termed Phred scores (Ewing & Green, 1998). Exceptions are the 454 and Ion Torrent error probabilities, which represent the likelihood that a base is an overcall and divide single quality values between two or more bases.

The adjustment of signal data for platform-specific anomalies is another important task that is performed by base callers (Sheikh & Erlich, 2012). Typically, this step is addressed by the use of a list of signal processing techniques, each tackling a specific error and error source. For example, most Illumina base-calling algorithms correct for fluorescent decay, fluorophore crosstalk, and errors caused by the incomplete removal or incorporation of reversible terminators (Kao *et al.*, 2009; Erlich & Mitra, 2008; Kircher *et al.*, 2009), whereas Sanger data is corrected for shifts in peak locations, fluorophore crosstalk, and background noise (Ewing *et al.*, 1998). Typically, a vendor-supplied base-calling method is used, because the base-calling process can require substantial amounts of processing time. However, third party programs have been developed as an alternative to vendor software and have been shown to improve the accuracy of base calls. Some popular third-party base-calling approaches for the Illumina (Rougemont *et al.*, 2008; Erlich & Mitra, 2008; Kao *et al.*, 2009; Kircher *et al.*, 2009), SOLiD (Wu *et al.*, 2010) and Roche 454 sequencing platforms (Quinlan *et al.*, 2008; Beuf *et al.*, 2012) are listed in Table 2 and Appendix Table 1.

databases is available in Appendix Table 1.	Appendix Table 1.
Category	Software
Base callers (Sanger)	Phred (Ewing & Green, 1998), KB Basecaller
Base callers (NGS)	Rolexa (Rougemont <i>et al.</i> , 2008), Alta-Cyclic (Erlich & Mitra, 2008), BayesCall (Kao <i>et al.</i> , 2009), Ibis (Kircher <i>et al.</i> , 2009), Pyrobayes (Quinlan <i>et al.</i> , 2008), HPCall (Beuf <i>et al.</i> , 2012), Rsolid (Wu <i>et al.</i> , 2010), All Your Base (Massingham & Goldman, 2012), BM-BC (Ji <i>et al.</i> , 2012).
Quality analysis and read manipulation	FastQC, PrinSeq (Schmieder & Edwards, 2011), BIGpre (Zhang et al., 2011), Cutadapt (Martin, 2011), fastx, Staden package (Staden et al., 1999), NGS QC Toolkit (Patel & Jain, 2012), NARWHAL (Brouwer et al., 2012)
Read correction	Reptile (Yang et al., 2010), HiTEC (Ilie et al., 2011), ECHO (Kao et al., 2011), Hybrid-SHREC (Salmela, 2010)
Greedy assemblers	SSAKE (Warren et al., 2007), VCAKE (Jeck et al., 2007)
Overlap-based genome assembly	Newbler (Margulies <i>et al.</i> , 2005), EDENA (Hernandez <i>et al.</i> , 2008), SGA (Simpson & Durbin, 2012), MIRA (Chevreux <i>et al.</i> , 2004)
De Bruijn graph based genome assembers	SPAdes (Bankevich et al., 2012), ALL-PATHS (Butler et al., 2008), SOAPdenovo (Li et al., 2010), Velvet (Zerbino & Birney, 2008), ABySS (Simpson et al., 2009), MaSuRCA (Zimin et al., 2013), RAY (Boisvert et al., 2010)
Reference-based genome assemblers	VAAL (Nusbaum <i>et al.,</i> 2008), Amos-Cmp (Pop <i>et al.,</i> 2004b)
Scaffolders	Bambus (Pop <i>et al.</i> , 2004a), SSPACE (Boetzer <i>et al.</i> , 2011), SOMA (Nagarajan <i>et al.</i> , 2008), OSLay (Richter <i>et al.</i> , 2007), BACCardl (Bartels <i>et al.</i> , 2005), PAGIT (Swain <i>et al.</i> , 2012)
Assembly integrators	Minimus2 (Sommer et al., 2007), MAIA (Nijkamp, et al., 2010), GAA (Yao et al., 2012)
Ab initio CDS predictors	Glimmer (Delcher <i>et al.</i> , 2007), GeneMark (Besemer <i>et al.</i> , 2001), EasyGene (Larsen & Krogh, 2003), Prodigal (Hyatt <i>et al.</i> , 2010), ZCURVE (Guo <i>et al.</i> , 2003)
Evidence-base CDS predictors	ORPHEUS (Frishman et al., 1998), CRITICA (Badger & Olsen, 1999)
CDS model integrators	Reganor (McHardy et al., 2004), YACOP (Tech & Merkl, 2003)
CDS model refinement tools	GenePRIMP (Pati <i>et al.</i> , 2010), Mugsy-Annotator (Angiuoli <i>et al.</i> , 2011), ORFCor (Klassen & Currie, 2013), MICheck (Cruveiller <i>et al.</i> , 2005)
ncRNA predictors	RNAmotif (Macke <i>et al.</i> , 2001), RNAmmer (Lagesen <i>et al.</i> , 2007), QRNA (Rivas & Eddy, 2001), RNAz (Washietl <i>et al.</i> , 2005), Aragorn (Laslett & Canback, 2004), tRNAscan-SE (Lowe & Eddy, 1997), Infernal (Nawrocki <i>et al.</i> , 2009), SRP-scan (Regalia <i>et al.</i> , 2002), Bcheck (Yusuf <i>et al.</i> , 2010), CMFinder (Yao <i>et al.</i> , 2006)
Intrinsic terminators	TransTermHP (Kingsford et al., 2007), RNIE (Gardner et al., 2011)

Ab initio protein functionCSS-Palm (Ren et apredictionGupta, 2008), SPAA	Context-based protein ContextMirror (Juan function prediction	Metabolism related PRIAM (Claudel-Re genes 2009), MEROPS (R:	Advanced functionBAGEL3 (van Heel of (Scaria et al., 2005), (Medema et al., 201	FunCut (Abascal & V General function (Vinayagam <i>et al.</i> , 2 classification 2005), InterProScan (Koskinen <i>et al.</i> , 201	Protein signature PFAM (Punta <i>et al.</i> , 2012), TigrFAN databases CDD (Marchler-Bauer <i>et al.</i> , 2011)	GenBank (Benson <i>e</i> Biological databases <i>et al.</i> , 2012), Rfam (<i>al.</i> , 2005)	Sequence database BLAST and PSI-BL/	Origin of replication Ori-Finder (Gao & Zhang, 2008)	Plasmids, integrative and conjugative elements, gene cassettes, integrons	Genomic islands (Vernikos & Parkhil)	Prophages ACLAME (Leplae et Mendez et al., 2008)	Insertion sequences ISfinder database (S	Repeats REPuter (Kurtz & Sc	CRISPR arrays CRT tool (Bland et a
CSS-Palm (Ren <i>et al.</i> , 2008), DISIS (Ofran <i>et al.</i> , 2007), MetalDetector (Lippi <i>et al.</i> , 2008), VirulentPred (Garg & Gupta, 2008), SPAAN (Sachdeva <i>et al.</i> , 2005)	ContextMirror (Juan et al., 2008), String (von Mering et al., 2005), Prolinks (Bowers et al., 2004)	PRIAM (Claudel-Renard <i>et al.</i> , 2003), KAAS (Moriya <i>et al.</i> , 2007), TCDB (Saier <i>et al.</i> , 2006), CAZy (Cantarel <i>et al.</i> , 2009), MEROPS (Rawlings <i>et al.</i> , 2004), REBASE (Roberts <i>et al.</i> , 2010)	BAGEL3 (van Heel <i>et al.</i> , 2013), RASTA-Bacteria (Sevin & Barloy-Hubler, 2007), TADB (Shao <i>et al.</i> , 2011), ARGO (Scaria <i>et al.</i> , 2005), MvirDB (Zhou <i>et al.</i> , 2007), ARDB (Liu & Pop, 2009), DBD (Wilson <i>et al.</i> , 2008), antiSMASH (Medema <i>et al.</i> , 2011), Mist2 (Ulrich & Zhulin, 2010), LOCP (Study I)	FunCut (Abascal & Valencia, 2003), CLAN (Kunin & Ouzounis, 2005), Gotcha (Martin <i>et al.</i> , 2004), GOPET (Vinayagam <i>et al.</i> , 2006), PFP (Hawkins <i>et al.</i> , 2006), ConFunc (Wass & Sternberg, 2008), SIFTER (Engelhardt <i>et al.</i> , 2005), InterProScan (Zdobnov & Apweiler, 2001), Argot2 (Falda <i>et al.</i> , 2012), BLANNOTATOR (Study II), PANNZER (Koskinen <i>et al.</i> , 2015), Sma3s (Muñoz-Mérida <i>et al.</i> , 2014)	PFAM (Punta <i>et al.</i> , 2012), TigrFAM (Haft <i>et al.</i> , 2003), HAMAP (Lima <i>et al.</i> , 2009), Interpro (Hunter <i>et al.</i> , 2012), CDD (Marchler-Bauer <i>et al.</i> , 2011)	GenBank (Benson <i>et al.</i> , 2013), UniProt (Bairoch <i>et al.</i> , 2008), PATRIC (Gillespie <i>et al.</i> , 2011), CharProtDB (Madupu <i>et al.</i> , 2012), Rfam (Griffiths-Jones <i>et al.</i> , 2003), COG (Tatusov <i>et al.</i> , 1997, Tatusov <i>et al.</i> , 2003), SEED (Overbeek <i>et al.</i> , 2005)	BLAST and PSI-BLAST (Altschul <i>et al.</i> , 1997), FASTA (Pearson & Lipman, 1988), HMMER3 (Eddy, 2011)	hang, 2008)	ICEberg (Bi <i>et al.</i> , 2012), INTEGRALL (Moura <i>et al.</i> , 2009), ACID (Joss <i>et al.</i> , 2009), cBar (Zhou <i>et al.</i> , 2010)	SIGI-HMM (Waack <i>et al.</i> , 2006), IslandViewer (Langille & Brinkman, 2009), PAI-IDA (Tu & Ding, 2003), Alien_Hunter (Vernikos & Parkhill, 2006), IslandPick (Langille <i>et al.</i> , 2008)	ACLAME (Leplae <i>et al.</i> , 2004), ProphageDB (Srividhya <i>et al.</i> , 2007), PHAST (Zhou <i>et al.</i> , 2011), Prophinder (Lima- Mendez <i>et al.</i> , 2008), PhiPsy (Akhter <i>et al.</i> , 2012), Phage_Finder (Fouts, 2006)	ISfinder database (Siguier et al., 2006), IScan (Wagner et al., 2007), ISsaga (Varani et al., 2011)	REPuter (Kurtz & Schleiermacher, 1999), RepeatScout (Price et al., 2005)	CRT tool (Bland et al., 2007), PILER-CR (Edgar, 2007), CRISPRFinder (Grissa et al., 2007)

(Darzentas, 2010)	Visualization
Genome atlas (Wassenaar et al. 2010) ACT (Carver et al. 2005) Combo (Engels et al. 2006) Circoletto	
IMG (Markowitz <i>et al.</i> , 2012), RAST (Aziz <i>et al.</i> , 2008), DOE-JGI MAP (Mavromatis <i>et al.</i> , 2009), CG pipeline (Kislyuk <i>et al.</i> , 2010), ERGO (Overbeek <i>et al.</i> , 2003), PGAAP, Broad Institute, BCM, JCVI (Tanenbaum <i>et al.</i> , 2010)	Annotation pipelines
Pathway tools (Karp <i>et al.</i> , 2002), FMM (Chou <i>et al.</i> , 2009), MetaCyc (Krieger <i>et al.</i> , 2004), UniPathway (Morgat <i>et al.</i> , 2012), KEGG (Kanehisa <i>et al.</i> , 2004)	Metabolic reconstruction
PhyML (Guindon et al., 2003), BioNJ (Gascuel, 1997), Phylip (Felsenstein, 1989)	Phylogenetic trees
GERP (Cooper et al., 2005), SiPhy (Garber et al., 2009)	Prediction of constrained elements
MUMmer (Darling et al., 2004; Darling et al., 2010), TBA (Blanchette et al., 2004), Pecan (Paten et al., 2008), Mauve aligner (Rissman et al., 2009), Gepard (Krumsiek et al., 2007), BLASTatlas (Wassenaar et al., 2010)	Whole-genome aligners
Muscle (Edgar, 2004), ProbCons (Do et al., 2005), Clustal Omega (Sievers et al., 2011)	Multiple sequence aligners
RBH (Tatusov <i>et al.</i> , 1996), RSD (Wall <i>et al.</i> , 2003), RIO (Zmasek & Eddy, 2002), OrthoStrapper (Storm & Sonnhammer, 2002), InParanoid (Remm <i>et al.</i> , 2001), EggNOG (Jensen <i>et al.</i> , 2008), OrthoMCL (Li <i>et al.</i> , 2003), Protein cluster (Klimke <i>et al.</i> , 2009), OMA (Roth <i>et al.</i> , 2008), PoFF (Lechner <i>et al.</i> , 2014), morFeus (Wagner <i>et al.</i> , 2014) 2014)	Orthology prediction
PSORTb v3.0 (Yu <i>et al.</i> , 2010), tatP (Bendtsen <i>et al.</i> , 2005), LipoP (Rahman <i>et al.</i> , 2008), SignalP (Petersen <i>et al.</i> , 2011), TMHMM (Krogh <i>et al.</i> , 2001), LocateP (Zhou <i>et al.</i> , 2008), EffectiveT3 (Arnold <i>et al.</i> , 2009), CoBaltDB (Goudenège <i>et al.</i> , 2010)	Subcellular location

Quality assessment is another important pre-processing step, which aims to pinpoint poor quality reads and typically includes the visualisation of base quality scores, sequence length distributions, and nucleotide distributions. In addition, sequence data can benefit from data cleaning and steps such as adapter and poor quality region trimming and the filtering of chimeric and short reads and other types of sequence artefacts. For example, the removal of bases with poor quality at the end of the reads has a positive impact on downstream analyses (Haridas *et al.*, 2011; Cox *et al.*, 2010). Adaptor cutting has also been shown to be advantageous and ensures that only the relevant part of the read is passed to and considered at the downstream analysis.

An alternative strategy for improving data quality is based on high base coverage and supposedly infrequent and random sequencing errors (Yang *et al.*, 2013). In this setting, sequencing errors are detected by aligning reads to a reference genome and by examining the alignment for uncommon base calls or, in its more generalised form, by decomposing reads into overlapping oligomers of the length k (*i.e.*, k-mers) and identifying infrequent k-mers that resemble frequent k-mers (Pevzner *et al.*, 2001; Yang *et al.*, 2013). Typically, k-mers that occur only a few times in the data are considered in this spectral alignment approach to represent errors and are rectified by making a minimum number of nucleotide edits to the reads from which they emerged. At present, several spectral alignment methods have been developed that mainly differ with regard to choosing the coverage threshold for the identification of infrequent k-mers. However, this feature is important because overly low thresholds result in uncorrected errors, whereas overly high thresholds affect correct k-mers. Among various spectral alignment tools, Reptile (Yang *et al.*, 2010), HiTEC (Ilie *et al.*, 2011) and ECHO (Kao *et al.*, 2011) have been shown to reliably detect and correct erroneous reads (Yang *et al.*, 2013).

1.1.3 Genome assembly

Genome assembly is the process of assembling short reads into the largest possible continuous sequences, thus providing a representation of the expected genome (Pop, 2009; Flicek & Birney, 2009; Miller *et al.*, 2010). In essence, this process relies on the assumption that highly similar reads originate from the same position within a genome and involves joining individual overlapping reads into contigs (Pop, 2009; Flicek & Birney, 2009; Miller *et al.*, 2010). In addition, the process can include a separate scaffolding step to produce larger scaffold structures comprising an ordered set of contigs with intervening gaps representing DNA stretches that are not present in the reads. The sequential phases of current genome assembly methodology are illustrated in Figure 3.

The first phase of genome assembly is contig construction. Currently, this is achieved in most cases using algorithms that are based on either the overlap-layout-consensus (OLC) or the de Bruijn graph approaches (Pop, 2009; Flicek & Birney, 2009). Alternatively, genome sequences can be reconstructed using the greedy extension approach, in which the best-matching reads are iteratively joined together into contigs until no more reads or contigs can be joined (Miller *et al.*, 2010). Many early assemblers relied on the greedy approach, as do some modern ones (Warren *et al.*, 2007; Jeck *et al.*, 2007); however, this approach is no longer utilised due to the inherently local assembly process, which can become stuck at a local minimum if the sequence under assembly is extended with a read that would have been more beneficial to other joining operations (Pop, 2009; Miller *et al.*, 2010). Finally, contig building can rely on aligning reads to the reference sequence and on grouping reads by continuity (Pop *et al.*, 2004b; Nusbaum *et al.*, 2008). However, the comparative approach is valid only in the presence of a reference genome with substantial sequence similarity (\geq 90%) to the genome of interest (Pop *et al.*, 2004b). Table 2 and Appendix Table 1 provide partial lists of the tools that are currently available for genome assembly and scaffolding.

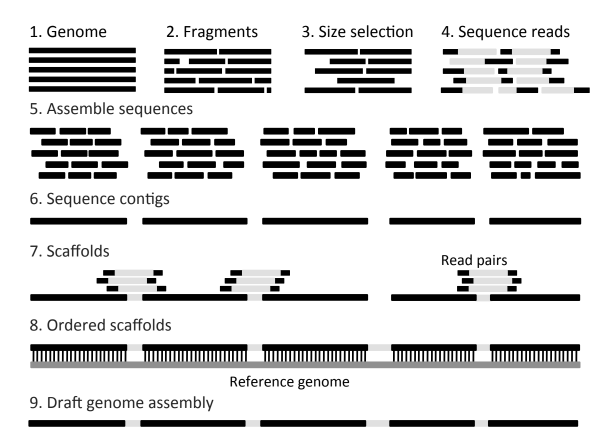


Figure 3. The sequential phases of current genome assembly methodology. Starting from a large amount of genomic DNA, the DNA is sheared (2) into random fragments, size selected (3), and amplified and sequenced from one or both ends (4). The sequencing reads are then assembled on the basis of sequence overlaps (5), thereby yielding sequence contigs (6). The contigs can then be oriented and ordered based on the read pairs that map to two contigs (7) or with the aid of reference genomes (8).

The OLC approach is one of the two main approaches used and is useful for the assembly of long reads (Pop, 2009; Miller *et al.*, 2010). The OLC process begins with an all-against-all read comparison. The reads and relationships between reads are then structured into a graph with a node for every read and an edge between any pair of reads

that overlap sufficiently well (Pop, 2009; Miller *et al.*, 2010). Using this structure, finding contigs becomes equivalent to finding a path through a graph that visits each node exactly once, termed a Hamiltonian circuit. The OLC approach is highly suitable for reads of varying length because it can employ all of the information obtained from long and short reads (Miller *et al.*, 2010). This approach captures repetitive regions in nodes with multiple connections, thus providing a means by which to exclude these regions and effectively handle sequencing errors. Regardless, identifying the Hamiltonian circuit is an intensive task, and storing each read in a separate node requires memory, thus making the OLC approach somewhat impractical for modern sequencing projects (Pop, 2009; Miller *et al.*, 2010; Flicek & Birney, 2009). An exception is the SGA assembly package, which exhibits significantly reduced memory requirements and computing time (Simpson & Durbin, 2012).

The second widely used assembly approach is known as the De Bruijn graph approach and is considered ideal for short read-length, high-coverage data (Pevzner *et al.*, 2001). This strategy relies on deconstructing reads into short k-mer fragments before assembling them into contigs with the help of De Bruijn graphs, in which nodes represents unique kmers and an edge connects all node pairs that overlap by k-1 bases (Pop, 2009; Miller et al., 2010; Flicek & Birney, 2009). In theory, the number of unique k-mers in the genome determines the number of nodes, thereby allowing the use of a small memory footprint for genomes of limited size and in the presence of repetitive elements. Moreover, De Bruijnbased assemblies are fast to compute because overlaps between the reads are implicitly captured by the graph rather than computed individually; furthermore, a linear-time algorithm exists for finding the Eulerian paths that visit each edge once (Fleischner, 1990). This approach is also considered useful for short-read data. Theoretically, 16-mers should yield reasonable assemblies, and larger k-mers should entail longer contigs; however, parameter optimisation tests have shown that the optimal performance is obtained using moderate (approximately 50) to large (approximately 100) k values (Peng et al., 2012; Jünemann et al., 2014) or a combination of multiple k-mers (Peng et al., 2012; Bankevich et al., 2012). The greatest disadvantage of the De Bruijn graph paradigm is its insensitivity to repetitive regions and read errors. The use of even large k-mers fails to resolve repeat regions, if repeats are longer than the given k-mer value. In the presence of such repeats, the graph will have branching vertices and contain multiple Eulerian paths. In these cases, continuity can however be improved by adding constrains to the graph and finding Eulerian superpaths that traverse the graph via predetermined sub-paths (Pevzner et al., 2001; Pop, 2009). Erroneously called bases near read ends on the other hand result in dead-end "tips", whereas errors in the middle lead to branching nodes and alternate paths termed "bubbles" that start and terminate in the same nodes, each providing an equally good solution. To compensate for these problems, many assemblers employ the spectral alignment approach to correct reads before beginning the assembly (Pevzner et al., 2001; Li et al., 2010; Butler et al., 2008). This approach has been shown to remove up to 66% of the k-mers (Li et al., 2010). The concept of De Bruijn graphs is implemented in several assemblers; of these, ALL-PATHS (Butler et al., 2008) and SPAdes (Bankevich et al., 2012) have performed well in recent genome assembler surveys (Earl et al., 2011; Magoc et al., 2013).

Repetitive regions that are longer than the read length are problematic for genome assembly and cannot be resolved by simply investigating read overlaps regardless of the coverage depth. The position of these repeats and other unrevealed genome regions in the genome can, however, be approximated by supplementing the assembly process with a scaffolding phase in which individual contigs are joined into larger sequence structures that comprise contigs and the intervening gaps (Pop, 2009; Miller et al., 2010). Chiefly, scaffolding exploits information that is derived from paired reads (libraries with insert sizes of approximately 300 to 1,000 bp) or mate-pair reads (libraries with insert sizes of 3-, 6-, or 8-kb; Figure 3). Pairs of contigs that are likely to reside side-by-side in the genome are detected by aligning reads to contigs and by identifying read pairs that match different contigs. The optimal order and orientation of contigs are then solved using heuristic or graph-based approaches that rely on majority voting from a large number of read pairs (Pop, 2009; Miller et al., 2010). Importantly, the length of the gaps can be estimated based on the positions of the paired reads in contigs and expected insert sizes (Pop, 2009). The major disadvantage of this approach is related to the quality of the data and to the difficulty in minimising inconsistency between the assembled contigs and read pair constraints; for this reason, the joining of contigs requires multiple coherently aligned read pairs (Li et al., 2010; Simpson et al., 2009).

Alternative approaches for organising contigs involve the use of PCR-assisted or other gap-closing techniques (Figure 2), optical maps, reference genomes, and multiple genome assemblies. Optical mapping is a technique for generating whole-genome ordered restriction endonuclease maps (Samad et al., 1995). These optical maps not only provide information about restriction fragment sizes, but also provide information regarding the order in which these fragments occur in the DNA. Regarding scaffolding, matches between in silico restriction-digested sequences and fragments in the optical map provide means for stitching contigs together (Nagarajan et al., 2008). The use of reference genome alignments is another method for organising contigs into larger units (Richter et al., 2007; Rissman et al., 2009; Bartels et al., 2005). This approach is increasingly common as the number of sequenced genomes in databases increases; however, the accuracy of the scaffolds depends on the alignment quality and the level of sequence similarity between the contigs and reference genomes. Indeed, even closely related reference genomes are useless in the presence of horizontal transfer and genomic rearrangements. Finally, scaffolds can be constructed by integrating assemblies. For example, methods such as Minimus (Sommer et al., 2007), MAIA (Nijkamp et al., 2010) and GAA (Yao et al., 2012) make use of multiple assemblies to produce meta-assemblies with increased contiguity and accuracy.

1.1.4 Structural annotation

Structural annotation is the aspect of genome annotation that consists of the identification of genomic features (Koonin & Galperin, 2003; Angelova *et al.*, 2010). In theory, this can be achieved using experimental techniques but is typically attained using computational approaches followed by manual curation due to time and cost. This chapter introduces

bioinformatic strategies for calling CDSs, ncRNA genes, and MGEs in bacterial genomes. Table 2 and Appendix Table 1 list some popular computational tools for structural annotation.

CDS calling is one of the most important steps of structural annotation (Angelova et al., 2010). In its simplest form, CDS calling could consist of scanning genomes for sufficiently long (\geq 90 bases) uninterrupted stretches of DNA between a start codon and a stop codon. However, the screening of such ORFs can yield a certain number of incorrect gene predictions (Koonin & Galperin, 2003) because true CDSs often echo long ORFs in neighbouring reading frames. An alternative strategy is to take advantage of the statistical properties of the coding sequences and estimate the coding potentials of ORFs or to infer genes based on the similarity of the encoded protein sequences to those of other proteins in public database (Koonin & Galperin, 2003). Ab initio programmes make predictions using a probabilistic model that distinguishes CDSs from noncoding sequences based on sequence composition and estimates the coding potentials of ORFs. The model employs, for example, the infrequencies of Gs and As at the first codon positions, the infrequency of Gs at the second codon position, and/or amino acid composition differences between coding and noncoding genome regions. Typically, the parameters used in probabilistic models are trained from separately prepared training datasets, which comprise sufficiently long and non-overlapping ORFs from the sequence in question (Delcher et al., 2007), ORFs from the sequence in question showing homology to known proteins in public databases (Larsen & Krog, 2003), and sequences of a known type. An alternative to the *ab* initio prediction of genes is to search the target genome for CDSs that are similar to extrinsic evidence (Koonin & Galperin, 2003). Extrinsic methods include the BLAST-type mapping of ORFs against known gene products and gene callers such as ORPHEUS (Frishman et al., 1998) and CRITICA (Badger & Olsen, 1999), which infer CDSs based on coding potential and sequence similarity. Finally, software packages exist for refining gene call anomalies (Pati et al., 2010; Cruveiller et al., 2005) and combining evidence from individual gene-finding systems into consensus CDS models (Tech & Merkl, 2003; McHardy et al., 2004).

To date, no systematic analysis of the gene calling accuracies of different gene finders is available. However, comparisons of methods given in the original papers show that bacterial gene finder algorithms boast an average accuracy of 90% or better (Delcher *et al.*, 2007; Hyatt *et al.*, 2010; Besemer *et al.*, 2001). Moreover, function assignments can be attached to most gene products, indicating that bacterial gene callers are likely accurate. The major hurdles in the use of modern *ab initio* gene callers involve the identification of short genes (\leq 150 bp), over-annotation, the false prediction of pseudo-genes, and the presence of longer than anticipated overlaps between CDS calls (Besemer *et al.*, 2001; Delcher *et al.*, 2007; Hyatt *et al.*, 2010). The scarcity of stop codons in GC-rich genomes can also impair the accuracy of gene callers that give value for gene length in estimating coding potentials (Hyatt *et al.*, 2010). It is also possible that the probabilistic model may fail in genomic islands with atypical base compositions. In contrast, evidence-based methods can ignore novel CDSs and have longer runtimes than *ab initio* methods, which scan millions of bases in minutes. Evidence-based gene callers are nonetheless useful in calling CDSs that are ignored by *ab initio* gene finders and that exhibit homology to known proteins. For example, current genome annotations appear to lack, on average, 30 *bona fide* genes that can be identified using a BLAST procedure (Warren *et al.*, 2010).

Another standard analysis in the process of structural genome annotation is the identification of tRNA, rRNA, and other types of ncRNA genes that function directly as RNA rather than being translated to proteins. Traditionally, ncRNA genes are called using comparative genomics methods (Eddy, 2002; Pichon & Felden et al., 2008). In the case of rRNAs, genes can be called using primary sequence similarity with known rRNA genes (Lagesen et al., 2007). However, other types of ncRNAs often lack common statistical signals in their primary sequences that could be exploited for detection. Instead, their calling often requires methods that use sequence and structural conservation, such as the tRNA prediction system tRNAscan-SE or the general ncRNA search suite Infernal. Both of these algorithms use covariance models to capture the primary consensus and secondary structure information of an RNA family and are very accurate (Lowe & Eddy, 1997; Nawrocki et al., 2009). However, this performance comes at the expense of runtime, and the analysis requires a template ncRNA structure. The more general approaches for identifying ncRNA genes rely on genomic variations in sequence composition statistics, predict transcripts without long ORFs and with initiation and termination sites, search for sequences that have the ability to adopt given secondary structure patterns (Macke et al., 2001), or use a combination of RNA structure prediction and comparative sequence analyses to test for a characteristic signal (Rivas & Eddy, 2001; Washietl et al., 2005). These programmes are considered useful for defining the structure of a sequence that is already known to be an RNA gene but are largely immature for use in genome-wide scans (Eddy, 2002; Pichon & Felden, 2008).

Approximately one tenth of a bacterial replicon is intergenic. Although this portion of the genome is commonly referred to as noncoding, it contains a variety of important sequence features (Madigan et al., 2010). Intergenic regions contain, for example, transcriptional regulatory elements and basal promoter elements that are key players in gene regulation. Such regions are also rich in repetitive elements and contain motifs contributing to the coordination of replication, cell division, DNA segregation, and DNA repair (Touzain et al., 2010). Also visible in the genome are mRNA stem-and-loop structures that control of gene expression. Depending on the type of stem-and-loop structure, these motifs can be called with the help of general ncRNA annotation algorithms and by inferring genomes for rho-independent terminators using software tools such as RNIE (Gardner et al., 2011) and TransTermHP (Kingsford et al., 2007). Intriguingly, stem-and-loop structure annotations provide valuable clues about genome structure and enable the marking of operon endpoints and start sites, because rho-independent terminators are mainly located at transcription termini and transcriptional attenuators are located between the basal promoter elements and the start codons of the 5'-most genes of operons. Finally, intergenic regions can be annotated for CRISPR arrays. Because CRISPR arrays consist of short (approximately 20-50 bp) direct repeats that are interspaced by variable sequences called spacers, they are routinely inferred using repeat finding algorithms or their modified versions (Bland et al., 2007; Edgar, 2007; Grissa et al., 2007).

MGEs are DNA segments that encode proteins that mediate the movement of DNA within genomes or between bacterial cells. These sequence features can have a tremendous impact on the transfer, recombination, and deletion of host genes, and traces of MGE activity are present in nearly all bacterial genomes (Frost *et al.*, 2005). MGEs are typically annotated based on their similarity to the known MGE members. This paradigm is implemented in the IScan (Wagner et al., 2007) and ISsaga (Varani et al., 2011) methods, enabling the identification of insertion elements using curated references from the ISfinder database (Siguier et al., 2006). This approach can also be used to identify clusters of genes that exhibit similarity to known phage genes (Lima-Mendez et al., 2008; Zhou et al., 2011), integrons and gene cassettes (Moura et al., 2009; Joss et al., 2009), and integrative and conjugative elements (Bi et al., 2012). Furthermore, some computational resources for genomic island annotation build on sequence similarity searches (Langille et al., 2008). Alternatively, the automated detection of MGEs can rely on sequence composition characteristics. The methods included in this category are based on the notion that MGEs are often acquired horizontally and that MGEs can be identified by searching for local variations in sequence composition, such as variations in the G+C ratio and dinucleotide bias. Several programmes for the automated detection of MGEs adopt this approach, including those designated for the annotation of plasmids (Zhou & Xu, 2010), phage-like regions (Srividhya et al., 2007), and genomic islands (Tu & Ding, 2003; Waack et al., 2006; Vernikos & Parkhill, 2006). Although these bioinformatic tools are also able to call novel MGEs, sequence composition skew appears to be a less reliable predictor than sequence similarity. For example, genomic island detectors that rely on base composition statistics exhibit less agreement with a dataset of known genomic islands than their homology-based counterparts (Langille et al., 2010).

1.1.5 Protein function prediction

Protein function prediction can be defined as the inference and assignment of specific biological and biochemical roles to proteins (Koonin & Galperin, 2003). This stage of genome annotation attempts to compile a definitive catalogue of the protein functions of the organism and provides researchers with specific testable hypotheses about the roles of proteins in the cell. This knowledge is critical for understanding life at the molecular level (Koonin & Galperin, 2003; Rost *et al.*, 2003; Friedberg, 2006; Valencia, 2005). However, genome-scale protein function prediction is a challenging process (Rost *et al.*, 2003; Valencia, 2005; Friedberg, 2006; Schnoes *et al.*, 2009) and involves the use of many functional classification schemes and computational procedures (Table 2 and Appendix Table 1), as discussed in detail below. As with structural genome annotation, the use of automated prediction methods is preferably performed in conjunction with manual annotation.

Functional classification schemes are used to capture biological knowledge in a form that is suitable for computational processing. Regarding bacteria, the functions of proteins are most often available as description lines (DEs, Bairoch *et al.*, 2008). These natural language function labels are the traditional way of describing functional information and

can be very informative. However, DEs, like the natural language itself, are rife with synonyms and ambiguity, making comparison of DEs notoriously difficult (Friedberg, 2006). Another challenge with regard to DEs such as 'DNA gyrase subunit B' is that protein function is a subjective concept, and different researchers may denote the functions of proteins differently (Friedberg, 2006). In addition to DEs, standardised functional labelling schemes have been developed to unify information about protein function. These schemes employ only certain words and punctuation and describe functional information in a controlled and computationally amenable fashion. Such resources include TIGRFAM (Haft et al., 2003), SEED (Overbeek et al., 2005), the keyword catalogues of UniProt (Bairoch et al., 2008), and orthologous groups of proteins (COG) (Tatusov et al., 1997; Tatusov et al., 2003) that cover general functional aspects and providing a means by which to overview and compare the functional contents of organisms. Additional classification schemes have also been developed for the classification of enzymes (Webb, 1992), enzyme-related functions (Rawlings et al., 2004; Roberts et al., 2010; Cantarel et al., 2009), and transporters (Saier et al., 2006). Noteworthy examples include the Enzyme Classification (EC, Webb, 1992) and Transporter Classification (TC, Saier et al., 2006) systems, which follow a hierarchical structure that allows the measurement of the functional similarity of genes. Another intensively used functional classification scheme is Gene Ontology (GO, Ashburner et al., 2000). Machine-readable GO encompasses three ontologies that describe three aspects of function: molecular function, biological process, and cellular location. These ontologies are non-redundant and are implemented as a directed acyclic graph (DAG) that represents general terms as nodes near the root of the ontology and specific terms as nodes near the leaves of the ontology. Differing from the hierarchical structure, a node can have multiple parents. By definition, if a gene is associated with a term, it is associated with all of its broader level GO terms. Furthermore, each GO term assignment has an evidence code attributed to it, thereby providing a means of separating computationally derived information from manually curated information (Ashburner et al., 2000). Although GO is the most widespread functional classification scheme in use today, its use in bacterial genome annotation is limited by the low number of specific GO descriptions that have been associated with bacterial proteins.

The conventional approach for protein functional classification involves the transfer of biological information on the basis of sequence similarity. The rationale for this approach is that sequences with a high degree of similarity are likely to have evolved from a common ancestor and will also share functional roles (Friedberg, 2006). This type of homology-based functional classification has been shown to be rather accurate and to be able to assign a function to approximately 73% of CDSs in the average genome (Raes *et al.*, 2007); however, this approach is not without problems. Principally, sequence similarity does not imply similarity of function (Eisen, 1998), and transferring annotations based on sequence similarity can propagate existing annotation errors (Schnoes *et al.*, 2009). This analysis method also fails with regard to orphan genes and often results in a set of differently characterised sequences. It also appears that the quality of the prediction depends on the sequence database used (Schnoes *et al.*, 2009), the sequence similarity search method used, and the type of function information to be transferred (Clark &

Radivojac, 2011). Another shortcoming is the need to separate useful and spurious sequences. Some studies have proposed that as low as 30% amino acid sequence identity is required for assigning complete enzyme function (Valencia, 2005; Devos & Valencia, 2001) and that inference by similarity performs approximately equally well for all matches above this sequence identity threshold (Clark & Radivojac, 2011; Altenhoff *et al.*, 2012). In contrast, other researchers argue that below a 70% sequence identity threshold, EC numbers start to diverge rapidly, such that at 30% sequence identity, only a tenth of pairs of proteins share all EC numbers (Rost, 2002; Rost *et al.*, 2003). It appears that the latter conclusions are more accurate regarding bacterial enzymes (Figure 4). Among the manually annotated proteins entries available in the UniProt database, the proportion of pairs in which both proteins are described using the same enzyme code became high above the 50% sequence identity level, whereas at the 30% identity level, only every fifth protein pair was observed to share the same enzyme class.

The functional classification of proteins on a genome-wide scale typically involves a direct sequence-sequence comparison of query proteins against large sequence databases using BLAST (Altschul et al., 1997), FASTA (Pearson & Lipman, 1988), or PSI-BLAST (Altschul et al., 1997). The function of the prototype sequence is then transferred to the protein under consideration. In its simplest form, predictions are based on information that is associated with the top hit (*i.e.*, the best-BLAST approach) or the most informative description. Based on Figure 4, this approach is accurate provided that it is performed for highly similar enzymes. More elaborate methods have also been proposed that make use of more than one sequence (Abascal & Valencia, 2003; Kunin & Ouzounis, 2005, Martin et al., 2004; Vinayagam et al., 2006; Hawkins et al., 2006; Wass & Sternberg, 2008) or that transfer functions within an evolutionary context (Engelhardt et al., 2005; Zmasek & Eddy, 2002; Storm & Sonnhammer, 2002). According to the method comparisons provided in the original papers, the pooling of information over multiple sequences and the additional use of weakly similar sequences as the source of function information appear to be beneficial and increase prediction accuracy. An alternative but fundamentally similar approach is to search for sequence-based signatures. Examples of tools that are designed for finding signatures and resources that archive signature profiles are listed in Table 2 and Appendix Table 1. Overall, the use of protein signatures appears to be more powerful and sensitive for detecting remote homologues than the use of pairwise sequence similarity search tools. Nevertheless, the combination of several methods for functional annotation is recommended because the most suitable method for a particular sequence set cannot be known *a priori*.

Homology transfer is also the principal method for identifying proteins with specific and predetermined functional roles, such as signal transduction system proteins, virulence factors, and adhesins. For example, function-specialised databases have implemented the possibility of conducting a BLAST-like search against their high-quality sequence sets. Some representative examples in the field of microbiology include databases and services that were developed for mining bacteriocins (van Heel *et al.*, 2013), carbohydrate-active enzymes (Cantarel *et al.*, 2009), *cas* genes (Haft *et al.*, 2005), signal transduction system proteins (Ulrich & Zhulin, 2010), antibiotic resistance factors (Scaria *et al.*, 2005; Zhou *et al.*, 2007; Liu & Pop, 2009), type II Toxin-antitoxin systems (Sevin & Barloy-Hubler,

2007; Shao *et al.*, 2011), pilus components (developed in Study I), restriction modification system components (Roberts *et al.*, 2010), DNA-binding transcription factors (Wilson *et al.*, 2008), and secondary metabolite biosynthetic loci (Medema *et al.*, 2011).

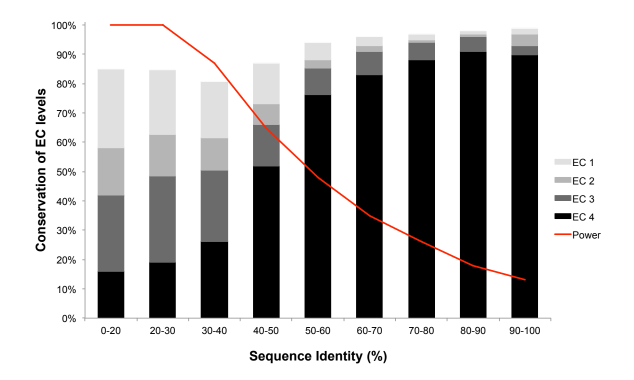


Figure 4. Conservation of EC number and the power of homology transfer for bacterial enzymes. Protein sequence data was extracted from the UniProt database, and only entries that had been verified either at the RNA level or at the protein level were accepted. Sequence-level identity was quantified using BLAST (default settings), and the fraction of pairwise sequence matches sharing a certain number of EC number digits was tabulated at different levels of pairwise sequence identity. The red line indicates the fraction of sequences that match other sequences at the given or a higher identity range.

Apart from evolutionary counterparts, functional classification can rely on the genomic context of genes. This approach utilises the co-localisation and/or co-evolution of genes to predict functional linkages between encoded proteins and is also practical for genes that match only other uncharacterised genes because the transfer of functional information between organisms is not necessary. Instead, the use of context information provides an opportunity for also transferring information between genes within a single organism. The four major methods for context-based prediction are the phylogenetic profile method (which employs the fact that functionally associated genes appear to be preserved or eliminated in concert during evolution; Pellegrini *et al.*, 1999), the gene fusion method (which relies on gene pairs that occur in parallel as a larger composite gene in other genomes; Marcotte *et al.*, 1999), the gene neighbour method (which searches for genes

with preserved physical genomic proximity; Dandekar *et al.*, 1998), and the co-evolution method (which uses the correlation between phylogenetic trees; Juan *et al.*, 2008). Among these methods, the phylogenetic profile method is considered the most precise and provides comparable performance to homology-based functional classification (von Mering *et al.*, 2005).

Ab initio protein function prediction methods predict protein function based on sequence information alone. Unlike other functional prediction methods, these methods are resistant to existing error-prone information in databases and have the advantage of being also suitable for orphan proteins as the analysis does not involve transfer of function from unreliable sources (Ofran et al., 2005; Punta & Ofran, 2008). However, ab initio methods currently provide only rough approximations of various aspects of function. Nevertheless, promising methods exist for the prediction of subcellular localisation and transmembrane topologies of proteins (Table 2 and Appendix Table 1); these methods providing valuable clues and rather accurate predictions about bacterial secretomes that comprise secreted or surface-associated proteins. With regard to bacterial genome annotation, other aspects of protein function that can be predicted using *ab initio* methods include amino acids that are involved in DNA binding (Ofran et al., 2007) and binding to various metals (Lippi et al., 2008), virulence-associated factors (Garg & Gupta, 2008), the probability of a protein being an adhesin (Sachdeva et al., 2005), and palmitoylation sites in protein sequences (Ren et al., 2008). In addition, specialised bioinformatic resources are available that locate proteins that are secreted by type III (Arnold *et al.*, 2009) or IV (Burstein et al., 2009) secretion systems.

1.1.6 Summarisation of genome annotation results

The summarisation and visualisation of genome data are key parts of any genome project and one of the first tasks to be performed following annotation. Typically, genome data are visualised to obtain an overview and to compare genome annotations. This step is also an important quality control step and can reveal assembly errors and genome regions that may need attention. Picture-based structural DNA analysis (e.g., the visualisation of GCskew, AT-content, GC-content, and gene-strand bias) can also be used to identify genomic features such as MGEs and origins of replication. Tools for data visualisation are listed in Table 2 and include standalone genome browsers, such as ACT (Carver et al., 2005), and web services, such as the BLASTatlas (Wassenaar et al., 2010). Additionally, genome annotation results can be recapitulated into genomic feature tables that, typically, provide information about the genome size, GC-content sequence elements, and coding density of the organism in question and of other, closely related organisms. These tables can also highlight the biological aspects of the organism. Genome size is, for example, an indicator of adaptive potential; large rRNA and tRNA gene numbers suggest a short doubling time, and codon usage can be used to shed light on the organism's likely environmental niche (Wassenaar et al., 2010).

1.1.7 Comparative genomics

Whole-genome comparisons are a powerful approach for understanding genomic diversity and the relatedness of organisms (Ali et al., 2013). This approach can reveal fascinating differences and similarities between genomes, is a prerequisite for profiling rapidly evolving sequences (Cooper et al., 2005; Garber et al., 2009), and provides a means to classify species and to describe horizontal gene transfer events (Darling et al., 2004). The two underlying paradigms that are used to identify regions of similarity are local and global sequence alignment (Frazer et al., 2003). The first of these reports all similar subregions of the sequence and also identifies homology in the presence of rearrangements. However, local alignment cannot suggest how the subregions have evolved from their ancestors (Frazer et al., 2003). In contrast to local alignments, global alignments describe an end-to-end alignment of sequences. Aligned regions need to be conserved in both order and orientation (Frazer et al., 2003); thus, this approach is optimal for the comparison of genomes with a high degree of synteny, but it is less good for outlining genomic relatedness of organisms in the presence of horizontal transfer and genomic rearrangements. Nevertheless, both alignment strategies can be useful, especially when explored graphically using comparative alignment viewers such as Combo (Engels et al., 2006) and ACT (Carver et al., 2005). Intuitive visualisation of the whole-genome homology can also be achieved by using dot-plot visualisation software (Krumsiek et al., 2007) and by mapping and visualising genome homology of genes and proteins within a reference strain in comparison to other prokaryotes (Wassenaar et al., 2010). Additionally, whole-genome comparisons can base on advanced genome alignment methods (Darling et al., 2010; Blanchette et al., 2004; Paten et al., 2008; Dubchak et al., 2009, Angiuoli et al., 2011; Rissman et al., 2009) that often mix global and local alignment procedures and can align long sequences while detecting the presence of inversions, translocations, duplications, and gains and losses. However, the use of even the most sophisticated software entails the selection of many mundane parameters (Frith et al., 2010), and these methods perform best with sequences that exhibit significant nucleotide-level similarity and colinearity.

The identification of orthologue groups is a central part of functional classification (Li *et al.*, 2003; Koonin & Galperin, 2003; Sonnhammer & Koonin, 2002) and underpins the delineation of phenotype-genotype relationships among bacteria by providing a means of listing genes that are present only in the genomes of those isolates that express the phenotype of interest (Korbel *et al.*, 2005). In general, methods for constructing orthologue groups are classified into distance and tree-based methods. Distance-based methods include the reciprocal best hit (RBH, Tatusov *et al.*, 1996) and reciprocal smallest distance (RSD, Wall *et al.*, 2003) approaches. These methods build on BLAST scores or maximum likelihood estimations of evolutionary distances and they resolve orthologous (*i.e.*, homologues that have evolved by speciation from a single ancestral gene) between two organisms by finding two-way best genome-wide similarities. The software InParanoid extends this concept further (Remm *et al.*, 2001) and exploits the RBH strategy to identify orthologues that arise from duplication after speciation (*i.e.*, inparalogues).

The rationale behind this approach is that orthologues and inparalogues are more likely to perform the same function than outparalogues that result from duplication preceding the speciation event (Sonnhammer & Koonin, 2002). Distance-based methods have also been proposed for finding orthologue groups across multiple genomes. These approaches typically use pairwise sequence similarity search methods and cluster RBHs that span multiple genomes using triangular linkage clustering (Tatusov et al., 1997; Jensen et al., 2008), the Markov clustering procedure (Li et al., 2003), maximum weight cliques (Roth et al., 2008), or fully connected sub-graphs (Klimke et al., 2009). Alternatively, orthologue detection can be based on tree reconstruction and on gene and species tree reconciliation (Zmasek & Eddy, 2002; Storm & Sonnhammer, 2002). Regardless, assessments of orthologue prediction tools have resulted in contradictory results. One study concluded that EggNOG is the best and OrthoMCL the worst prediction method (Trachana et al., 2011); however, another study endorsed OrthoMCL and InParanoid over other methods (Chen et al., 2007). Confusingly, a recent study proposed that the RBH method might be more accurate and specific than any complex method (Salichos & Rokas, 2011). Overall, phylogenetic approaches are powerful for capturing evolutionary relationships; however, the computational costs of multiple sequence alignment, the lack of an accurate species trees for a given collection of bacteria, and the complexity of tree reconciliation preclude the use of this approach for the genome-wide identification of orthologues (Rentzsch & Orengo, 2009; Trachana et al., 2011).

Molecular phylogenetic analyses are frequently used to depict the evolutionary history of a given set of organisms (Williams & Sarah, 2014). In addition, they can be applied to a single gene family, with each copy of the gene in each organism being included in the analysis (Yang & Rannala, 2012). The reconciliation of the gene tree with a species tree can then be used to time gene gains, duplications, and losses (Eisen, 1998). Some methods have even been proposed for the genome-scale inference of gene gains and deletions and for the inference of ancestral gene inventories (Mirkin et al., 2003). Relatedness among organisms has typically been estimated by comparing molecular sequences, mostly smallsubunit ribosomal RNAs (Woese, 1987) or ubiquitous housekeeping genes (Konstantinidis & Tiedje, 2005). The process begins with the extraction of sequences from all species under examination. After obtaining a multiple sequence alignment for the sequences, several phylogenetic methods can be used to infer the phylogeny. These methods can be broadly classified into maximum parsimony, maximum likelihood, Bayesian inference, and distance methods (Yang & Rannala, 2012), which differ largely in the way in which they choose the best among all possible trees. Maximum parsimony selects the tree that requires the minimum number of character changes from the common ancestral sequences. In maximum likelihood methods, the probability that a certain tree with a set of parameters produces a given set of data is computed, and the tree that makes the given data most probable is chosen. In Bayesian analysis, inferences of phylogeny are based upon the posterior probabilities of phylogenetic trees, whereas distance-based methods calculate the evolutionary distance among sequences of interest and construct a distance matrix that is used to cluster sequences hierarchically (Yang & Rannala, 2012). Additionally, organism-level evolutionary histories can be estimated from genome-scale datasets. These methods are believed to generate a more accurate picture of evolution,

especially for bacteria that have a high incidence of gene movement from one lineage to another (Brown *et al.*, 2001) and include the derivation of phylogenies from gene content (Snel *et al.*, 1999), gene order (Korbel *et al.*, 2002), and compositional signatures (Fox *et al.*, 1980). Further, phylogenies can be determined on the basis of multiple source-trees (Sanderson *et al.*, 1998) or concatenated single-copy orthologue sequences (Brown *et al.*, 2001; Rokas *et al.*, 2003).

1.1.8 Metabolic and regulatory reconstructions

Metabolic reconstructions provide a starting point for the study of the metabolism of an organism and attempt to include all of the relevant metabolic information of an organism (Oberhardt et al., 2009; Durot et al., 2009). In general, the metabolic information of an organism is represented in terms of metabolic pathways, each of which describes a set of chemical interactions and transformations for the conversion of compounds (Durot et al., 2009). Metabolism-related data can also be conceptualised, and pathways can be integrated into genome-wide networks that usually envision metabolites as nodes and reactions as edges of the network. Although metabolic networks are not generated as often as metabolic pathways in bacterial genome studies, this approach offers a realistic view of metabolism, whereby, in theory, any reaction can have implications for other reactions (Oberhardt et al., 2009; Durot et al., 2009). Typically, metabolic pathway and network reconstructions stem from genome annotations (Oberhardt et al., 2009). Initially, a preliminary list of metabolic and transport reactions relevant for the given organism are harvested; gene name, TC number and EC number assignments and transmembrane protein predictions being the main information sources. Metabolic pathways and networks are then formed by linking individual reactions into increasingly complex structures. Reaction sets can be assembled *ab initio* by tracking the movement of atoms through the network (Arita, 2004; Heath et al., 2010) or by projecting the metabolic data onto reference metabolic pathways (Moriya et al. 2007; Karp et al., 2002). Finally, the initial metabolic networks are refined based on literature information, manual inspection, experimental data, and bioinformatic tools that annotate inconsistencies between metabolic models and substrate utilisation predictions and that can resolve metabolites or reactions that are disconnected from the rest of the metabolism. For example, Pathway Tools can include missing reactions in pathways if a significant fraction of the remaining reactions of the pathways are supported by genome annotations (Karp et al., 2002). In addition to reconstructing metabolic networks, genome annotation allows the reconstruction of transcriptional regulatory networks (Barabási & Oltvai, 2004). These networks provide a global picture of the transcriptional machinery of the cell and are constructed by integrating existing knowledge of regulons, operons, and transcriptional regulation interactions with the results of operon and transcription factor binding site screens (Baumbach et al., 2009; Ravcheev et al., 2013). Bioinformatics resources that are relevant to the study of metabolism and transcriptional regulation are listed in Table 2 and Appendix Table 1.

1.1.9 Genome annotation pipelines

Several genome annotation systems that are intended for the automated, in-depth annotation of prokaryotic genomes have been designed and presented in recent years, including the IMG (Markowitz et al., 2012), RAST (Aziz et al., 2008), DOE-JGI MAP (Mavromatis et al., 2009), CG pipeline (Kislyuk et al., 2010), ERGO (Overbeek et al., 2003), and JCVI (Tanenbaum et al., 2010) genome annotation pipelines. Some of these systems are completely automatic online services that have the advantage of simplicity, whereas others are standalone tools that require maintenance but provide an extra level of confidentiality. Some popular bacterial genome annotation systems are listed in Table 2 and Appendix Table 1. Typically, genome annotation pipelines involve a wide array of methods, through which they identify genomic features and assign functional information that describe the biological role of these features. For example, the annotation service of the Broad Institute calls ncRNA genes and CDSs using three automated methods for each. Gene models are clustered, and representative models are selected using heuristics, such as the relative overlap with BLAST hits. Finally, gene models with problems are filtered out (as described in the human microbiome web page; Nelson et al., 2010). The JCVI genome annotation system (Tanenbaum et al., 2010) is another popular annotation system and shares similarities with that of the Broad Institute. CDS prediction employs one ab initio and two evidence-based prediction methods, and the pipeline runs three RNA gene callers over the genome: tRNAScan-SE, ARAGORN, and BLAST searches against Rfam. Other modern annotation pipelines call genes in a highly similar fashion. After gene calling, most bacterial genome annotation systems search the set of predictions against one or more protein databases using BLAST (Markowitz et al., 2012; Aziz et al., 2008; Mavromatis et al., 2009; Kislyuk et al., 2010; Overbeek et al., 2003; Tanenbaum et al., 2010). Further, the gene products are usually also searched via InterProScan against a set of sequence profile databases. In addition to these basic analyses, some annotation pipelines include additional analysis modules and perform protein subcellular localisation prediction (Kislyuk et al., 2010; Tanenbaum et al., 2010; Markowitz et al., 2012) and reconstruct metabolic pathways (Aziz et al., 2008). Some annotation pipelines also have viewers that permit users to rectify old calls and introduce new gene and function calls. Surprisingly, modern genome assemblers and assembly pipelines are scarce in bacterial genome annotation systems; the CG pipeline is the sole exception (Kislyuk et al., 2010). Although the various annotation servers are largely based on the same bioinformatic tools, pipelines appear to produce rather different annotation results. Importantly, the one study that has systematically compared the results of annotation services has documented distinct differences in annotation outputs (Bakke et al., 2009).

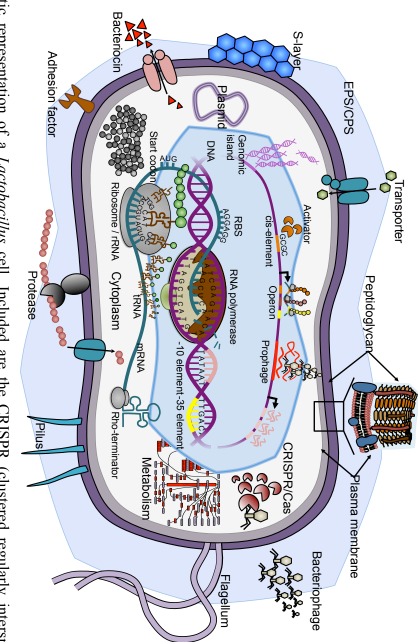
1.2 Lactobacilli

The genus *Lactobacillus* comprises a large, heterogeneous group of gram-positive, nonsporulating, rod-shaped bacteria that have complex nutritional requirements and a low GC-content genome (less than 50 mol%). These bacteria are acid-tolerant, aero-tolerant or anaerobic, and aciduric or acidophilic and employ a strictly fermentative metabolism (Hammes & Vogel, 1995; Felis & Dellaglio, 2007; Salvetti et al., 2012). They are part of the lactic acid bacteria (LAB) group, which is characterised by the production of lactic acid as the main by-product of carbohydrate fermentation (Kandler & Weiss, 1986). In general, lactobacilli are encountered in an array of plant-, food-, and animal-related habitats that are rich in carbohydrates (Pot et al., 1994; Hammes & Vogel, 1995; Felis & Dellaglio, 2007; Salvetti et al., 2012). Some species, such as Lactobacillus iners, are restricted to specific niches, whereas others demonstrate a notable ability to adapt to a diverse set of environments. Lactobacilli also have a beneficial effect on our daily life and are of great economic importance. Several species are encountered on and in the human body, including the oral cavity, gastrointestinal tract (GIT), and vagina (Hammes & Vogel, 1995; Walter, 2008; Salvetti et al., 2012); other species are essential in the fermentation of food, beverage, and feed products (Leroy & Vuyst, 2004; Bernardeau et al., 2006; Giraffa et al., 2010). Recently, members of Lactobacillus have been added to dietary and dairy products for probiotic purposes and to offer benefits for health and wellbeing (Saxelin et al., 2005; Giraffa et al. 2010).

1.2.1 Cellular characteristics of lactobacilli

The cell structure of lactobacilli is typical for that of a gram-positive bacterium and is often organised into three basic architectural regions: the cytoplasm, cell envelope, and surface appendages (Figure 5). The innermost region is the cytoplasm, which is largely the site of metabolism and replication. The cytoplasm contains the nucleoid, which is an irregularly shaped and non-membrane bound region that contains the chromosomal DNA, ribosomes, which are essential for protein synthesis, and an array of non-coding RNA molecules (ncRNA), which are not translated into proteins but function at the RNA level (Madigan et al., 2010). Also included in the cytoplasm are various proteins that are responsible for important functions, such as forming structural components (structural proteins), the catalysis of biochemical reactions (enzymes), and the transmission of molecular signals (transcription factors and signal transducers). The innermost compartment can also carry one or more independently replicating extra-chromosomal DNA molecules (plasmids) that are not essential for survival but can comprise a notable fraction of the total genome (Madigan et al., 2010). Of note, a sizable part of the genome is present as fragments of DNA that are capable of moving around within the genome or between genomes. A wide variety of such mobile genetic elements (MGEs) have been characterised in a number of different lactobacillus strains and include plasmids (Claesson et al., 2006), genomic islands (genome regions that exhibit evidence of horizontal origins; Kleerebezem et al., 2003), transposons (Callanan et al., 2008), and prophages (lysogenic phages that can switch under some conditions to a lytic lifestyle and then infect other bacteria; Ventura et al., 2006).

The cell envelope is a structural compartment that protects the cytoplasm and mediates interactions with the host and environment. In gram-positive lactobacilli, the cell envelope contains a cytoplasmic cell membrane and a thick peptidoglycan layer that is decorated



are attached to surfaces), and flagella (which are involved in cell movement). Genomic islands, clusters of co-transcribed genes repeats)-Cas (CRISPR-associated) adaptive immunity system (which predominantly targets foreign genetic material), bacteriocins Figure 5. Schematic representation of a Lactobacillus cell. Included are the CRISPR (clustered regularly interspaced palindromic in width RNA hairpin formation) are also indicated in the figure. The cells are between 1 and 10 microns in length and are approximately 1 micron (operons), prophages, *cis*-elements (DNA-binding sites for transcription factor proteins), -10 and -35 elements (attachment sites for σ -(proteinaceous antibacterial compounds that inhibit the growth of mainly closely related bacteria), pilus fibres (by which bacterial cells factor of RNA polymerase), ribosomal binding sites (RBSs), and rho-terminator sites (which control transcription termination through

with proteins, teichoic acids, and polysaccharides (Lebeer *et al.*, 2008; Madigan *et al.*, 2010). In addition, some lactobacilli express an outermost coat, the Surface-layer (S-layer), which is composed of a single protein that completely encases the cell and appears to regulate bacterial contacts with the human dendritic cell (Konstantinov *et al.*, 2008). Some lactobacilli also surround themselves with an exopolysaccharide (EPS), which is either tightly associated with the cell wall or secreted into the surroundings (Lebeer *et al.*, 2008). Interestingly, these ubiquitous components of the cell envelope of lactobacilli have been shown to exert some immune responses and are therefore attractive candidates as probiotic effector molecules (Lebeer *et al.*, 2008). Finally, molecular and genomic studies of lactobacilli have characterised the presence of surface appendages in lactobacilli as being involved in movement (flagella; Forde *et al.*, 2011) or adhesion to surfaces (sortase-dependent pili and fimbriae; von Ossowski *et al.*, 2011; Lebeer *et al.*, 2012; Pridmore *et al.*, 2004). The cell envelope also contains membrane-associated proteins that are responsible for such processes as nutrient acquisition, adhesion, cell communication, microbe-host interactions, and stress sensing (Lebeer *et al.*, 2008).

1.2.2 Sugar fermentation

Lactobacilli exhibit a strong ability to degrade various carbohydrates and derive energy mainly from the conversion of sugars into lactic acid. In general, these bacteria degrade hexoses through homo- or heterofermentative carbohydrate fermentation pathways (Hammes & Vogel, 1995; Pot et al., 1994). First, homofermentative lactobacilli use glycolysis (Embden-Meyerhof-Parnas pathway) to ferment hexoses primarily into lactic acid; most Lactobacillus species fall into this group (Salvetti et al., 2012). In comparison, obligately heterofermentative lactobacilli ferment hexoses and pentoses via the pentose phosphate pathway. During this process, half of the substrate is converted into lactic acid and the rest is metabolised in equimolar amounts to ethanol (or to acetic or formic acid) and carbon dioxide. Finally, facultatively heterofermentative species utilise both pathways and are almost as common as homofermentative lactobacilli species (Salvetti et al., 2012). Genetically, the three modes of sugar fermentation are explained by the absence or presence of genes encoding aldolase and phosphoketolase. Aldolase is present in homofermentative and facultatively heterofermentative species; phosphoketolase is present in obligately and facultatively heterofermentative lactobacilli (Salvetti et al., 2012).

1.2.3 Taxonomy

The genus *Lactobacillus*, as currently circumscribed, contains over 152 species that exhibit wide phenotypic and genotypic variation. The genus is polyphyletic with the genus *Pediococcus* and is the largest genus within the family *Lactobacillaceae*, which in turn belongs to the order Lactobacillales, class Bacilli, and phylum Firmicutes. According to the most recent systematic study (Salvetti *et al.*, 2012), *Lactobacillus* species can be

subdivided into 29 distinct phylogenetic groups (Figure 6). These groups represent a distinct cluster in the 16S rRNA gene phylogeny and were named in the study according to the first recognised species of the given group. Some economically and biomedically important phylogenetic *Lactobacillus* groups are introduced below.

The Lactobacillus delbrueckii group is the largest of the phylogenetic Lactobacillus groups (Salvetti et al., 2012) and contains many species that are essential in food production. For example, *L. delbrueckii* is widely used as a starter culture in yoghurt manufacturing, whereas Lactobacillus helveticus is important in the manufacture of a range of Swiss- and Italian-type cheeses (Leroy & De Vuyst, 2004; Giraffa et al., 2010). Included in the group are also GIT-associated species, such as Lactobacillus acidophilus and Lactobacillus johnsonii (Altermann et al., 2005; Pridmore et al., 2004), and species such as Lactobacillus crispatus, Lactobacillus jensenii, and L. iners, which are major constituents of the healthy adult female urogenital tract and important agents of urogenital health (Ma et al., 2012; Martin, 2012). Notably, the L. delbrueckii group includes several commercially distributed probiotic strains that appear to benefit health (Saxelin et al., 2005; Giraffa et al. 2010).

The *Lactobacillus salivarius* group is a heterogeneous group of 16 homofermentative and 9 facultatively heterofermentative species. The GC content within this group varies widely from 32 to 47 mol% (Figure 6), reflecting the fact that the members of this group occupy a wide variety of habitats, including human saliva, vertebrate intestine, soil, water, plants, and food (Forde *et al.*, 2011). Some species of the group appear to be motile and contain genes that encode the flagellar apparatus (Forde *et al.*, 2011).

The Lactobacillus casei group comprises three species; namely, L. casei, Lactobacillus paracasei, and Lactobacillus rhamnosus. These species have a broad ecological distribution and are frequently found in plant material as well as in the oral cavity and GIT of humans and animals (Kandler & Weiss, 1986). Industrially, L. paracasei, L. rhamnosus, and L. casei have applications as acid-producing starter cultures for milk fermentation and as starter adjunct cultures for the intensification and acceleration of flavour development in bacterial-ripened cheeses (Broadbent et al., 2012; Mäyrä-Mäkinen & Bigret, 1998). Selected strains, such as L. rhamnosus GG, L. casei Shirota, and L. casei DN114-001, are commercially important probiotic that are added to various products for their potential to enhance the health of humans (Saxelin et al., 2005; Giraffa et al., 2010; Siezen & Wilson 2010).

The *Lactobacillus reuteri* group contains 15 species that cover a broad host range. These species have been isolated from foods such as rye-bran fermentations and sourdough, and some are frequent in the GIT of birds, pigs, mice, and rats (Forde *et al.*, 2011; Frese *et al.*, 2011). *L. reuteri* is also considered indigenous to humans according to some investigations (Walter, 2008).

The *Lactobacillus sakei* group contains four facultatively heterofermentative species. The best known of the four species is *L. sakei*, a psychrotrophic bacterium that is found naturally on fresh meat and fish and that is used widely in their fermentation (Chaillou *et al.*, 2005). The *Lactobacillus plantarum* is another well-studied phylogenetic lactobacilli group. Out of its five facultatively heterofermentative species, the most noteworthy is the genetically heterogeneous *L. plantarum*, which exhibits remarkable ecological adaptability

and can be recovered from a variety of habitats including fermented foods, vegetables, and the human GIT (Siezen *et al.*, 2010; Siezen & van Hylckama Vlieg, 2011). Of note, some strains of *L. plantarum* have probiotic applications (Kleerebezem *et al.*, 2003).

	Metabolism	GC content (%)	Growth T (°C)	Growth pH
0.02	C/B	39-46	10-45	3-9
L. senioris	С	42-42	15-45	4-8
69 L. collinoides-group (5)	С	40-49	4-50	4-9
<u>94</u> L. brevis-group (10)	C/B	44-55	15-37	5-7
L. malefermentans	С	41-42	10-37	4-7
L. fructivorans-group (5)	C/B	35-42	15-35	3-8
60 Pediococcus genus (12)	-	-	-	-
L. kunkeei	С		15-37	4-8
97 L. ozensis	С	41-41	20-30	-
<u>100</u> L. plantarum-group (5)	В	44-47	10-40	4-7
99 L. alimentarius-group (11) A/B	34-41	8-45	3-9
83 L. manihotivorans-group	(3) A	48-59	15-45	4-8
L. camelliae	А	52-52	15-15	8-8
L. sharpeae	А	53-53	15-30	-
L. pantheris	A	52-53	10-37	5-8
¹⁰⁰ L. thailandensis	А	49-49	15-15	8-9
L. brantae	В	-	25-40	-
L. saniviri	В	49-49	10-37	4-9
100 L. casei-group (3)	В	45-47	10-45	5-5
100 L. sakei-group (4)	В	41-44	0-42	5-5
69 L. vaccinostercus-group	(3) C	35-41	4-40	3-8
100 L. rossiae	С	44-44	15-37	
L. siliginis	С	46-46	20-37	5-5
99 <i>L. reuteri</i> -group (15)	C/B	38-56	15-45	4-8
95 L. composti	В	47-47	10-37	4-9
45 L. coryniformis-group (3)) В	45-45	13-37	4-8
100 L. perolens-group (3)	B/C	49-56	10-45	4-9
L. selangorensis	А	45-47	15-37	5-5
100 L. concavus	А	46-47	10-42	4-8
L. dextrinicus	С	-	32-45	5-8
L. algidus	В	36-37	0-25	-
76 L. salivarius-group (25)	A/B	32-47	10-45	3-8
L. floricola	А	48-48	20-30	-
96 L. delbrueckii-group (27)	A/B/C	33-51	15-52	3-8
Eggerthia catenaformis	-	-	-	-
100 Kandleria vitulina	-	-	-	-

Figure 6. A phylogenetic tree illustrating the evolutionary relationship between *Lactobacillus* and *Pediococcus* species based on 16S rRNA gene sequence similarity. The tree was calculated using Tamura Three Parameters as the distance matrix formula and minimum evolution as the tree reconstruction method. The scale bar represents the number of substitutions per site. Bootstrap values are reported in percentages at nodes if $\geq 60 \%$. Clusters containing more than three species were condensed and given the name of the first species described. The number of species in each group is indicated in parentheses, followed by the fermentation mode (A, homofermentative; B, facultatively heterofermentative; and C, obligately heterofermentative), GC content, temperature growth range, and pH growth range. The tree and phenotype data were adapted from Salvetti *et al.*, 2012.

1.2.4 Industrial applications

The large-scale production of dairy products such as cheeses, yogurts, and fermented milks is the best-known industrial application of lactobacilli (Leroy & De Vuyst, 2004; Bernardeau *et al.*, 2006; Giraffa *et al.*, 2010). However, lactobacilli also play an important role in other types of food fermentation (du Toit *et al.*, 2010, Leroy & Vuyst, 2004; Bernardeau *et al.*, 2006) and are used for the production and preservation of foods of plant (*e.g.*, pickles, olives, sauerkraut, sourdough bread, and Korean kimchi) and animal (*e.g.*, fermented and dry sausages, salami, and fermented fish) origin, even though lactobacilli can in some cases cause spoilage of meat and seafood products (Varnam, 2002). These processes typically employ lactobacilli as starter cultures that are added to a raw material to accelerate and drive fermentation. Alternatively, preservation can be based on microbes that are naturally present in the raw material (Pfeiler & Klaenhammer, 2007; Leroy & De Vuyst, 2004). However, the use of spontaneous fermentation results in less control over the fermentation process and varying product quality.

Lactobacilli are mainly used in food production for the fermentative conversion of sugars into organic acids, mostly lactic acid. This reduces the sugar content of the product and acidifies the raw material below the pH ranges within which most food-spoilage microbes can grow (Leroy & De Vuyst, 2004). Acid production can also influence the organoleptic properties of the final product by contributing to the coagulation of milk proteins (Heller, 2001; Giraffa *et al.*, 2010). In addition to acids, lactobacilli are known to produce a variety of other compounds of relevance to food industry. Among these are aroma compounds, lipases, and proteases that play a role in flavour development in cheese (Steele *et al.*, 2012), EPS molecules that can enhance the mouthfeel of yogurts (Vuyst & Degees, 1999), and proteinaceous antibacterial peptides (*i.e.* bacteriocins) that can be used as natural food preservatives (Leroy & De Vuyst, 2004). Some lactobacilli are also able to produce health-enhancing ingredients (*e.g.*, vitamins, bioactive peptides, and antioxidants) that have potential applications as biotherapeutic agents (Saxelin *et al.*, 2005; Giraffa *et al.*, 2010).

Chiefly, lactobacilli are generally regarded as safe (GRAS) organisms and have a long and safe history of application and consumption in the production of fermented foods. Under rare and unusual circumstances, the consumption of *Lactobacillus* products has been associated with infections in humans (Bernardeau *et al.*, 2008). However, the risk of *Lactobacillus* infection was estimated in that study to be unequivocally negligible, with approximately only one case per 10 million people over more than a century. In addition to food-related applications, lactobacilli are used to ferment animal feeds, to produce chemicals (Saxena *et al.*, 2009), to produce antibiotics, and as live vaccine carriers (Giraffa *et al.* 2010).

1.2.5 Lactobacilli in and on animals and humans

Lactobacilli are closely associated with humans and animals. In humans, they are notably abundant in breast milk (Collado *et al.*, 2009), the oral cavity (Walter, 2008), and the

mucosal surfaces of the vagina (Ravel *et al.*, 2011). Importantly, *Lactobacillus* species that are prevalent in breast milk, such as *L. gasseri* and *Lactobacillus fermentum* (Martín *et al.*, 2005), or in the vagina, such as *L. crispatus*, *Lactobacillus gasseri*, *L. iners*, and *L. jensenii* (Ravel *et al.*, 2011), are often regarded as offering a number of benefits for health and wellbeing (Martín *et al.*, 2005; Martin, 2012).

The human GIT is another site that hosts lactobacilli (Molin *et al.*, 1993; Ahrne *et al.*, 1998; Reuter, 2001; Vaughan et al., 2005; Dal Bello & Hertel, 2006; Walter, 2008; Ryan et al., 2008; Matsuda et al., 2009) and at least 12 Lactobacillus species have been associated with the human GIT (Table 3). However, the Lactobacillus species composition varies among subjects, and approximately 25% of human faecal samples lack lactobacilli entirely (Walter, 2008). Furthermore, in faecal samples in which lactobacilli are detected, these organisms are found at low levels and account for a minor portion (approximately 0.01% to 0.6%) of the microbiota present (Lebeer et al., 2008). It is therefore believed that only a small number of GIT-associated species are genuine residents of the GIT, and most are considered simply as allochthonous members that are passing through the GIT after originating from fermented food, the oral cavity, or more proximal parts of the GIT (Walter, 2008). In contrast, the presence of lactobacilli in animals and their GITs is more pronounced than in humans. For example, lactobacilli form stable populations in the GITs of pigs, mice, rats, and chickens at sites that are lined with stratified squamous epithelium (Walters, 2008). Although this epithelium type is absent from the human GIT, it is present in the buccal and vaginal cavities, sites at which lactobacilli are abundant (Walter, 2008).

Table 3. *Lactobacillus* species that are commonly detected in food, human faeces, and in various parts of the human GIT. The data were compiled from Molin *et al.*, 1993; Ahrne *et al.*, 1998; Reuter, 2001; Vaughan *et al.*, 2005; Dal Bello & Hertel, 2006; Walter, 2008; Ryan *et al.*, 2008; and Matsuda *et al.*, 2009. Brackets indicate the overall size of the resident lactobacillar populations according to Walter, 2005.

	Food	Oral cavity (<10 ³ -10 ⁶)	Stomach (<10 ³)	Small intestine (<10 ³ -10 ⁸)	Colon (<10 ³ -10 ⁹)	Faeces (<10 ³ -10 ⁹)
L. acidophilus	+	+				+
L. crispatus		+				+
L. casei	+	+				+
L. fermentum	+	+				+
L. gasseri		+	+	+		+
L. paracasei	+	+			+	+
L. plantarum	+	+			+	+
L. reuteri	+		+	+		+
L. rhamnosus	+	+		+	+	+
L. ruminis			+			+
L. sakei	+					+
L. salivarius		+				+

1.2.6 Probiotic lactobacilli

By definition, probiotics are live microorganisms that have a beneficial effect on the health of the host when administered in adequate amounts (Lee & Salminen, 1995; Saxelin et al., 2005; Giraffa et al. 2010). Although in theory any microorganism could be identified as probiotics, most probiotics in use today are lactobacilli and bifidobacteria (Siezen & Wilson, 2010). However, not all members of these genera are equally useful as probiotic additives. Preferably, the strains used should be of human origin and should have been proven safe for consumption (Lee & Salminen, 1995). In addition, probiotic additives should tolerate bile and acid to survive passage through the upper GIT and adhere to human tissues (Giraffa et al. 2010). Commercially distributed strains also should not adversely affect taste and should exhibit good growth characteristics and survive the production and storage processes used (Lee & Salminen, 1995). The last characteristic is particularly important because a bacterial concentration of $\geq 10^8$ colony-forming units per gram appears to be the efficacious dosage (Aureli et al., 2011). Although the list of functional requirements is lengthy, at least some Lactobacillus strains appear to fulfil these criteria; see Table 4 for selected examples. These strains are typically delivered to customers as fermented dairy product or dietary supplements (Saxelin et al., 2005; Giraffa et al. 2010); vaginal suppositories, cereals, and skin lotions representing other examples of products to which probiotic lactobacilli have been added (Krutmann, 2009; Rivera-Espinoza & Gallardo-Navarro, 2010; Salvatore et al., 2011).

Many types of health benefit have been associated with the consumption of probiotic lactobacilli. These include the treatment of gastrointestinal infection and inflammatory bowel disease, the prevention of respiratory tract infections, the treatment of atopic diseases and allergy, the suppression of Helicobacter pylori infection, the prevention of urinary tract infections, anti-diarrheal properties and the treatment of bacterial vaginosis (Saxelin et al., 2005; Uehara et al., 2006; Anukam et al., 2006; Siezen & Wilson, 2010; Aureli et al., 2011). The potential mechanisms of action involved include a strengthening of the cell barrier function, vitamin supply, the enhancement of healthy microbiota, and antagonism against pathogens via the production of antimicrobials and competitive exclusion (Lebeer et al., 2008; Ventura et al., 2008; Oelschlaeger, 2010). Many probiotics are also believed to secrete immunomodulatory molecules that interact directly with the host (Yan et al., 2007; Lebeer et al., 2008). The physical interaction of cell-surface components with host tissues can also reinforce immune stimulation and provide positive health benefits. However, despite extensive research in the area of probiotics and the fact that probiotic products have been on the market since the creation of Yakult in 1935, none of the numerous commercial probiotic *Lactobacillus* products have been officially approved for their health-promoting claims in Europe. To date, the European Food Safety Authority has deemed only one general health claim valid; namely, "live cultures in yogurt or fermented milk improve lactose digestion".

Table 4. Representative *Lactobacillus* strains associated with health-enhancing products. The data were compiled from Siezen & Wilson 2010 and Saxelin *et al.*, 2005.

Strain	Brand name	Claimed effect
<i>L. casei</i> Shirota	Yakult®	Alleviation of acute diarrhoea
L. rhamnosus GG	Gefilus®	Immune stimulation, alleviation of atopic eczema, prevention of diarrhoea, aleviation of symptoms associated with irritable bowel syndrome
L. acidophilus NCFM	Howaru®	Improvement of intestinal health, alleviation of symptoms associated with irritable bowel syndrome, gastrointestinal ecology
<i>L. casei</i> DN114-001	Actimel®	Diarrhoea treatment, gut infections, strengthening of the body's natural defences
L. reuteri 55730	Boost®	Alleviation of colic, pathogen inhibition

1.3 Lactobacillus genomes

The first 135 *Lactobacillus* genomes to be published included 38 finished and 97 draft genomes (Table 5 and Appendix Table 2). These genomes have enabled researchers to gain insight into the ecology, evolution, and biological role of lactobacilli and represent in total 46 different species that vary widely in GC content (32-53%), tRNA gene number (25 to 98), coding efficiency (49-91%), and genome size (1.2-3.8 Mb). Below, some of the most valuable discoveries from the first 135 *Lactobacillus* genome projects are outlined in detail.

L. delbrueckii is the largest of the 29 phylogenetic Lactobacillus groups (Salvetti et al., 2012). The first genomes of this phylogenetic group to be resolved were the approximately 2.0-Mb genomes of L. johnsonii NCC 533 (Pridmore et al., 2004) and L. acidophilus NCFM (Altermann et al., 2005). Notably, these two GIT-associated isolates possess a bile salt hydrolase and various types of adhesins that support their persistence in the GIT. Their ability to synthesise cofactors and vitamins is in contrast limited and both isolates can synthesise only some amino acids de novo. These deficiencies are, however, alleviated by their broad repertoires of peptidases, proteases, and transporters, which allow efficient amino acid acquisition from the surrounding medium. As for these GITassociated strains, genomes are available for several dairy-related isolates of the L. delbrueckii group. These include L. delbrueckii ATCC11842, which is known for its worldwide application in yogurt production (van de Guchte et al., 2006), and L. helveticus DPC 4571, a Swiss cheese isolate that is recognised for its ability to reduce bitterness and increase flavour development in cheese (Callanan et al., 2008). Intriguingly, L. helveticus DPC 4571 has few cell-surface-protein-encoding genes and does not encode bile salt hydrolase, despite its striking genome conservation with L. acidophilus NCFM (Callanan et al., 2008). Analysis of the L. delbrueckii ATCC11842 genome has revealed an exceptionally high number of rRNAs, tRNAs, and partial carbohydrate-utilisation pathways, suggesting that this genome has undergone a recent phase of size reduction (van

76	20	1181	1.3	Human	1/1	-	Unclassified
	368	2534	2.7	Apple mash	1/1	_	L. caccinostercus
	17-440	1552-2642	1.9-2.7	Apple juice, dairy, poultry, human, kimchi	5/25	12	L. salivarius
	145-173	1862-1885	1.8-1.9	Sausage	2/4	N	L. sakei
	3-320	1051-2818	1.7-2.8	Beets, plant material, human, kimchi, mouse, rat, silage, porcine	9/15	23	L. reuteri
	27-153	2755-3154	3.2-3.8	Olives, silage, human, kimchi, cabbage	2/5	თ	L. plantarum
	196	1968	2.0	Beer	1/1		L. malefermentans
	127-163	1284-1358	1.4-1.4	Sake, sourdough	2/5	2	L. fructivorans
0-126	3-421	1144-2330	1.2-2.4	Beer, dairy, fermented yak milk, human, kefir grain, porcine, poultry, starter culture	11/27	61	L. delbrueckii
	15-942	2719-3255	2.5-3.1	Dairy, corn steep liquor, human, koumiss, laboratory strain, starter culture	3/3	17	L. casei
	134-574	2392-3325	2.6-3.0	Corn steep liquor, ethanol production plant, human, wine, oral cavity	4/12	ហ	L. buchneri
	235-254	2218-3041	2.3-3.1	Wine, silage	1/10	2	L. brevis
	290-349	2354-2440	2.5-2.5	Sausage	2/11	N	L. alimentarius
RNA genes	Strain specific orthologue groups	CDSs	Length (Mb)	Source	Species with sequenced strains	Sequenced strains	Clade name as in Salvetti <i>et al.</i> , 2012

genome.

Table 5. Characteristics of sequenced Lactobacillus groups and their genomes. See Appendix Tables 2 and 3 for a full description of each

de Guchte *et al.*, 2006). Finally, genomes for *L. crispatus*, *L. jensenii*, and *L. iners* isolates have provided novel perspectives on the genomic basis of urogenital lactobacilli. These genomes have, for example, revealed the lack of a complete bacteriocin synthesis apparatus and most known adhesion factors in *L. iners* AB-1 (Macklaim *et al.*, 2011) and the presence of three bacteriolysin and seven adhesion and colonisation-related protein encoding loci in the *L. crispatus* core-genome (Study V).

The genomes of the sequenced L. casei group members are approximately 2.8 Mb in size (Table 5) and appear to all harbour a repertoire of genes that are involved in sugar uptake, carbohydrate utilisation, and amino acid biosynthesis (Makarova et al., 2006; Cai et al., 2009; Morita et al., 2009). Importantly, pilus gene clusters similar to those that were disclosed in the genome study of L. rhamnosus GG (Study III) are also widespread and have hitherto been identified in the genomes of all probiotic (Douillard et al., 2013a) and various other (Douillard et al., 2013b; Kant et al., 2014) members of this Lactobacillus phylogenetic group. These genome investigations have also provided insights into the genetic complexity of these species and have been helpful in defining the scale and scope of biomedically important effector molecules in the L. casei group that have previously been associated with a cytokine production (Péant et al., 2005), promotion of in vitro intestinal epithelial homeostasis (Yan et al., 2007; Lebeer et al., 2008), and adaptation of strain GG to the host environment (Lebeer et al., 2009). Other genome studies have on the other hand provided valuable insights into the genomic niche-associated evolution of L. *casei* and have determined the scale and scope of genetic variation of this versatile and important species (Cai et al., 2009).

The *L. salivarius* group is the second most speciose group of the genus *Lactobacillus*, and includes representatives of both non-motile (13) and motile (12) *Lactobacillus* species (Salvetti *et al.*, 2012). The genomes associated with this phylogenetic group range from 1.9 to 2.7 Mb and can consist of multiple large replicons. For example, a megaplasmid comprises 11% of the genome of *L. salivarius* UCC118 (Claesson *et al.*, 2006). Although this megaplasmid contains no essential genes, it encodes functions that are beneficial to the host cell, such as the ability to use additional sugars, to hydrolyse bile salt, and to produce supplementary amino acids. The plasmid also includes a locus that encodes and is required for the synthesis of bacteriocin (Claesson *et al.*, 2006), which is active against *Listeria monocytogenes* (Corr *et al.*, 2007). Analysis of the genome of *Lactobacillus ruminis* ATCC 27782 has been highly useful in understanding the nature of flagellum-mediated motility in the genus *Lactobacillus* and has revealed a complete set of flagellum biogenesis genes (Forde *et al.*, 2011). The genome study has also described genes that share similarity with known pilin genes, suggesting that *L. ruminis* ATCC 27782 might contain a sortase-dependent pilus organelle.

Genomes in the *L. reuteri* group are on average 2.1 Mb and contain on average 2,055 CDSs (Table 5). They appear to be rich in genes encoding putative cell-surface-associated proteins (Båth *et al.*, 2005; Saulnier *et al.*, 2011), and some strains possess an EPS gene cluster that is involved in modulating host immune responses (Saulnier *et al.*, 2011). In addition, genomic island coding for the production of a broad-spectrum antimicrobial substance termed reuterin has been identified in selected *L. reuteri* strains. This reuterin island is particularly interesting because the production of reuterin appears to be linked to

the prevention of gastrointestinal infections (Saulnier *et al.*, 2011). Moreover, genomic analyses have revealed loci that are associated with the production of vitamins B_1 (Saulnier *et al.*, 2011) and B_{12} (Morita *et al.*, 2008), although it remains an open question whether *L. reuteri* produces an active (Mohammed *et al.*, 2014) or inactive (Santos *et al.*, 2007) form of vitamin B_{12} .

Other economically or biologically important lactobacilli are found in the *L. sakei* and *L. plantarum* groups (Table 5). These include the psychrotrophic bacterium *L. sakei* 23K (Chaillou *et al.*, 2005). Based on its genome, this organism has metabolic pathways for arginine catabolism and purine nucleoside scavenging. Other notable features include genes implicated in dealing with the harsh conditions associated with food processing and allowing growth on meat during refrigeration and in the presence of curing salts. Genome analysis have also revealed auxotrophy for all amino acids except aspartate and glutamic acid (Chaillou *et al.*, 2005). Another noteworthy strain is *L. plantarum* WCFS1, which is the first organism from the *Lactobacillus* genus to be sequenced (Kleerebezem *et al.*, 2003). The genome of *L. plantarum* WCFS1 is large, close to 3.3 Mb in size, and appears to encode a number of transporters and enzymes for the uptake and utilisation of sugars, thus providing an explanation for the widespread distribution of the *L. plantarum* species in nature (Kleerebezem *et al.*, 2003). The authors also described a number (>200) of extracellular proteins that could enable exchange signals with the environment and adherence to surfaces (Kleerebezem *et al.*, 2003).

1.3.1 Computational genomics of Lactobacillus

The current Lactobacillus genome data were largely obtained using Sanger and Roche 454 sequencing technologies (Appendix Table 3). The first 15 genomes were chiefly determined using Sanger technology (Kleerebezem et al., 2003; Pridmore et al., 2004; Altermann et al., 2005; Chaillou et al., 2005; Makarova et al., 2006; Claesson et al., 2006; van de Guchte *et al.*, 2006; Frese *et al.*, 2011; Macklaim *et al.*, 2011; Morita *et al.*, 2008); after these genomes, sequencing has typically relied on the Roche 454 platform. For example, in 2009, most *Lactobacillus* genome projects were obtained using the Roche 454 platform (Appendix Table 3). Surprisingly, sequence data were generated using the Illumina sequencing platform alone in only three cases (part of the human microbiome project (HMP); Nelson et al., 2010). As expected, Newbler represents the most popular assembler. In addition, Phrap, Jazz, Velvet, and CLC bio genome assemblers have been employed in more than two genome projects (Appendix Table 3). Among the 97 draft assemblies, the median contig number is 75. Given their high degree of sequence similarity and synteny with finished assemblies, draft assemblies are however presumed to offer a near-complete picture of the genome. The unresolved regions of these genomes most likely comprise primarily long, repetitive sequences, such as those encoding ribosomal genes, MGEs, and repetitive structures of some surface-protein genes (Seepersaud et al., 2005; Edelman et al., 2012). The most notable exception is the genome assembly of L. rhamnosus MTCC 5462, which comprises 2,543 contigs and might include a notable number of contigs that end in incomplete gene sequences (Prajapati et al., 2012).

As expected for sequences with GC contents ranging from 32 to 59% (Marine *et al.*, 2011), the genomic GC content has not affected assembly quality, and no obvious correlation was detected between the contig numbers and the GC contents of different *Lactobacillus* genomes.

Regarding genome annotation, no single bioinformatic tool or approach has gained supremacy (Appendix Table 3). Structural annotation has relied on several gene callers, mainly Glimmer, GeneMark, tRNAscan-SE, and RNAmmer, as was the case in the 55 genome projects that were processed using the BCM, JCVI, or PGAAP genome annotation pipelines. Functional classification has primary been performed using BLAST and domain search tools. On occasion, protein functions have been amended manually (Kleerebezem et al., 2003; Pridmore et al., 2004; Altermann et al., 2005; Chaillou et al., 2005; Makarova et al., 2006; Claesson et al., 2006; van de Guchte et al., 2006; Frese et al., 2011; Macklaim et al., 2011; Morita et al., 2008; Cai et al., 2009; Study III; Morita et al., 2009; Study IV; McNulty et al., 2011; Forde et al., 2011). Overall, the level of manual curation is higher for finished genomes and genomes that were processed before 2010, as reflected by their higher rates of start site consistency and exact DEs. In particular, more attention was given to the annotation of the first 15 Lactobacillus genomes. The initial annotations were in these projects generated using a variety of methods and curated manually. Many of the more recent genomes have on the other hand chiefly relied on the non-human assisted use of protein function prediction services. However, annotation processes are generally poorly documented, which complicated the comparison of annotation processes. For 14 genomes, no information was offered; for another 67 genomes, only the general protocol of the sequencing centre that had performed the sequencing was listed. Significantly, in various instances, neither genome annotations nor sequences have been updated since the data release by means other than the automatic annotation systems used by the databases. Annotations between different genomes can thus be inconsistent, even for strains of the same species. The most notable exception is the genome of L. plantarum WCFS1, which was re-sequenced and re-annotated by Siezen et al. recently (Siezen et al., 2012).

1.3.2 Comparative genomics of Lactobacillus

Comparative analyses of *Lactobacillus* have expanded our understanding of the molecular evolution, diversity, and function of lactobacilli. Importantly, these studies have revealed significant functional and genomic variance between *Lactobacillus* genomes (Boekhorst *et al.*, 2004; Pridmore *et al.*, 2004; Canchaya *et al.*, 2006; Berger *et al.*, 2007; Ventura *et al.*, 2008; Canchaya *et al.*, 2006; Claesson *et al.*, 2008; Morita *et al.*, 2008; Azcarate-Peril *et al.*, 2008; O'Sullivan *et al.*, 2009; Kant *et al.*, 2011a; Lukjancenko *et al.*, 2012). For example, only 28 large regions of conserved gene order, ranging in size from 7 to 75 genes, were found in a comparative analysis of *L. plantarum* WCFS1 and *L. johnsonii* NCC 533 genomes (Boekhorst *et al.*, 2004). A lack of gene order synteny has also been shown for other distant lactobacilli, whereas the genomes of closely related lactobacilli tend to exhibit a high degree of gene order conservation across their entire genomes

(Canchaya et al., 2006; Berger et al., 2007; Ventura et al., 2008). However, even the genomes of closely related strains of the same lactobacilli species can differ by one or more genomic islands (Pridmore et al., 2004; Study III; Morita et al., 2008; Azcarate-Peril et al., 2008). These regions of diversity typically lack similarity with other lactobacilli regions and contain genes that might be relevant for environmental adaptation. For example, genomic islands contain genes coding for EPS biosynthesis in L. gasseri ATCC 33323 (Azcarate-Peril et al., 2008), fimbrial components in L. johnsonii NCC 533 (Pridmore et al., 2004), pilus fibres in L. rhamnosus GG (Study III), and B₁₂ and reuterin biosynthesis in L. reuteri JCM 1112^T (Morita et al., 2008). In addition, horizontal gene transfer has been linked to the ability of L. plantarum WCFS1 to adapt to a variety of niches, to the adaptation of L. delbrueckii to dairy environments (Liu et al., 2009; van de Guchte *et al.*, 2006), and to the acquisition of amino acid metabolism, lipid biosynthesis, and restriction endonuclease genes during the evolution of L. helveticus DPC 4571 (Callanan et al., 2008). However, predicting laterally acquired genome regions is not always straightforward. For example, the sequencing and analysis of the genomes of L. ruminis ATCC 27782 and ATCC 25644 identified a bacteriocin cluster in ATCC 27782, although it remained unclear whether ATCC 25644 also contained a complete bacteriocin locus; some genes associated with bacteriocin production were missing and the others did not assemble into a single contig in the latter strain (Forde et al., 2011).

Comparative genomics studies have revealed interesting biological similarities and differences among lactobacilli. The lactobacilli genomes, for example, appear to contain a constant fraction of flavour-related genes, independently of their isolation. The approximately 2.0-Mb genomes of L. delbrueckii ATCC 11842 and L. acidophilus NCFM, for example, code for 15 and 14 flavour-related enzymes (Liu *et al.*, 2008), even though the first is a dairy isolate and the second is a human isolate. Analysis of Lactobacillus secretomes has revealed that on average, 8% of a lactobacilli proteome represents secreted or surface-associated proteins (Zhou et al., 2010; Kant et al., 2010). The largest predicted secretome (approximately 9% of the predicted proteome) is that of L. acidophilus NCFM, whereas the smallest predicted secretome (approximately 5% of the predicted proteome) was that of L. reuteri DSM 20016 (Kant et al., 2010). Comparative genomics has been used to measure the diversity of CRISPR-cas systems, revealing that approximately two thirds of the analysed *Lactobacillus* strains have a CRISPR locus (Horvath et al., 2009). Mucus-binding domain screens have been used to measure the prevalence of mucusbinding proteins in lactobacilli; the most abundant mucus-binding domain-containing proteins were found in GIT-associated lactobacilli, supporting the idea that mucus-binding proteins are involved in adherence to the intestinal mucus that covers intestinal epithelial cells (Boekhorst et al., 2006).

1.3.3 Comparative core and pan-genomics of Lactobacillus

The microbial pan-genome comprises a core and an accessory gene pool (Tettelin *et al.*, 2005). Core genes are conserved across all isolates of a given group of organisms, whereas accessory genes are present in some but not all strains (Tettelin *et al.*, 2005). In general,

core genes are responsible for basic cellular processes and the main phenotypic traits of the group, whereas accessory genes contribute to diversity within the group and enable adaptation to specific environments (Medini et al., 2005). When applied to Lactobacillus genomes, core genome investigations have defined differing sets of 593 (Canchaya et al., 2006), 141 (Claesson et al., 2008), 383 (Kant et al., 2011a), and 363 (Lukjancenko et al., 2012) core genes that are shared by all lactobacilli. These apparently contradictory results are tentatively explained by methodological differences and by the number of genomes analysed in the individual studies; 5 by Canchaya et al., 2006, 12 by Claesson et al., 2008, 20 by Kant et al., 2011a, and 20 by Lukjancenko et al., 2012. Surprisingly, the impact of genome number and the consequence of sequentially adding more genomes and the core genome size of an infinite number of Lactobacillus genomes were not estimated in any of the studies, as is typically performed in core and pan-genome studies (Medini et al., 2005). Nevertheless, these and other comparative studies have provided key insights into the genome evolution of Lactobacillus and revealed that the gene complements are the results of extensive gene losses and gains during evolution (Claesson et al., 2008; Kant et al., 2011a) and are devoid of habitat-specific genes (Claesson et al., 2008; O'Sullivan et al., 2009), and that the core genome contains only a few genus-specific genes (Claesson et al., 2008; Canchaya et al. 2006).

The core genomes of specific Lactobacillus species have also sparked interest. The establishment of the level of intraspecies diversity in seven Lactobacillus species using comparative genomic hybridisation and the mapping of short-read sequences to reference genomes revealed that individual strains lack 3-24% of the genes of the given reference genome and that the size of the core genome is correlated with the total size of a single reference genome (Siezen et al., 2010; van Hemert et al., 2010; Meijerink et al., 2010; Berger et al., 2007; Cai et al., 2009; Raftis et al., 2011; Nyquist et al., 2011; Frese et al., 2011; Douillard et al., 2013b). Additionally, whole-genome assembly comparisons have been used for the investigation of genetic diversity in L. paracasei (Smokvina et al., 2013), L. rhamnosus (Kant et al., 2014), and L. casei (Broadbent et al., 2012). This approach is more powerful than read-mapping and microarray-based approaches and can call genes that are present in genomes other than the reference genome. Using this approach, gene loss and gain was found to be the dominant force in genome evolution within both Lactobacillus species, although some variation took the form of differences in universally conserved genes. For example, L. casei strains exhibited >99% identity to the 16S rRNA sequence of L. casei ATCC 334. Nevertheless, on average, 119 strain-specific gene families are present in each genome, and only 61% of any individual genome is shared by all 17 strains. Mathematical modelling of the data indicates extensive genome diversity and that the L. casei pan-genome contains 9,072 gene families, of which 1,600 are common to all individuals (Broadbent et al., 2012).

2 Aims of the study

The aim of this study was to develop algorithms for the automated function prediction of bacterial protein sequences and to advance our understanding of *Lactobacillus* physiology by annotating the genomes of *L. rhamnosus* GG and LC705 and *L. crispatus* ST1. Genes in the genomes that exhibited functions that are involved in interactions with the host and that responsible for strain-specific characteristics were of particular interest. To determine the scale and scope of genetic variation in *L. crispatus* and to infer physiological traits that are common for all *L. crispatus*, pan-genomic strategies were also considered. Overall, the goal was to develop efficient and accurate ways to annotate bacterial genomes.

3 Materials and methods

The bioinformatics methods developed are described in detail with appropriate references in original publications I and II. Detailed descriptions of materials and methods used in the genome investigations are in the original papers III-V.

3.1 Evaluation of bioinformatics methods

A comprehensive assessment of the newly developed bioinformatics methods is available in the original publications I and II. Briefly, the performance of the LOCP software tool (Study I) was evaluated based on 20 completely or partially sequenced genomes with known pilus operons. In this process, the genes in these genomes were assigned scores of their highest ranked LOCP predictions or were assigned a score of zero indicating that the gene was not reported by LOCP. The ability of LOCP to distinguish genuine pilus-related from other genes was then assessed using receiver operating characteristic analysis. In Study II, the data used in the development process of BLANNOTATOR were obtained from UniProt (Bairoch et al., 2008). Protein function labels were restored to the state preceding the functional characterisation of the test sequences using scripts that were developed in-house. The details of the restoring process as well as the use of competing protein function classification methods and statistical tests are in the original paper II. In addition, a retrospective assessment of the classification accuracy of nine gene-calling systems was performed, and this assessment is described in this thesis. Specifically, CDSs in L. rhamnosus GG, L. rhamnosus LC705, and L. crispatus ST1 were called using nine gene-calling tools, and the sensitivity, specificity, and F-score (*i.e.*, the harmonic mean of the sensitivity and specificity) were computed at the single-base level. The gold standard sets that were used in the comparison were the manually annotated gene models from the genomes as well as regions with and without transcriptional activity in L. rhamnosus GG. Transcriptome data were compiled from three previous L. rhamnosus GG gene expression studies (Laakso et al., 2011; Koskenniemi et al., 2011; Koponen et al., 2012) and comprised 626,450 bases with evidence of transcription (defined as bases that are covered by probes with an intensity value ≥ 5 standard deviations above the mean intensity of negative control probes in ≥ 18 of the 360 RNA samples described in these three studies).

3.2 Strains and growth conditions

The *Lactobacillus* strains that were sequenced in Studies III-IV included *L. rhamnosus* GG (ATCC 53103), *L. rhamnosus* LC705 (DSM 7061), and *L. crispatus* ST1. To prepare genomic DNA, each strain was grown in the de Man, Rogosa, and Sharpe (de Man *et al.*, 1960) broth at 37°C. DNA was extracted as previously described (Pitcher *et al.*, 1989).

3.3 Sequencing and assembly

Whole-genome sequencing was done at the DNA sequencing and Genomics Laboratory at the Institute of Biotechnology, University of Helsinki. Briefly, DNA from *L. rhamnosus* GG and LC705 was processed according to publication III. Plasmid and fosmid libraries were sequenced using an ABI 3730 DNA sequencing instrument and Big Dye chemistry (Applied Biosystems, Foster City, CA, USA), whereas genomic fragment libraries were sequenced using Roche GS 20 pyrosequencing (454 Life Sciences/Roche Applied Biosystems, Branford, CT, USA). In Study IV, genomic DNA from *L. crispatus* ST1 was used for 454 library construction and sequenced using a Roche 454 instrument with GS FLX chemistry (454 Life Sciences/Roche Applied Biosystems, Branford, CT, USA). All DNA reads were processed and assembled using the Staden Package (Staden *et al.*, 1999) and/or Newbler (454 Life Sciences/Roche Applied Biosystems, Branford, CT, USA). For gap closure, the PCR-amplified fragments obtained using genomic DNA were sequenced using an ABI 3730 instrument and Big Dye chemistry (Applied Biosystems, Foster City, CA, USA).

3.4 Accession numbers for the submitted data

The genome sequences of *L. rhamnosus* GG and LC705 and the plasmid pLC1 have been deposited in the EMBL nucleotide sequence database under accession numbers FM179322, FM179323, and FM179324, respectively. The genome sequence of *L. crispatus* ST1 has been deposited in the EMBL nucleotide sequence database under accession number FN692037.

3.5 Publicly available genome sequences

The genome data used in Studies III-V were downloaded from the GenBank database (Benson *et al.*, 2013) and the PATRICK database (Gillespie *et al.*, 2011), as indicated in the original publications. The genome data referenced in this dissertation were downloaded in April 2012 from GenBank (Benson *et al.*, 2013) and the PATRICK database (Gillespie *et al.*, 2011). The NCBI database was preferred over the PATRICK database when a genome was available at both databases, provided the NCBI entry was properly annotated. Where possible, scaffold assemblies were preferred.

3.6 Structural and functional annotation

The genomes in Studies III and IV were scanned for CDSs, tRNAs, rRNAs, CRISPRs, genomic islands, prophage-like clusters, and rho-independent transcription terminators

using an array of computational methods (Table 6). Predicted protein sequences were then searched against a variety of databases and further processed using a set of bioinformatics tools with the aim of assigning function (Table 6). Automated computer annotations were verified, and discrepancies were resolved manually. Detailed descriptions of databases and software tools used are in the original papers III-IV. In Study V, genome sequences were mined for CRISPRs, genomic islands, plasmids, and prophage-like regions using a variety of methods (Table 6). A functional annotation update was also performed to ensure that protein function predictions were of identical quality for all of the investigated *L. crispatus* genomes. An overview of the computational methods used in Study V is presented in Table 6. The use of these tools is described in detail in the corresponding publication. Where feasible, the methods developed in Studies I and II were used to obtain details about pilus-like gene clusters and protein functions, respectively, in Studies III-V.

3.7 Metabolic pathway reconstruction

Using the KAAS tool (Moriya *et al.*, 2007), CDSs within the genomes of *L. rhamnosus* GG, *L. rhamnosus* LC705, and *L. crispatus* ST1 were annotated for EC numbers describing enzymatic activity. Metabolic pathway reconstructions were then realised by associating enzymatic activities with combined KEGG (Kanehisa *et al.*, 2004) and MetaCyc (Krieger *et al.*, 2004) reference reaction pathways and by manually investigating reaction maps. To resolve the precise metabolic activity of genes with partial EC codes, the enzymatic activity supported by the API 50 CH (BioMérieux, Marcy l'Etoile, France) carbohydrate-fermenting patterns was chosen where possible. In Study V, EC codes were determined using KAAS software (Moriya *et al.*, 2007) and the FMM server was used to assemble these EC codes to metabolic pathways (Chou *et al.*, 2009).

3.8 Comparative analyses

The methods used in Studies III-V for orthologue analysis, phylogenetic tree construction, and whole-genome alignment are summarised in Table 6. The *L. rhamnosus* whole-genome nucleotide alignments that are presented in this thesis were generated using BLASTN (Altschul *et al.*, 1997) and ACT (Carver *et al.*, 2005). The *L. rhamnosus* draft genomes were ordered and oriented with respect to the genome sequence of *L. rhamnosus* GG using progressive Mauve (Rissman *et al.*, 2009). To produce the *L. crispatus* whole-genome nucleotide alignments that are presented in this thesis, matching genome blocks between the genome of *L. crispatus* ST1 and the genomes of *L. crispatus* CTV-05, *L. helveticus* DPC 4571, *L. acidophilus* NCFM, *L. johnsonii* NCC 533, *L. gasseri* ATCC 33323, and *L. delbrueckii* ATCC 11842 were identified using PROmer and visualised using a MUMmer plot (Kurtz *et al.*, 2004). The draft genome of *L. crispatus* ST1 using progressive Mauve (Rissman *et al.*, 2009).

3.9 Core and pan-genome analyses

Orthologue and paralogue groups among the L. crispatus genomes were in Study V identified using BLASTP (Altschul et al., 1997) and OrthoMCL (Li et al., 2003). To estimate the development of the size of the core and pan-genome as a function of the number of sequenced strains, orthologue and paralogue groups were determined iteratively for increasing numbers of sequenced genomes. At each sample size, the analysis was repeated 50 times with different random sets of L. crispatus genomes. The core genome trend was extrapolated by fitting an exponential decay (Tettelin et al., 2005) to the medians of the core orthologue groups using a weighted least-squares regression. The number of pan-groups in an infinite number of L. crispatus genomes was predicted by fitting a power-law (Tettelin et al., 2005) to the pan-group medians using a weighted leastsquares regression. The regression analyses were performed using the *nls* function as implemented in the statistical software R (Ihaka & Gentleman, 1996). The methods used for identifying the orthologue and paralogue groups among the Lactobacillaceae genomes and for estimating the Lactobacillaceae core and pan-genome sizes were the same as those described in Study V with the exception of using a double exponential decay (Bottacini et al., 2010) for the core genome data. The results of Lactobacillaceae analyses are presented in this thesis.

	L. rhamnosus GG	L. rhamnosus LC705	L. crispatus ST1	L. crispatus
Structural annotation				
CDS	Glimmer, BLAST, ERGO	Glimmer, BLAST	Glimmer, BLAST, RAST	
rRNA	RNAmmer, ERGO	RNAmmer	RNAmmer, RAST	
tRNA	tRNAscan-SE, ERGO	tRNAscan-SE	tRNAscan-SE, RAST	
CRISPR	PILER-CR	PILER-CR	PILER-CR	PILER-CR
Intrinsic terminators	TransTermHP	TransTermHP	TransTermHP	
Plasmid related contigs				cBar
Genomic islands	PAI-IDA	PAI-IDA	PAI-IDA	IslandViewer
Functional classification				
General function prediction	BLAST, ERGO, InterProScan, COG	BLAST, PFAM, InterProScan, COG	RAST, COG InterProScan, BLAST, BLANNOTATOR	RAST, BLAST, COG, 45 BLANNOTATOR
Gene Ontology terms	InterProScan to GO	InterProScan to GO	InterProScan to GO	
Transporters	TransAAP, TCDB	TransAAP, TCDB	TCDB	TCDB
Enzymes	Merops, CaZY, KAAS	Merops, CaZY, KAAS	Merops, CaZY, KAAS	Merops, KAAS
cas genes	TIGRfam	TIGRfam	TIGRfam	TIGRfam
Bacteriocins	InterProScan	InterProScan	InterProScan, BLAST	BAGEL
Prophages	Prophinder	Prophinder	Prophinder	Prophinder
Adhesion factors	PFAM	PFAM	PFAM (revised)	PFAM (revised)
Secretome	SignalP, LipoP, TMHMM	SignalP, LipoP, TMHMM	LocateP	
Other analysis				
Ortholog assignment	InParanoid	InParanoid	OrthoMCL, Cliquer	OrthoMCL
Phylogenetic analysis	BLASTP, PHYLIP	BLASTP, PHYLIP	Muscle, Gblock, PhyML	Mauve, PhyML
Whole-genome alignment	BLASTN, ACT, Gepard	BLASTN, ACT, Gepard	BLASTN, ACT, Gepard, Mauve	BLASTN, ACT, Mauve
Metabolic pathways	KEGG, MetaCyc	KEGG, MetaCyc	KEGG, MetaCyc	FMM

Table 6. Methods used for annotation of the L. rhamnosus GG, L. rhamnosus LC705, and L. crispatus genomes.

4 Results and discussion

The main objective of this study was to develop algorithms for protein classification that go beyond the present practice in several aspects and to advance our understanding of *Lactobacillus* physiology with the aid of genome sequencing and comparative genomics approaches. In particular, novel insights into the physiology, ecology, and biochemistry of lactobacilli were provided by comparing the genome of the widely studied (Bernardeau et al., 2006) and commercially significant (Saxelin et al., 2005; Giraffa et al. 2010) probiotic bacterium L. rhamnosus GG with the genome of the industrial dairy strain L. rhamnosus LC705 (Suomalainen & Mäyrä-Mäkinen, 1999). Furthermore, the comparative analysis of L. crispatus genomes unveiled cross-species conserved adhesion components that could protect the vagina from pathogen attack. The comparison also highlighted an array of other interesting differences and similarities between the vaginal L. crispatus isolates and L. crispatus ST1, which is known for its strong adherence to the chicken alimentary canal and to human vaginal and buccal cells (Edelman et al., 2012). In the following chapters, the main findings of these studies are presented in detail. First, the bioinformatics approaches used are described; then, the outcomes of the Lactobacillus genome studies are summarised.

4.1 Novel tools for predicting the function of bacterial proteins

Recently developed protein function prediction methods provide a comprehensive and efficient means for inferring protein function (see for example, Abascal & Valencia, 2003; Kunin & Ouzounis, 2005; Martin *et al.*, 2004; Vinayagam *et al.*, 2006; Hawkins *et al.*, 2006; Wass & Sternberg, 2008; Engelhardt *et al.*, 2005; Zdobnov & Apweiler, 2001) but can sometimes fail to characterise pilins (Scott & Zähner, 2006) and produce unreliable functional calls (Friedberg, 2006; Rost, 2002; Rost *et al.*, 2003). To overcome these difficulties, new computational approaches were developed for pilus operon (Study I) and protein function (Study II) prediction. As described in the following chapters, these newly developed bioinformatics tools provided many interesting insights into the physiology of *L. rhamnosus* and *L. crispatus*.

Pili are long filamentous protein assemblies that are located on the surface of bacteria and are often involved in the adhesion of bacteria to host cells (Madigan *et al.*, 2010). In gram-positive bacteria, these structures typically comprise one major pilin protein and two auxiliary pilin proteins that are cross-linked by a sortase enzyme (Telford *et al.*, 2006; Scott & Zähner, 2006). It appears that the pilin genes are usually located in an operon with a sortase gene (Scott & Zähner, 2006) and that their protein products display various features that are characteristic of gram-positive pilins, such as a positively charged tail and a membrane-spanning domain at the C terminus, an E box, a sortase recognition site (a LPXTG motif), a pilin motif, and a Sec-dependent secretion signal peptide (Telford *et al.*, 2006). In addition, sequence comparisons with gram-positive pilins have revealed conserved sequence motifs that stabilise the structure (Kang *et al.*, 2009). Although these

motifs have enabled the search for putative pilus operons (Ton-That & Schneewind, 2003), a novel bioinformatic algorithm was proposed in Study I for the systematic screening of pilus operons in gram-positive genomes. This tool, LOCP, which is written in Perl, scans sequences with five pilin-, five sortase recognition site-, and three sortase enzyme profile HMMs. Each sequence is labelled either as a hit (if at least one HMM model is matched) or as a miss, after which genome regions that are enriched with hits are identified based on a hypergeometric distribution. The use of a discrete distribution to locate gene runs that are statistically enriched in sequence features borrows from the concept of a previously described prophage-finding program (Lima-Mendez *et al.*, 2008) and was shown to distinguish genuine pilus clusters precisely from other genome regions. Specifically, LOCP identified all 28 genuine pilus clusters and made no false predictions in the given 20 evaluation genomes (the area under the curve was approximately 0.99). To date, no other bioinformatics tools other than LOCP have been developed for locating pilus operons in bacterial genomes.

To facilitate bacterial genome annotation, a new protein function classification system was created in Study II. Leveraging the advantages of DEs and GOs, the software BLANNOTATOR generates predictions based on database hits that are associated with consistent protein function information. This strategy is similar to those employed in the CLAN (Kunin & Ouzounis, 2005) and ConFunc (Wass & Sternberg, 2008) protein classification systems and is known to be less error-prone to annotation anomalies than methods that rely on a single or all database hits, such as the PFP (Hawkins et al., 2006) and the ARGOT2 (Falda et al., 2012) tools. Importantly, BLANNOTATOR was exceedingly helpful in systematising DEs and generated precise function predictions in evaluation tests. When applied to the predicted proteins of L. crispatus ST1, the algorithm assigned a biologically acceptable DE for 85% of the query sequences. In comparison, RAST- (Aziz et al., 2008) or BLAST-based approaches provided a valid function prediction for approximately only 58 and 69% of the proteins in the test set. This method was particularly useful for predicting the function of proteins for which the top database hits were uninformative, as discussed in detail in Study II. The accuracy of BLANNOTATOR was further benchmarked by simulating the annotation process for more than 3,000 high-quality annotated bacterial protein entries and by assessing the ability of BLANNOTATOR to reproduce current annotations based on annotation information that predated the functional characterisation of the test entries. For this dataset, the method produced more precise predictions than any of the five other function classification approaches (i.e., most significant BLAST match, the top informative BLAST match, the most common annotation among BLAST hits, the annotation associated with the highest cumulative BLAST bit score, and a word-based scoring scheme) that were tested. However, performance differences between the protein function prediction approaches were marginal, and even the worst performing approaches provided reasonable accuracy, indicating that major improvements in the field of homology-based function transfer are less likely to occur in future.

4.2 L. rhamnosus and L. crispatus genome sequencing

The objectives of Studies III (initiated in 2004) and IV (initiated in 2008) were to produce finished genomes for the previously undescribed isolates of L. rhamnosus and L. crispatus using the WGS sequencing and assembly strategy (Fleischmann et al., 1995). To achieve this, the genomes of L. rhamnosus GG, L. rhamnosus LC705, and L. crispatus ST1 were sequenced using a combination Roche pyrosequencing and Sanger sequencing. These platforms provided approximately 17-19× read sequence coverage for each genome (Table 7) and were preferred in the process over Illumina sequencers because of their long read length (Shendure & Ji, 2008). A collection of PCR-assisted techniques was then applied to improve genome assemblies, providing to resolve all the gaps in the draft genomes of both L. rhamnosus GG and LC705 and all but one gap in the draft genome of L. crispatus ST1. Despite a single gap region remaining in the L. crispatus ST1 assembly, each assembly provided valuable information about the general genome features of the organism (Table 7). They also represented an important step forward in the biology of that species because, before their release, only two L. rhamnosus and five L. crispatus genome sequences had been deposited in public sequence databases (mainly generated by the HMP; Nelson et al., 2010).

L. rhamnosus GG is one of the most extensively studied lactobacilli strains (Saxelin et al., 2005; Bernardeau et al., 2006) and has shown promising results for the treatment or prevention of respiratory tract infections (Hatakka et al., 2001; Hojsak et al., 2010), certain types of diarrhoea (Isolauri et al., 1991; Guandalini et al., 2000; Szajewska & Mrukowicz, 2001), and atopic diseases (Kalliomäki et al., 2001; Kalliomäki et al., 2003; Kalliomäki et al., 2007). The identity of the specific effector molecules behind these beneficial effects was however chiefly lacking prior to the Study III, illustrating the value of knowing its genome sequence. Indeed, the determination and annotation of the genome of L. rhamnosus GG uncovered several genes of potential biomedical importance and expanded our knowledge of the L. rhamnosus bacterial components far beyond the few earlier in vitro verified instances (Vélez et al., 2007; Yan et al., 2007; Iliev et al., 2008; Lebeer et al., 2009). At the summary statistics level (Table 7), strain GG was identified to be comparable to other L. casei and L. paracasei and L. rhamnosus strains (reflecting the close phylogenetic relationships among these strains). It was observed to have one of the largest Lactobacillus genomes and it was predicted to contain only a slightly fewer CDSs and tRNA genes than the largest known lactobacilli genome from L. plantarum WCFS1 (Kleerebezem et al., 2003). A large majority (78%) of CDSs were predicted to start with an ATG and approximately 20% of all CDSs were preceded by a putative RBS (Figure 7), which is defined here as a DNA sequence located at a maximum of -20 bases from the start codon and showing strong similarity to the genome-specific RBS position weight matrix motif model. Genome mining of other genomic features revealed three prophagelike regions and a CRISPR locus that was notably similar to that of L. salivarius UCC118 and L. casei BL23 (Horvath et al., 2009).

	L. rhamnosus GG	L. rhamnosus LC705	L. rhamnosus	L. crispatus ST1	L. crispatus
ACC	FM179322	FM179323	NA	FN692037	NA
Rank (clade / all lactobacilli)	3 / 51	3 / 51	NA	6 / 63	NA
Read coverage	17	19	NA	18	NA
Genome size, Mbp	3010111	3033106	2892564	2043161	2234576
Scaffolds	_		419	_	114
Plasmids	0	-	NA	2	NA
GC content %	47	47	47	37	37
Coding efficiency %	85.00	85.00	80.14	88.00	83.57
CDS	2944	2992	2946	2024	2194
ATG/TTG/GTG %	77/11/12	78/11/11	74/10/11	86/9/5	83/10/7
Function Predicted %	73	72	69	77	67
Enzyme %	24	25	25	24	22
Transporter %	14	15	14	13	13
Transmembrane %	27	28	29	26	N
Average CDS size bp	870	865	800	892	852
rRNA operons	J	ഗ	2-5	4	1-4
tRNA genes	57	61	51	64	57
CRISPR loci		0		2	-

Table 7. Comparison of the genomic features of *L. rhamnosus* GG, *L. rhamnosus* LC705, and *L. crispatus* ST1. Values presented for *L. rhamnosus* and *L. crispatus* are averages for the remaining seven *L. rhamnosus* and seven *L. crispatus* genomes, which are listed in Appendix Table 2.

The dairy-associated strain L. rhamnosus LC705 is widely used in the manufacture of cheese products (Saxelin et al., 2011). LC705 also is one of the main components of a bacterial multispecies product that appears to alleviate irritable bowel syndrome symptoms (Kajander *et al.*, 2005; Kajander *et al.*, 2008) and appears to be immunologically active by inducing the expression of a diverse array of immune response genes in human macrophages (Miettinen et al., 2012) and mast cells (Oksaharju et al., 2011). Intriguingly, the dairy strain LC705 adheres poorly to human mucus (Tuomola et al., 2000) and Caco-2 cells (Jacobsen et al., 1999) and is markedly less effective in colonising humans than strain GG (Saxelin *et al.*, 2010), which originates from the stool specimen of a healthy human (Silva et al., 1987). In Study III, the genome of L. rhamnosus LC705 was revealed to be slightly larger than that of strain GG. This strain was observed to contain a 2.97-Mb circular chromosome and a 64.5-kb circular plasmid that together included 2,992 CDSs, 61 tRNA genes, and 5 rRNA operons (Table 7). Approximately 77% of the CDSs in L. rhamnosus LC705 were found to begin with ATG and approximately 20% of them were preceded by an RBS (Figure 7), highlighting the similarity between the genomic compositions of L. rhamnosus LC705 and L. rhamnosus GG. Notably, LC705 was predicted to contain three prophage-like regions, indicating that transduction might have been an important mechanism for genome evolution in this species, as has been proposed for L. casei (Broadbent et al., 2012).

The chicken isolate L. crispatus ST1 has been shown to colonise various areas of the chicken alimentary canal (Edelman et al., 2002) and has in the literature been documented to strongly adhere to human vaginal epithelial cells, apparently through the highmolecular-mass *Lactobacillus* epithelium adhesin known as LEA (Edelman et al., 2012). Other noteworthy traits associated with L. crispatus ST1 include its inhibition of the adhesion of avian pathogenic E. coli (Edelman et al., 2003) and its ability to secrete proteins that enhance the cleavage of plasminogen into biologically active fragments (Hurmalainen et al., 2007). In an effort to advance our understanding about the physiology of L. crispatus ST1 and to catalogue its adhesion factor potential, the genome of L. crispatus ST1 was determined and analysed in Study IV. The final genome assembly is delimited by a small gap of approximately 590 bp in the lea gene and was estimated to be approximately 2.04 Mb in size, representing the smallest L. crispatus genome reported to date. The genome was predicted to be devoid of plasmids and was found to have a low GC content (37%), which is similar to the GC contents of its closest relatives, L. acidophilus (Altermann et al., 2005; 35%) and L. helveticus (Callanan et al., 2008; 38%) but is different from those of L. rhamnosus GG and LC705 (47%). In terms of functional features, strain ST1 resembled the previously described L. crispatus strains and was predicted to contain 2,024 CDSs (Table 7), several of which might contribute to the maintenance of vaginal health (Study IV). An in-depth analysis of the CDS models revealed that ATG is the most common start codon in strain ST1 and that the use of alternative start codons is consistent with observations made in other L. crispatus, whereby TTG and GTG were associated with 10 and 9% of CDSs, respectively (Table 7). Only 6% of CDSs exhibited an RBS motif, similar in number to the observations for L. crispatus 125-2-CHN, JV-V01, and 214-1 but less than half of the number of observations for the remaining four L. crispatus strains. In Study V, a comparative genomics analysis of *L. crispatus* ST1 and nine vaginal *L. crispatus* isolates was carried to investigate the scale and scope of the pan- and core genomic potential of and genomic diversity in *L. crispatus*.

Collectively, the results indicate that *L. rhamnosus* GG and LC705 contain a genome of approximately 3.0 Mb and that *L. crispatus* ST1 has a genome that is a bit over 2.0 Mb. Although these were not the first *L. rhamnosus* or *L. crispatus* strains to be determined at the genome level, they were the first for these two species to be assigned into a single scaffold, thereby extending the collection of high-quality genomes that are available for the *L. casei* clade beyond *L. casei* (Makarova *et al.*, 2006; Cai *et al.*, 2009) and the collection of high-quality genomes that are available for the *L. delbrueckii* clade beyond *L. johnsonii* (Pridmore *et al.*, 2004; Wegmann *et al.*, 2009), *L. acidophilus* (Altermann *et al.*, 2005), *L. delbrueckii* (van de Guchte *et al.*, 2006), *L. delbrueckii* (Makarova *et al.*, 2006), *L. helveticus* (Callanan *et al.*, 2008), and *L. gasseri* (Azcarate-Peril *et al.*, 2008); thus, this research opened new avenues for the comparative genomics of *Lactobacillus*.

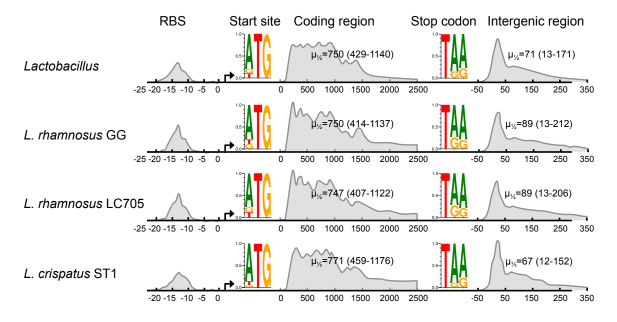


Figure 7. Comparison of consensus CDS models in the *Lactobacillus*, *L. rhamnosus* GG, *L. rhamnosus* LC705, and *L. crispatus* ST1 genomes. From left to right: the spacing distribution between the 5' A residue of the RBS and the translational start point, the sequence logo of the translation start point, the length distribution of CDSs, the sequence logo of the translational stop point, and the length distribution of intergenic regions. The median (and interquartile ranges) of the CDS and intergenic region lengths is indicated.

4.3 Gene calling in *L. rhamnosus and L. crispatus* sequences

Identifying CDSs in genomes is one of the first and most crucial steps in any genome sequence analysis (Angelova *et al.*, 2010) and should receive a great deal of care because annotation errors made at this step of the analysis can have detrimental consequences on

our understanding of the functional capacity of the organism. To ensure that as few annotation errors as possible had occurred at the initial automated gene calling step in Studies III and IV, an extensive set of gene callers was applied to the L. rhamnosus GG and LC705 and L. crispatus ST1 genomes to evaluate their performance at locating genes within their genomes. Of the nine tested gene-calling systems, Glimmer (Delcher et al., 2007) was among the top performers in terms of sensitivity and specificity (Table 8). However, the performance of Glimmer was comparable to those of GeneMark (Besemer et al., 2001) and Prodigal (Hyatt et al., 2010), and only marginal differences existed between these three methods, as has been observed in previous gene-calling performance tests (Hyatt et al., 2010; Angelova et al., 2010). The accuracy of the remaining methods was slightly lower; however, their gene models also matched fairly well with the manually reviewed CDS models and regions in the GG genome with transcriptional activity. An exception to the above was ERGO's gene-calling software (Overbeek et al., 2003), which produced gene calls that did not match those obtained using the other methods and those observed in other lactobacilli. Specifically, the ERGO system preferred CDSs that started from codons other than ATG and produced CDS models that were aligned only partially with those found in sequence databases, suggesting that this service might not provide optimal results for lactobacilli. This result is consistent with the results of an earlier genome investigation, wherein the re-annotation of the genome of Caulobacter crescentus strain NA1000 resulted in a reduction in the use of rare codons and led to the improved annotation of 7% of the original 3,879 ERGO gene models (Ely & Scott, 2014). Another interesting finding was that low sensitivity values were evident in the gene expression test (Table 8). This is partially explained by the presence of polycistronic transcripts that can include intergenic regions (and are thus expressed) but also relates to the finding that various probes targeting an antisense region produced an expression signal that was higher than the chosen threshold.

Table 8. Comparison of CDS calls of nine gene callers. CDS calls were compared against manually refined annotation data and regions with and without transcriptional activity in *L. rhamnosus* GG. The sensitivity (Sn) is the fraction of gold standard bases that was captured by each prediction, and specificity (Sp) is the fraction of non-gold standard bases that was captured by each prediction.

	GG (DSs	LC705	CDSs	ST1 (CDSs	GG mic	croarray
Method	Sn	Sp	Sn	Sp	Sn	Sp	Sn	Sp
Glimmer	98.2 %	99.4 %	98.5 %	99.3 %	99.1 %	99.8 %	61.7 %	95.5 %
Prodigal	98.3 %	99.2 %	98.4 %	99.3 %	99.0 %	99.6 %	61.8 %	95.5 %
GeneMark	98.1 %	99.2 %	98.3 %	99.2 %	99.3 %	99.3 %	61.8 %	95.5 %
EasyGene	96.8 %	99.7 %	97.0 %	99.7 %	98.6 %	99.7 %	58.3 %	96.0 %
YACOP	96.2 %	99.6 %	96.3 %	99.5 %	97.4 %	99.6 %	58.4 %	96.2 %
RAST	98.3 %	98.4 %	96.7 %	97.5 %	98.8 %	98.6 %	64.1 %	95.3 %
ERGO	59.0 %	95.5 %	-	-	-	-	58.1 %	80.4 %
Critica	95.6 %	99.7 %	95.4 %	99.6 %	97.4 %	99.7 %	56.8 %	96.5 %
Zcurve	98.1 %	96.9 %	98.0 %	97.1 %	99.2 %	98.5 %	63.5 %	92.0 %

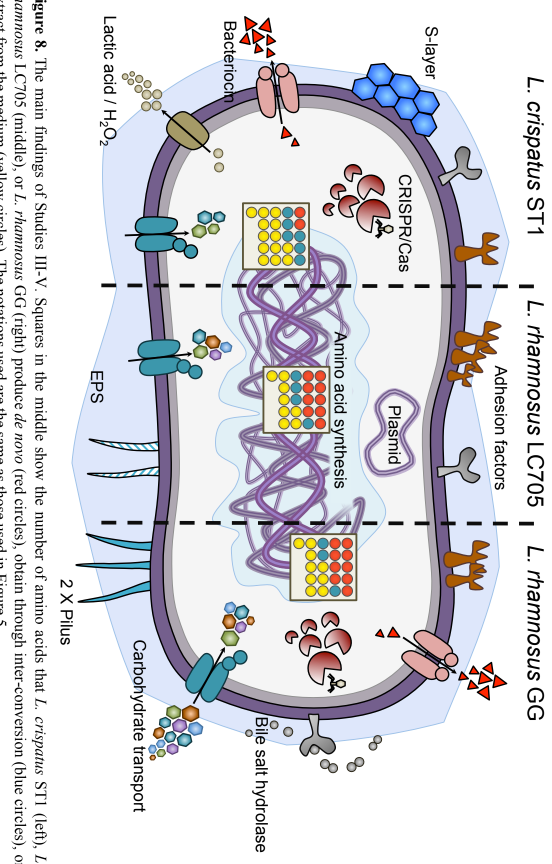
Overall, the three top-performing gene-calling systems were almost equal in quality. However, the rank of Glimmer as the performing method in five out of the eight performance tests provide a strong justification for the choice of Glimmer for initial gene calling. It is, however, possible that Glimmer might have missed some CDSs residing in genome regions with an abnormal base composition and failed to characterise the correct start site for some CDSs. However, the manual annotation phase, which constituted the manual editing of start sites and the screening of intergenic regions for missed CDSs, should have rectified these problems in the *L. rhamnosus* GG and LC705 and *L. crispatus* ST1 genomes.

4.4 Functional annotation of L. rhamnosus and L. crispatus

Protein functional prediction consisted of running a battery of automatic, mostly homology-based function prediction tools, followed by manual curation of the results. The tools developed in Studies I and II were also used. A specific focus was placed on CDSs that code for EPS biosynthesis, extracellular proteins, antimicrobial peptides, proteinaceous adhesion factors, enzymes, prophage-like proteins, and CRISPR-Cas system components. Figure 8 outlines the main findings of these analyses.

4.4.1 General functional prediction of L. rhamnosus and L. crispatus genes

The application of bioinformatic methods constituted a central component of the functional classification of the predicted proteins. In general, the bioinformatic analysis provided important insights into the physiology of L. rhamnosus GG, L. rhamnosus LC705, and L. crispatus ST1 and enabled the assignment of initial functions to 72-77% of the predicted proteins; 10-13% of the predicted proteins represented conserved proteins with unknown functional roles, and 13-15% of the predicted proteins lacked homology information and remained classified as hypothetical. The success rate of the annotation processes was similar to those described in various earlier (Kleerebezem et al., 2003; Pridmore et al., 2004; Altermann et al., 2005) and more recent (Forde et al., 2011; Macklaim et al., 2011) Lactobacillus genome studies, indicating the presence of a rather constant fraction of classifiable CDSs in any Lactobacillus genome regardless of publication date. Automatic protein function prediction was especially successful in defining genes that encode transcription factors, two-component regulatory systems, and enzymes, suggesting that the functional assignment of these gene types did not require much manual input. In contrast, the automated protein function prediction was less useful for proteinaceous adhesins, host-interaction factors, CRISPR-Cas systems, and transporters. This situation appears to be reminiscent of the results obtained in other Lactobacillus genome sequence studies, given the number of studies focusing on rediscovering these types of proteins in lactobacilli (Boekhorst et al., 2006; Zhou et al., 2008; Horvath et al., 2009; Kleerebezem et al., 2010; Kant et al., 2010;). Moreover, the concordance between the results generated by different protein function prediction



extract from the medium (vellow circles). The notations used are the same as those used in Figure 5 rhamnosus LC705 (middle), or L. rhamnosus GG (right) produce de novo (red circles), obtain through inter-conversion (blue circles), or Figure 8. The main findings of Studies III-V. Squares in the middle show the number of amino acids that L. crispatus ST1 (left), L.

procedures was low. In Study II, BLANNOTATOR generated a correct prediction (an annotation that was accepted by the human operator) for approximately 85% of the *L. crispatus* gene products for which it assigned a function. In contrast, the two other homology-based protein function prediction tools that were applied to this dataset generated supported annotations for approximately 58% (RAST) and approximately 69% (the best BLAST approach) of the gene products with approved functional annotation, indicating the importance of choosing the correct annotation method. However, it should be noted that different methods failed with regard to different proteins and that pooling the results was necessary in several instances. For example, the best-BLAST approach was poor at producing component composition annotations for phosphotransferase system (PTS) transporter genes, unlike the InterProScan; RAST failed at calling CRISRP-cas components, unlike BLANNOTATOR; and GO term predictions were overly general when calling adhesins but were highly useful in classifying proteases.

4.4.2 Host-interaction molecules in L. rhamnosus and L. crispatus strains

The computational prediction of localisation sites of L. rhamnosus GG and LC705 and L. crispatus ST1 proteins revealed that close to 10% of these predicted proteins are secreted and/or membrane-associated and are therefore potentially involved in processes such as nutrient acquisition, cell communication, microbe-host interaction, and adhesion (Lebeer et al., 2008; Kleerebezem et al., 2010; Segers & Lebeer, 2014). Although the values are in good agreement with those seen for closely related lactobacilli (Kleerebezem et al., 2010), these predictions provide only an initial guess of the secretomes. Particularly, a relatively limited number of concordances appeared between the predictions and experimentally determined cell envelope (Koskenniemi et al., 2011) or secreted (Sanchez et al., 2009) proteins in L. rhamnosus GG. There also is a noted disagreement between the results from different studies. For example, two subsequent studies have defined that the secretomes of strains GG and LC705 could contain over 80 (Zhou et al., 2008) and 700 (Kant et al., 2010) more proteins than observed in Study III. In the first case, most disagreements were related to proteins that are N-terminally (Study III defining 149-159 fewer predictions) or C-terminally (Study III describing 18-19 more predictions) anchored, despite of the obvious similarities in the underlying computational procedures. Which of the approaches is preferable remained unclear; however, the high rate of false-positives (84%) that were reported according to the N-terminally anchored sorting predictions of LocateP (Berlec et al., 2011) argues against the other study. Despite the uncertainty of the sorting predictions, subcellular-location predictors were used in Studies III and IV because homology-based functional inference failed to provide clues about protein sorting. GO annotation data from the cellular component ontology were assigned to approximately only 30% of the CDSs, whereas the ambiguous naming of known extracellular proteins excluded the use of gene names.

Bacterial surface polysaccharides are ubiquitous components of the cell envelope of lactobacilli and are purportedly involved in determining host-microbe interactions (Lebeer *et al.*, 2008). In particular, a long galactose-rich EPS molecule from *L. rhamnosus* GG is

important for required for optimal survival of strain GG inside the murine GIT (Lebeer et al., 2011) and an EPS with high rhamnose content from another L. rhamnosus strain has been shown to stimulate various cytokines in human cell-line experiments (Chabot et al., 2001; Péant et al., 2005). The EPS gene cluster of LC705 that was discovered in Study III exhibited high similarity to EPS loci present in four other L. rhamnosus strains and shown to produce an EPS with high rhamnose content (Péant et al., 2005). In contrast, the EPS locus in L. rhamnosus GG was genetically different from that of the L. rhamnosus LC705 and verified the presence of a previously identified EPS gene cluster in strain GG (Lebeer et al., 2008), providing groundings for the comparison of the genomic neighbourhoods of these two EPS gene clusters. Regarding L. crispatus, some strains have been observed to produce EPS (Donnarumma et al., 2014). However, the genetic composition of EPS loci in L. crispatus has remained uncharacterised, despite of the vast amount of literature on EPS clusters in closely related Lactobacillus genomes (Pridmore et al., 2004; Altermann et al., 2005; Callanan et al., 2008; van de Guchte et al., 2006; Azcarate-Peril et al., 2008). In Study V, eight L. crispatus strains were identified to include a gene cluster associated with EPS biosynthesis. Each of these eight clusters was predicted to comprise a set of five highly conserved genes encoding a transcriptional regulator, a polymerisation and chain length determination protein, a tyrosine protein kinase, a protein-tyrosine phosphatase, and a priming glycosyltransferase. In contrast, differences in the glycosyltransferase genes situated at the 5' end of the L. crispatus EPS clusters suggested that the EPSs might contain different sugar monomers and glycosidic linkages.

Adhesion to host tissues has long been considered a central factor and a prerequisite for the long-term colonisation by and realisation of the health benefits of probiotic bacteria (Lee & Salminen 1995; Pfeiler & Klaenhammer, 2007; Siezen & Wilson, 2010; Segers & Lebeer, 2014; Lebeer et al., 2008;). However, prior to Studies III-IV, few adhesins been identified in L. rhamnosus (Chan et al., 1985) or in L. crispatus (Antikainen et al., 2002; Hurmalainen et al., 2007; Edelman et al., 2012), suggesting that new insights into L. *rhamnosus* and *L. crispatus* adhesion factors were enabled by the analysis of their genome sequences. In Studies III and V, proteins were classified into adhesion- or colonisationrelated protein domain families. The domains were collected from PFAM, and their potential adhesion or colonisation associations were determined by manual examination of the corresponding literature. In Study III, this approach revealed over 30 putative proteinaceous adhesion and colonisation factors in both L. rhamnosus GG and LC705, including many that have later been verified experimentally (Vélez et al., 2010; Lebeer et al., 2012; von Ossowski et al., 2011) and some others with physiological roles that have yet to be verified, such as the 382.1 KDa protein in L. rhamnosus LC705 that contains several collagen-binding domains. The search also revealed SpaCBA pilin subunits in L. rhamnosus GG that has been proven to be critical for its efficient adherence to human cells (Study III; von Ossowski 2010; Lebeer et al., 2012). Repeating the analysis for L. crispatus using a revised domain list revealed nine to 13 adhesion- and colonisationrelated proteins in each L. crispatus strain, many of which are part of the L. crispatus core genome. Notably, this search failed to characterise the LEA-protein, which is critical for the adhesion of strain ST1 to vaginal epithelial cells (Edelman et al., 2012) and that was in Study IV shown to displace *Gardnerella vaginalis* from vaginal cells, indicating that LEA-mediated adhesion might involve some yet undisclosed bacterial adhesion domain. Intriguingly, application of the revised domain set to *L. rhamnosus* GG and LC705 returned only eight and ten proteins, respectively. A detailed investigation of the functional annotations of the two sequence sets revealed differences in carbohydrateactive enzymes, indicating that the first domain set might have included erroneous, nonadhesion-related PFAM models. In addition to the PFAM search, protein functional information was used in the search for adhesins. However, the scarcity of adhesion-related GO annotations and the inconsistent naming of known bacterial adhesins in public databases precluded the use of these search strategies at full power.

Although pilins were successfully reported by searching for adhesion- or colonisationrelated protein domain families, the approach used in Study III was found to be laborious for the genome-scale mining of hundreds of genomes. Thus, a new tool was developed in Study I for the systematic screening of pilus operons in bacterial genomes and then used to investigate the distribution of pilus clusters in all complete gram-positive prokaryotic genomes that had been deposited in NCBI database. Interestingly, putative pilus operons were found in 67 out of the 181 genomes analysed, including four lactobacilli genomes. Further analysis with all available Lactobacillus genomes that are listed in Appendix Table 2 demonstrated the presence of putative pilus genes clusters in 29 strains. Putative pilus gene clusters were identified in selected L. gasseri, L. reuteri, and L. ruminis strains. In addition, pilus operons were found to be an almost universal feature of the L. casei group, with the exception of L. rhamnosus MTCC 5462. The wide distribution of pilus gene clusters within the L. casei group has recently been described in other studies and for a greater number of L. rhamnosus genomes, revealing (i) the occurrence of SpaCBA pilin subunits in 4 out of 13 (Kant et al., 2014) and in 34 out of 100 (Douillard et al., 2013b) L. rhamnosus strains and (ii) the presence of SpaFED pilin subunits in all strains that were investigated (Douillard et al., 2013b; Kant et al., 2014). Of note, LOCP returned some false predictions based on the gene annotations. The fraction of false positives in the analysis was however tolerable and substantially less than the number of sequences in these protein collections that contained an LPXTG-motif or an E-box (typically used to search pilins).

4.4.3 Bacteriocins in L. rhamnosus and L. crispatus strains

Lactobacilli produce an extensive set of antimicrobial substances including metabolic byproducts (Daeschel, 1989; Leroy & Vuyst, 2004; Nes & Johnsborg, 2004) such as lactic acid, acetic acid, ethanol, diacetyl, and hydrogen peroxide, as well as bacteriocins (peptides or small proteins that exhibit antimicrobial activity; Riley & Wertz, 2002; Jack *et al.*, 2005). Notably, bacteriocin-producing lactobacilli are of great importance for use in food preservation (Leroy & Vuyst, 2004; Nes & Johnsborg, 2004; Cotter *et al.*, 2005; Mills *et al.*, 2011) because they can protect food against contamination with specific pathogenic and spoilage organisms, such as *L. monocytogenes* (Corr *et al.*, 2007) and *Clostridium tyrobutyricum* (Jiménez-Díaz *et al.*, 1993) without affecting harmless LAB. In Study III, an 8.7-kb putative type IIb bacteriocin locus was identified in the GG genome. In addition to two short bacteriocin structural genes, the locus appeared to encode five genes implicated in immunity, bacteriocin production, and bacteriocin processing, indicating that strain GG (like several other L. rhamnosus strains, Jacobsen et al., 1999) produces a bacteriocin. However, contradictory evidence exists regarding the role of bacteriocin in the antimicrobial activity of strain GG and whether the inhibition of Salmonella typhimurium by strain GG is due to bacteriocin activity (Silva et al. 1987; Jacobsen et al., 1999) or to lactic acid accumulation (De Keersmaecker et al., 2006). A similar locus was identified in the LC705 genome. However, this region appeared to be non-functional because two of its genes were truncated and were probably pseudogenes. The *in silico* analysis of the *L. crispatus* genomes described in Study V revealed loci in each strain, which encoded bacteriolysins that are similar to the previously described enterolysin A (Nilsen et al. 2003) and helveticin J (Joerger & Klaenhammer, 1990), which lyse sensitive cells by catalysing cell wall hydrolysis. In addition, vaginal L. crispatus isolates contained genes that are implicated in the production of class II bacteriocins, indicating that these strains might produce an active bacteriocin, as previously reported for L. crispatus ATCC 33820 (Kim & Rajagopal, 2001) and L. crispatus JCM 2009 (Tahara & Kanatani, 1997).

Although bioinformatics services exist for mining bacteriocin loci from genomes (van Heel et al., 2013), the annotation of bacterial genomes for bacteriocins is not an easy task due to the small size and low degree of conservation of bacteriocins and the fact that they are often omitted from genome annotations and/or lack descriptive functional descriptions. Furthermore, genome regions surrounding bacteriocin genes and genes that are implicated in both the production and processing of bacteriocins are often enriched in pseudogenes. During manual annotation, these problems can be tackled by searching the surroundings of bacteriocin-related genes for small ORFs and by comparing these ORFs to HMMs that correspond to bacteriocin-related sequences, as was performed in Studies III and IV. Alternatively, bacteriocin calling can be based on bacteriocin prediction systems, such as BAGEL (van Heel et al., 2013), as was done in Study V. In general, these two approaches should be very similar and result in comparable protein sets; however, differences are possible. For example, BAGEL revealed four class III bacteriocin loci for ST1, two of which were also identified in a manual search. However, analysis of the GG and LC705 genomes using BAGEL revealed seven bacteriocins, including only half of the bacteriocin type II leader motif proteins that were described in Study III; these findings indicate the difficulty of the prediction task involved, at least for these genomes.

4.4.4 Prophage elements and CRISPR loci

Based on the *in silico* identification of prophage-like regions (Lima-Mendez *et al.*, 2008), the genome of *L. rhamnosus* GG had two large and one short putative prophage element, whereas the genome of LC705 had one large and two short putative prophage elements. The prophage-like regions in the LC705 genome shared a low similarity to those in GG and resided at different locations, suggesting an important role for bacteriophage in *L. rhamnosus* evolution, as has previously been proposed for *L. casei* (Cai *et al.*, 2009) and is

reminiscent of the high degree of diversity seen among prophages that have been identified in the genomes of *L. gasseri*, *L. salivarius*, and *L. casei* previously (Ventura *et al.*, 2006). Consistent with the high level of lysogeny (77%) in vaginal *L. crispatus* strains (Damelin *et al.*, 2011), each vaginal *L. crispatus* isolate was predicted to have at least one putative prophage element in Study V. Conversely, the genome of *L. crispatus* ST1 revealed no putative prophage elements, possibly due to the fact that it was predicted to have a different type of CRISPR locus than the vaginal isolates. Overall, computational tools were successful in identifying prophage-like clusters, and the methods used appeared to capture all relevant prophage elements and resulted in only one false prediction: a prophage like-region in *L. crispatus* ST1 that harboured housekeeping genes. Moreover, the phage-finder services helped in the annotation of phage-related genes.

CRISPR-Cas systems constitute a widespread class of RNA-based immunity systems that control invasions of bacteriophages and plasmids in prokaryotes (Deveau et al., 2010; Marraffini & Sontheimer, 2010). These systems are present in approximately two-thirds of Lactobacillus strains (Horvath et al., 2009) and provide an exceptional tool for the control of phage infections (Marraffini & Sontheimer, 2010); a significant and prevalent threat that disrupts dairy fermentation cycles, thus stalling the manufacturing chain and lowering the quality of the end product (Mc Grath et al., 2007). In Study III, a genomic screen for CRISPR repeats identified the presence of one CRISPR array in the genome of strain GG. This region comprised of 24 perfect repeats, and the spacers showed substantial sequence identity with various L. rhamnosus-specific phages, indicating that they might be its phage targets. Further, the CRISPR array located next to four Type I cas genes, corroborating the possibility of a functional system. Intriguingly, the Cas-proteins showed notable similarity to those described in L. salivarius UCC188 and L. casei BL23 but not to those described in L. casei ATCC 344 (Horvath et al., 2009), indicating that these Cas-proteins have not followed an evolutionary development similar to that of their bacterial hosts. No CRISPR-Cas systems were detected in strain LC705. In Study V, full or partial CRISPR-Cas systems were identified in each of the studied L. crispatus strains. All vaginal L. crispatus isolates contained at least one Type II cas gene and a CRISPR array comprising 36-bp direct repeats and at least two to six spacer sequences each. Homology searches among the spacers, and public virus and plasmid sequences did not reveal the putative targets of the crRNAs, suggesting a pool of undisclosed vaginal bacteriophages and plasmids. However, many of the spacers were shared by different vaginal strains, indicating that these isolates might have encountered common invaders in the past. Unlike the vaginal L. crispatus isolates, L. crispatus ST1 was predicted to carry eight Type I cas genes and two CRISPR arrays comprising 15 and 16 repeats. The repeats were highly similar and resembled a repeat that was recently described as present in vaginal metagenome samples (Rho et al., 2012). In contrast, the spacers of these systems did not match known plasmid or virus sequences. Similar to the prophage search tools described above, the CRISPR array scanners were accurate and reliable. The tools uncovered all CRISPR-appearing genomic regions, accurately predicted CRISPR array boundaries, and resulted in only one false prediction; namely, a CRISPR array situated within the LCRIS 01228 gene in L. crispatus ST1. Thus, genome sequencing and computational biology provide a powerful means for annotating prophage elements and CRISPR arrays in Lactobacillus.

4.5 Genomics of *L. rhamnosus* and *L. crispatus* metabolism

Metabolic pathway reconstructions were used to investigate carbohydrate metabolism and amino acid biosynthesis pathways in L. rhamnosus GG, L. rhamnosus LC705, and L. crispatus ST1. Overall, substantial percentages of genes were identified as involved in transport (13-15%) or enzymatic reactions (22-25%; Studies III and IV), percentages that are consistent with those described for other L. rhamnosus and L. crispatus genomes (Table 7). The assignment of these genes into reference pathways provided some of the first insights into the biosynthetic capabilities of these organisms and indicated that strains GG and LC705 can synthesise nine amino acids de novo and synthesise three through inter-conversion. Additionally, genes for the conversion of serine to cysteine were annotated for plasmid pLC1, extending the biosynthetic potential of LC705 beyond that of L. rhamnosus GG, which appeared to lack the genes that are implicated in this conversion. These biosynthetic capabilities are reminiscent of those of L. casei ATCC 344, which was annotated to have nine amino acid biosynthesis routes (Makarova et al., 2006), and are almost comparable to the collection of metabolic pathways that was described for the versatile L. plantarum, which was predicted to contain complete pathways for the biosynthesis of most amino acids (Kleerebezem et al., 2003). Through de novo synthesis and amino acid inter-conversions, L. crispatus ST1 was predicted to be able to produce eight amino acids (Study IV). Bioinformatic analysis of vaginal L. crispatus genomes suggested that these and the strain ST1 share the same biosynthetic potential, except for CTV-05, which was predicted to be auxotrophic for aspartate (Study V). These biosynthetic capabilities are similar to those found in L. johnsonii NCC 533, L. acidophilus NCFM, and L. helveticus DPC 4571, which have been reported to have the ability to produce 4 (Pridmore et al., 2004), 10 (Altermann et al., 2005), and 4 (Callanan et al., 2008) amino acids (either de novo or as derivatives), respectively.

To compensate for their limited amino acid biosynthetic capabilities, many strains of *Lactobacillus* have developed sophisticated proteolytic and transport systems to obtain amino acids from their habitats (Pridmore *et al.*, 2004; Altermann *et al.*, 2005). Indeed, based on *in silico* analyses, *L. rhamnosus* GG, *L. rhamnosus* LC705, and *L. crispatus* ST1 contain arrays of extracellular proteinases, cytoplasmic peptidases, and peptide and amino acid transporters. This result suggests that these bacteria exhibit enhanced abilities to utilise exogenous amino acids and peptides. Notably, *L. rhamnosus* GG and LC705 were predicted to have nearly identical casein degradation systems, although only LC705 has the ability to degrade milk protein (Study III). Regarding pyrimidine and purine biosynthesis, *L. crispatus* ST1, *L. rhamnosus* GG, and *L. rhamnosus* LC705 resembled their close relatives. Similar to *L. johnsonii* NCC 533 (Makarova *et al.*, 2006), *L. crispatus* ST1 needs to obtain pyrimidines from its environment, whereas *L. rhamnosus* GG and LC705 were described for *L. casei* ATCC 334 (Makarova *et al.*, 2006).

Genomic analysis revealed that both *L. rhamnosus* GG and LC705 include a series of genes that code for sugar metabolism and that these strains exhibit rather similar carbohydrate metabolism. Two noteworthy exceptions were the maltose and rhamnose pathways, which were found to be intact and functional only in LC705. In addition,

analyses presented in Study III revealed GG-specific frameshifts in two genes that act in lactose utilisation, providing a plausible explanation for the results obtained from a sugar utilisation assay, which indicated that GG cannot utilise lactose, unlike its dairy counterpart, LC705 (Study III). Moreover, similar sets of glycosidase genes were predicted to be present in the genomes of *L. rhamnosus* GG and LC705. Based on functional annotations and protein-sorting analyses, approximately ten of these genes contribute to peptidoglycan hydrolysis and the decomposition of complex polysaccharides.

The *in silico* reconstruction of *L. crispatus* sugar utilisation pathways suggested that *L. crispatus* can use a range of carbohydrates (Study V), as has previously been reported for several other members of the *L. delbrueckii* group (Pridmore *et al.*, 2004; Altermann *et al.*, 2005; Azcarate-Peril *et al.*, 2008). As with amino acid biosynthesis, CTV-05 differed the most and was predicted to lack various sugar utilisation pathways that were present in the other strains, most likely because of the sequencing gaps that are present in the corresponding genomic loci. Interestingly, *L. crispatus* pathway data argue against the classical grouping of *L. crispatus* as a homofermentative species (Salvetti *et al.*, 2012). Instead, pathways for both homofermentation (the Embden-Meyerhof-Parnas pathway) and heterofermentation (the pentose phosphoketolase pathway) were observed in all *L. crispatus* strains investigated in Study V, as is typical of facultatively heterofermentative species (Pot *et al.*, 1994; Hammes & Vogel, 1995; Felis & Dellaglio, 2007; Salvetti *et al.*, 2012).

Practically, the reconstruction of metabolic pathways involved the use of a battery of bioinformatics tools and resources (Table 6). In Study III, KEGG reference pathways were noted to be useful for obtaining an overall view of metabolism but were often insufficient for explaining specific metabolic capabilities due to their complexity. In contrast, the reaction maps in the MetaCyc database contain on average only 4.4 reactions (Altman et al., 2013) and allowed the efficient examination of whether genes for a particular bioconversion were present in the organism (Studies III and IV). Overall, the data in these reaction pathway collections agreed well, which is consistent with the reported high degree of overlap (63%) between the KEGG and MetaCyc reaction spaces (Altman et al., 2013). However, inconsistencies were also observed. According to the KEGG, reaction 4.4.1.8 transforms pyruvate into cysteine. In contrast, MetaCyc found that this transformation involved another reaction and an enzyme that was not found in L. crispatus ST1, indicating a lack of cystathionine beta-lyase reaction in ST1. Other discrepancies involved reactions 1.1.1.351 (which catalyses the reduction of 6-phosphogluconate to ribulose 5-phosphate) and 1.2.1.59 (which catalyses the sixth step of glycolysis) that were associated with the phosphoketolase and Embden-Meyerhof-Parnas pathways only in MetaCyc and KEGG maps, respectively. In Study V, the presence of metabolic routes was tested by matching each strain's EC complement against the EC sets that are annotated to enable the conversion of a given starting compound to a particular end product. Metabolic routes between two given compounds were retrieved using the FMM web-server (Chou et al., 2009), which connects different KEGG maps and reconstructs metabolic pathways between metabolites. This approach greatly reduced the amount of work involved in understanding metabolic activities, although the test was found to yield some false

positive calls due to an unrealistic linking between some reactions. The FMM service also ignored some metabolic conversions that were recognised by the KEGG web service, possibly causing some pathways to remain undetected. Overall, use of the KEGG and MetaCyc databases and the reference-pathway approach were crucial for understanding the metabolic capabilities of *L. rhamnosus* GG, *L. rhamnosus* LC705, and *L. crispatus* ST1. The FMM server aided the process but at the cost of yielding some errors. However, the level of errors is acceptable considering the workload involved with reconstructing pathways manually, especially if there is a need to investigate tens or hundreds of genomes.

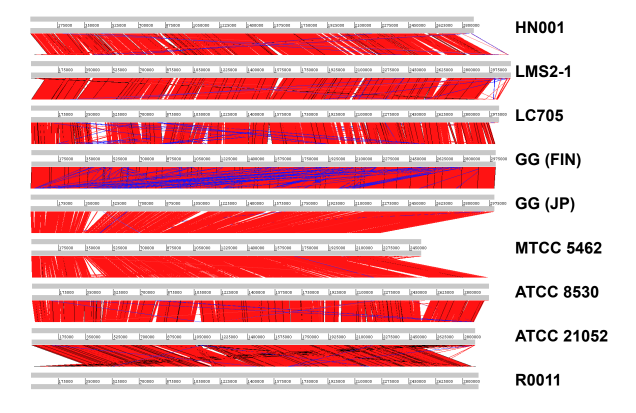


Figure 9. Alignment of nine *L. rhamnosus* genomes using ACT. The vertical bands between the genomes represent BLASTN matches (bit score \geq 500) between the two sequences. Forward and reverse matches are indicated by red and blue, respectively. The draft genome sequences were ordered and oriented according to the genome sequence of *L. rhamnosus* GG (FIN) using progressive Mauve.

4.6 Comparative genomics of *L. rhamnosus* and *L. crispatus*

Whole-genome alignment is a powerful tool for understanding the genetic forces that have shaped genomes (Ali *et al.*, 2013; Darling *et al.*, 2004). It has promoted the discovery of key lactobacilli effector molecules (Morita *et al.*, 2008) and has vastly expanded our

knowledge of the diversity and complexity that exists among Lactobacillus species (Forde et al., 2011; Boekhorst et al., 2004; Canchaya et al., 2006; Berger et al., 2007; Ventura et al., 2008; van de Guchte et al., 2006; Broadbent et al., 2012). In Study III, a wholegenome comparison revealed a high degree of stability among the genomes of L. casei group bacteria, a finding that has recently been confirmed in several other studies (Cai et al., 2009; Broadbent et al., 2012; Douillard et al., 2013a). The high level of genome relatedness and synteny is further underscored by a genome comparison of nine L. rhamnosus isolates, which revealed notable whole-genome synteny and no evidence of chromosomal rearrangements between these nine L. rhamnosus strains (Figure 9). Specifically, the nine L. rhamnosus isolates shared approximately 88% of their DNA. The colinearity was however noted to be punctuated by 1-182 genomic islands, which were generally consistent with genomic island predictions and exhibited abnormal sequence compositions, further attesting to their foreign origin and suggesting a key role for lateral gene transfer during L. rhamnosus evolution. In particular, the genomes of L. rhamnosus GG and L. rhamnosus LC705 shared extensive synteny, which was found to be punctuated by four (strain LC705) to five (strain GG) genome regions that displayed nucleotide composition deviations relative to the remainder of the genome. An examination of these genome regions in Study III revealed many genes of biomedical importance and implicated in EPS biosynthesis, host colonisation (SpaCBA pilins), prophage, and metabolism-related functions. It was further concluded that the GG-specific EPS and prophage and LC705-specific sugar utilisation islands were most likely acquired by horizontal gene transfer at a late point of divergence. In contrast, the phylogenetic distribution of the SpaCBA cluster in the lineage was best explained by a horizontal gene transfer by the common ancestor of L. casei and L. rhamnosus, as was apparent in more recent and broader comparative genome analyses that confirmed the presence of SpaCBA genes in the genomes of many L. casei strains (Broadbent et al., 2012) but only in a few L. rhamnosus strains (Douillard et al., 2013a; Douillard et al., 2013b; Kant et al., 2014).

As expected, genome alignments between L. crispatus strains revealed a high level of similarity and synteny (Study V). The genomes of 214-1 and SJ-3C-US were the most conserved; approximately 97% of their sequences were conserved in at least one other strain. The least related strain was the L. crispatus ST1; only approximately 82% of the genome of this isolate being alignable with the genomes of the nine vaginal isolates, underscoring its non-human origin. The data also indicated that the genome size differences observed in L. crispatus are not due to chromosomal insertions, inversions, deletions, or re-arrangements. Instead, horizontally acquired genomic islands and putative prophage elements explain a notable portion of the genomic differences in L. crispatus (Study V). Interestingly, the conservation of gene order in L. crispatus genomes has survived over a long evolutionary timescale and is also present in other lactobacilli of the L. delbrueckii group. Specifically, comparisons of the genomes of selected lactobacilli from the L. delbrueckii group revealed extensive sequence similarity and genome synteny between L. crispatus ST1 and other strains from this group (Figure 10) and indicated that the same overall gene order known to exist between most other members of the subgroup (Canchaya et al., 2005; Berger et al., 2007; Callanan et al., 2008) is also valid for strain ST1. The most noteworthy exception was observed by comparing the L. crispatus ST1 and

L. gasseri ATCC 33323 genomes; this comparison revealed sequence scrambling around the replication terminus, which is best explained using the fork replication theory (Tillier & Collins, 2000; Canchaya *et al.*, 2005).

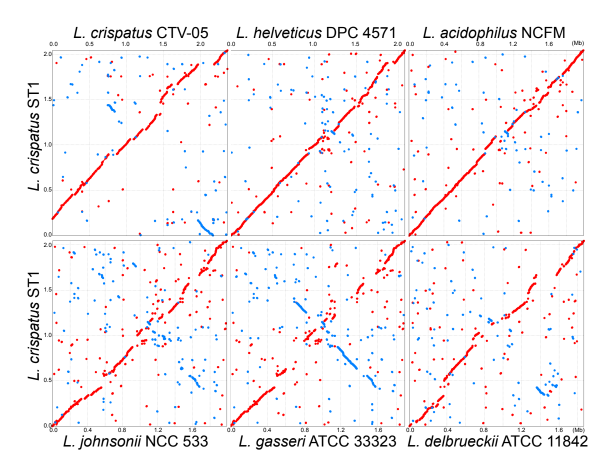


Figure 10. Comparison of the genome of *L. crispatus* ST1 with the genomes of selected lactobacilli of the *L. delbrueckii* group. PROmer alignments of the genome of *L. crispatus* ST1 with the genomes of *L. crispatus* CTV-05, *L. helveticus* DPC 4571, *L. acidophilus* NCFM, *L. johnsonii* NCC 533, *L. gasseri* ATCC 33323, and *L. delbrueckii* ATCC 11842 are shown. Red dots indicate conserved DNA sequences in the same orientation. Blue dots indicate conserved DNA sequences orientation.

In addition to pairwise alignment approaches, sophisticated genome alignment tools were tested to address the problem of whole-genome alignment. The *L. rhamnosus* GG, *L. rhamnosus* LC705, and *L. casei* ATCC 334 genomes were aligned using Mauve (Darling *et al.*, 2004; Rissman *et al.*, 2009), TBA (Blanchette *et al.*, 2004), and MUMmer (Kurtz *et al.*, 2004). Of these sophisticated genome alignment tools tested, the Mauve alignment tool (Darling *et al.*, 2004; Darling *et al.*, 2010) was considered the most suitable alignment package for studying these lactobacilli genomes. However, even Mauve did not provide an additional level of detail over simpler alignment approaches. Moreover, because the genomes in question were small (Table 7) and alignments were largely collinear, manual

interpretation of the results was not laborious. Thus, it appeared that simple alignment techniques were sufficient for collinear and closely related bacterial genomes, such as those that were investigated in Studies III-V. Among the pairwise alignment approaches examined, ACT (Carver et al., 2005) was most practical due to its ability to visualise several genomes simultaneously (see Figure 9). An obvious defect of this approach was that the results depend on the piling order and that an incorrect piling order can hide interesting genome differences. In contrast, summarisation of the alignment information over several dotplots was found laborious (see Figure 10), even though figures generated using dotplot visualisation utilities, such as Gepard (Krumsiek et al., 2007) and MUMmer (Kurtz et al., 2004), provided more information. Although not considered comparative genomics approaches, methods for sequence composition based genomic island prediction were noted to be useful for whole-genome alignment validation and for explaining the emergence of alignment-free genomic regions. However, sequence composition based genome island predictors were found to produce inconsistent results. For example, the IslandViewer resource (Langille & Brinkman, 2009) identified 102 genomic islands that constituted approximately 7% of the L. crispatus pan-genome. Shockingly, no genomic islands were recovered by all three methods included in the IslandViewer resource, and only six were supported by two methods (Study V).

4.7 Orthologue grouping of L. rhamnosus and L. crispatus genes

Three orthologue grouping tools were evaluated in Studies III-V. In Study III, orthologue groups were identified using a search strategy similar to that used in the InParanoid tool (Remm et al., 2001). Although accurate, conversion of the multiple pairwise orthologue groups into multi-species clusters was found to be a laborious process; thus, this approach was abandoned. Instead, orthologue groups were identified in Studies IV and V using the OrthoMCL tool (Li et al., 2003), which was noted to exhibit a good balance of sensitivity and specificity. In addition, a maximal clique approach based on RBHs was tested but not further used because long run time requirements (Study IV). Similar to previous ortholog assessment studies (Salichos & Rokas, 2011; Trachana et al., 2011), the three orthologue grouping tools were noted to produce comparable results. For example, L. rhamnosus GG and LC705 were predicted to have 364 and 430, respectively, CDSs lacking orthologous counterparts in the other L. rhamnosus strain or in L. casei ATCC 344 according to the InParanoid search strategy (Study III). In contrast, OrthoMCL identified 273 and 292 such CDSs, respectively. More recently, an RBH comparison was conducted on the protein collections of 13 L. rhamnosus strains (Kant et al., 2014). However, this analysis revealed only 94 and 26 unique elements for GG and LC705, partly because of methodological differences and partly because of the inclusion of new L. rhamnosus genomes, such as that of another strain GG isolate (Morita et al., 2013).

In Study V, the extent of the core and pan genomic potential of ten *L. crispatus* isolates was calculated. It was predicted that pan-genome of these ten strains comprised 3,929 orthologue groups, 1,224 of which were present in each strain (Study V). This set of core groups captured approximately 31% of the given collection of orthologous groups and was

comparable to those observed previously for L. casei (approximately 29%; Broadbent et al., 2012), L. paracasei (approximately 43%; Smokvina et al., 2013), and L. rhamnosus (approximately 43%; Kant et al., 2014). Based on the regression analysis, the core was also considered a good estimate of the final orthologue group repertoire of an unlimited number of L. crispatus strains. Regarding strain-specific functions, on average each L. crispatus strain was predicted to contain 131 orphan orthologous groups. However, the number of orphan groups ranged widely from 51 for strain MV-1A-US to 287 for strain FB077-07. Surprisingly, even the more phylogenetically distant L. casei and L. rhamnosus species have been reported contained approximately a 100 strain-specific elements (Broadbent et al., 2012; Kant et al., 2014), which might indicate that strain-specific gene pools of different Lactobacillus species are relatively same in size irrespective of the phylogenetic distance, life habitat, and genome size of the given species. Methodologically, the current Lactobacillus pan and core-genome investigations were comparable. However, some studies involved the use of relatively simple orthologue grouping approaches (Kant et al., 2014), whereas others were based on more sophisticated procedures (Broadbent et al., 2012; Smokvina et al., 2013; Study V). Some studies also failed to address functional annotation anomalies and inconsistencies that resulted from differences in the original protein function predictions. Given the relative high number of orthologue groups that contained sequences with inconsistent original protein function annotations (approximately 36% for L. crispatus; Study V), studies omitting the reannotation phase might have misidentified the correct biological role for some orthologues. Resolving the annotation anomalies among the L. crispatus using the BLANNOTOATOR tool at least improved the annotation consistency and allowed the assignment of more similar functional descriptions to orthologous sequences in Study V.

Comparative genomics data were also used to resolve gene-phenotype relationships. For example, this approach was applied in Study III to proteinaceous adhesion factors of L. rhamnosus GG and LC705 to aid in understanding the role of these adhesins in microbe-host interactions. Specifically, the initial screen for adhesion factors identified several types of proteins in L. rhamnosus GG (31 adhesins) and LC705 (37 adhesins) with adhesion- or colonisation-related protein domain families. Using comparative genomics, this protein set was narrowed to eight GG-specific candidates purportedly involved in determining its host-microbe interactions. Importantly, the gene-phenotype correlation analysis exhibited high specificity and sensitivity and was able to identify SpaC, which has recently been shown to be the single most important adhesin of GG (Lebeer *et al.*, 2012), as well as other proteins for which roles in adhesion and biofilm formation (Vélez et al., 2010) or mucus binding (Ossowski et al., 2011) have been verified. Interestingly, the current L. rhamnosus genome data would have resulted in a stronger association between SpaCBA genes and host cell binding if L. rhamnosus strains LMS2-1 and E800 had shown levels of adherence that were comparable to that of GG, based on the notion that the spaCBA genes are present only in the genomes of L. rhamnosus strains GG, LMS2-1, and E800 (Kant et al., 2014). All other GG-adhesins found in Study III have a counterpart in at least one other L. rhamnosus strain.

At the time of Study V, ten *L. crispatus* genomes were available in public databases. However, the full exploitation of these data was limited by the lack of commensurable phenotype information for the studied bacteria. Nevertheless, the genome comparisons that were conducted in Study V revealed interesting relationships between the isolation source and putative prophage elements and between the natural habitat of these strains and adaptive immunity systems, suggesting that different types of CRISPR-Cas systems are beneficial in different niches (Study V). In addition, ortholog assignments of the *L. crispatus* and *G. vaginalis* protein complements and projection of the adhesion factor information across species was able in Study V used to identify *L. crispatus* core-genome encoded proteins that are implicated in the competitive exclusion of *G. vaginalis*, providing an explanation for the inverse association between *L. crispatus* and *G. vaginalis* colonization in the human vagina (Fredricks *et al.*, 2007; Srinivasan *et al.*, 2012; Shipitsyna *et al.*, 2013).

4.8 Phylogenetic reconstructions

Similar to molecular phylogenetic approaches (Felis & Dellaglio, 2007; Salvetti *et al.*, 2012), the phylogenomic strategy adopted in Study III revealed that *L. rhamnosus* is closely related to *L. casei*. In particular, the study revealed a close phylogenetic relationship between *L. rhamnosus* GG and LC705 and demonstrated that among the 25 LABs that were tested, these strains were particularly phylogenetically close to *L. casei* ATCC 334. The relationship between *L. casei* and *L. rhamnosus* was supported with 100% confidence, corroborating the known assignment of *L. rhamnosus* to the *L. casei* and *L. paracasei* as well as *Lactobacillus zeae*, which was recently reclassified as *L. casei* by Salvetti *et al.*, 2012.

In Study V, a *Lactobacillus* phylogeny was derived from the concatenated alignment of 72,019 single-nucleotide polymorphisms from the core regions of three *L. acidophilus*, ten *L. crispatus*, five *L. helveticus*, and one *Bacillus subtilis* strains. The analysis yielded a tree with a high confidence and indicated that *L. crispatus* and *L. helveticus* were sister species to *L. acidophilus*. This finding contradicted some previous reports indicating that *L. crispatus* and *L acidophilus* cluster together first (Canchaya *et al.*, 2006; Felis & Dellaglio, 2007; Salvetti *et al.*, 2012) but agreed with some other phylogenomic studies (Kant *et al.*, 2011a). Among the *L. crispatus* cluster, the chicken isolate ST1 was the first to branch off from the others.

4.9 Intrafamily variation in Lactobacillaceae

In addition to the genome analyses presented in Studies III-V, a comparative analysis of the available *Lactobacillus* and *Pediococcus* genomes is presented here to better describe this family and to determine the scale and scope of the pan and core genomic potentials of these bacteria. Using BLAST (Altschul *et al.*, 1997) and OrthoMCL (Li *et al.*, 2003), the full complement of *Lactobacillaceae* protein sequences was assigned to 30,693 orthologue

groups. The pan-genome was found to be approximately 15-fold the size of a single genome and was predicted to be open, based on a power-law regression (positive exponent β =0.526±0.003, Figure 11). Notably, the pan-genome was found to grow by at least one orthologue group per additional genome until 3.18 million isolates have been sequenced, emphasising the need to sequence more Lactobacillaceae genomes. A relatively large fraction of the orthologue groups of any given isolate was conserved and shared by at least one other organism in this group. Genomes that were the only representative of their species, such as Lactobacillus kisonensis F0435, had the largest strain-specific gene pools (over 20% of their orthologue groups are orphans), whereas strains such as L. reuteri DSM 20016^T and JCM 1112^T, which originated from the same isolate, included only a few strain-specific elements. The average number of orphan orthologue groups in each Lactobacillaceae was 114, which is approximately the same average value as that reported for individual Lactobacillus species (Broadbent et al., 2012; Kant et al., 2014; Study V). As expected, the size of the core genome decreased with the addition of genomes: on average, ten genomes produced a core genome of 530 orthologue groups, whereas a set of twenty genomes produced a collection of 389 core orthologue groups. This trend is reminiscent of that observed in previous core genome analyses (Tettelin et al., 2005; Bottacini et al., 2010; Broadbent et al., 2012). It is also interesting that the Lactobacillaceae core genome curve was consistent with two previous total core estimates, which suggested 383 and 363 orthologue groups for a set of 20 and 21 Lactobacillus genomes, respectively (Kant et al., 2011a; Lukjancenko et al., 2012). Notably, extrapolation of the core genome curve showed that the core genome reaches a plateau at 59±10 orthologue groups for an infinite number of Lactobacillaceae (Figure 11), whereas the current Lactobacillaceae core genome included 66 orthologue groups, a value that approximates well the predicted core genome. Therefore, the strains included in the analysis can be judged to have represented the common features of the family Lactobacillaceae quite well. However, because some draft genomes contain up to hundreds of sequence gaps, genuine gene products might have been ignored in this analysis, thereby causing an under-estimation of the size of the core genome and a slight over-estimation of the size of the pan-genome. The extent of this error is unknown and remains to be elucidated. Moreover, because this extrapolation curve was the first to be fitted to Lactobacillaceae core genome data, the validity of the goodness of estimation remains open. Nevertheless, the data undoubtedly indicate the presence of large repertoires of undiscovered genes in Lactobacillaceae genomes that are yet to be sequenced and suggest that the Lactobacillaceae core is perhaps smaller than previously anticipated (Canchaya et al., 2006; Kant et al., 2011a; Lukjancenko et al., 2012).

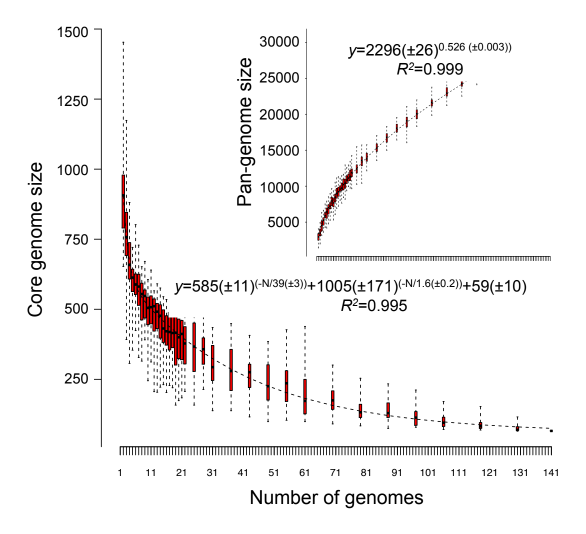


Figure 11. Pan- and core genomes of *Lactobacillaceae* according to the number of considered genomes. At each step, N genomes were chosen 50 times randomly, and the pan-genome and the core genome were recalculated. The dashed lines represent least squares fits to the medians, and R^2 describes the suitability of the fit. The boxes represent the 25–75 percentiles, the horizontal lines represent the medians, and the whisker data represent the extremes estimated from 50 iterations. The pan-genome curve is a least squares fit of the power law $y = k N^{\beta}$ to medians. An exponent $\beta > 0$ indicates an open pan-genome. The regression analysis for the core genome was performed by fitting a double exponential decay $y = \kappa_1^{(-N/\tau I)} + \kappa_2^{(-N/\tau 2)} + \Omega$ to the medians with a least square regression. Ω is the asymptotic core genome size.

5 Conclusions

Whole-genome sequencing of bacteria and genome mining is a powerful approach for addressing microbiological questions (Fleischmann et al., 1995; Mardis, 2008; Loman et al., 2012). However, the extraction of biological knowledge involves the handling of vast amounts of sequence data and is largely beyond human limits; thus, researchers rely on sophisticated computational procedures (Edwards & Holt, 2013; Richardson & Watson, 2013; Ali et al., 2013). In this thesis, two new algorithms were developed for laborious genome annotation tasks: LOCP and BLANNOTATOR. The LOCP service is based on the detection of genomic segments that are enriched in sortase and pilin-resembling genes and provides a major step forward in understanding the distribution of pilus appendixes in gram-positive bacteria. The other novel tool was designed for the functional annotation of bacterial protein sequences. Based on sequence similarity searches and clustering the annotation space, BLANNOTATOR yields DEs that are accurate and that do not suffer from annotation anomalies. While developing the algorithm, DEs were preferred over GO annotations, because of their great value in inferring host-interaction factors. In addition to automated protein function prediction methods, this study focused on the genomics of Lactobacillus and demonstrated the power of computer-assisted genome annotation for improving our understanding of the biochemistry, niche-adaptation, and host-interactions of lactobacilli. In particular, three Lactobacillus genomes were sequenced and annotated. Genome analysis identified genes that are instrumental for survival and of industrial value. Importantly, for each of the investigated species, comparative genomics identified hostinteraction factors, the roles of which were validated using experimental methodologies.

First, two software tools were designed for laborious bacterial genome annotation processes for which bioinformatics solutions did not exist prior to these studies: one for locating pilus operons and another for DE prediction. In Study I, a new algorithm was presented to manage the cumbersome and slow process of locating pilus operons in grampositive genomes. Unlike the error-prone and arduous manual curation of pilus gene regions, LOCP was designed for ease of use and to accurately locate pilus operons from any set of sequences. For example, LOCP analysis of the 135 Lactobacillus genomes that are listed in Appendix Table 2 revealed 29 potential pilus carriers among these organisms, many of which have been recently verified using other approaches (Broadbent et al., 2012; Kant et al., 2014). In Study II, an effort was made to explore the possibilities for improving annotation anomalies and inconsistencies, which are rather common in sequence databases (Andorf et al., 2007; Schnoes et al., 2009). During the manual curation processes, existing protein function prediction tools were noted to inconsistently classify orthologues and genes that were judged to share the same protein functions. Because manual correction of these annotation errors was laborious, BLANNOTATOR was developed in Study II. Comparisons based on simulated data indicated that BLANNOTATOR was better than the five other protein function prediction methods that were tested in Study II and that this method made function calls with a highly consistent nomenclature. However, even the simplest protein function prediction strategies, such as the best BLAST approach, performed well in Study II; this illustrates that substantial improvements using homology-based annotation alone are unlikely to be made in the future. However, the most important benefit of the method developed in Study II resulted from its ability to systematise DE calls, thereby influencing the manual curation process and the comparative analysis of protein function. The ability to classify proteins into consistent classes was especially useful for determining host-interaction factors and when dealing with multiple organisms, such as in Study V, where the functional catalogues of ten *L. crispatus* strains needed to be commensurable.

Second, the genomes of L. rhamnosus GG and LC705 were sequenced and annotated in Study III. These organisms were chosen for genomic investigation to map molecules that might explain their successful use in preventing and treating various diseases (Hoisak et al., 2010; Hatakka et al., 2001; Isolauri et al., 1991; Guandalini et al., 2000; Szajewska & Mrukowicz, 2001; Kalliomäki et al., 2001; Kalliomäki et al., 2003; Kalliomäki et al., 2007; Kajander et al., 2005; Kajander et al., 2008) and to identify the bacterial components that are responsible for their adhesion to human tissues (Jacobsen et al., 1999; Tuomola et al., 2000; Saxelin et al., 2010). Among the approximately 3,000 genes in each strain, Study III disclosed those that are implicated in EPS biosynthesis, sugar and peptide usage, and the production of a putative bacteriocin. The study also revealed the presence of two gram-positive pilus gene clusters in L. rhamnosus genomes. Using immunoblotting and immunogold electron microscopy, the expression of a spaCBA gene cluster in strain GG was in Study III shown, and the presence of this pilus structure on the cell surface was confirmed. This finding represented the first description of such a pilus structure in lactobacilli and established the presence of these adhesion structures in non-pathogenic bacteria. Furthermore, the mucus-binding capacity of the SpaC subunit and its importance for the adhesion between strain GG and human intestinal mucus was demonstrated in Study III. Recently, further evidence has accumulated that describes the role of SpaC pilin in adhesion (von Ossowski et al., 2010; Lebeer et al., 2012), confirming that this pilin plays a crucial role in the adhesion of strain GG to human tissues. Although recent studies have demonstrated the absence of SpaFED pili on the cell surface of L. rhamnosus GG (Reunanen et al., 2010), the fact that the SpaFED genes are present in the genomes of many L. rhamnosus (Douillard et al., 2013b; Kant et al., 2014) and L. casei (Broadbent et al., 2012) strains corroborates their potential importance in the adhesion processes of L. casei group bacteria. Notably, the discovery that the SpaC pilin is essential for the efficient adherence of strain GG to human tissues was based on ortholog analysis and represents an impressive demonstration of the power of comparative genomics in predicting host-interaction factors in lactobacilli. Specifically, 31 proteins purportedly involved in adhesion or colonisation were detected in the GG. Only eight of these proteins were however predicted to be GG-specific, thus providing a plausible explanation for the differing adhesion characteristics of the two L. rhamnosus strains to human mucus (Jacobsen et al., 1999; Tuomola et al., 2000; Saxelin et al., 2010). It is clear that without this type of analysis, all 31 of the genes identified in Study III would have had to been validated experimentally. In contrast to the successful detection of adhesion factor components, the search for molecules mediating immune responses failed, because not much known about the role of LC705 in disease alleviation. This failure indicates the role of commensurable phenotype information in comparative genomics. Overall, the L. rhamnosus GG and LC705 genomes are the first two genomes of free-living organisms to

have been sequenced in Finland and have provided a notable framework for various *Lactobacillus* comparative genomics studies and other studies that attempt to understand the mechanism underlying the interaction of probiotics with host tissues.

Finally, the genomic potential of L. crispatus ST1 was determined in Study IV and compared to those of nine vaginal L. crispatus isolates in Study V. Chicken-isolated ST1 was subjected to whole-genome sequencing to reveal the coding regions of genes encoding LEA and other adhesins. Although a complete genome could not be produced in the study, the project generated a high-quality reference genome with only one unresolved region (within the *lea* gene) and provided a valuable insight to the genomic foundations of an important urogenital species. Bioinformatic analysis of the nearly complete genome of ST1 identified approximately 2,100 genes, including genes that are implicated in EPS biosynthesis, antimicrobial activity, acquired resistance against bacteriophages, and adhesion. The metabolic pathways constructed in Study V revealed auxotrophy for 12 amino acids. The analysis also highlighted the presence of both pentose phosphate and in L. crispatus, defining species glycolytic pathways the as facultatively heterofermentative; this finding contradicts earlier assumptions (Salvetti et al., 2012; Salvetti et al., 2013). Extensive comparative genomic analysis provided evidence for considerable sequence identity and synteny among the genomes of the ten L. crispatus isolates that were investigated in Study V. Moreover, the study revealed genes in the L. crispatus core genome coding for adhesins that are involved in the competitive exclusion of G. vaginalis from vaginal cells, thus providing an attractive explanation for the proven inverse association between L. crispatus and G. vaginalis colonisation in the human vagina (Fredricks et al., 2007; Srinivasan et al., 2012; Shipitsyna et al., 2013). Importantly, the competitive exclusion process was demonstrated to result from the coreencode LEA protein (Edelman et al., 2012) by measuring the adhesion capacity of vaginal L. crispatus and G. vaginalis isolates to vaginal cells in the presence and absence of pretreatment with LEA-specific Fab fragments. Collectively, Studies IV and V established the genetic landscape of key urogenital lactobacilli. They suggested significant relationships between the life environments of different L. crispatus isolates and their adaptive immunity systems and revealed proteins that might protect the vagina from G. *vaginalis* and bacterial vaginosis. These studies also demonstrated the benefit of including phenotypically characterised strains in sequencing projects and indicated that the linking of bacterial traits to genes is easier when using well-characterised strains (as in Study III).

In conclusion, Studies III-V highlighted the power of whole-genome sequencing for generating new hypotheses about lactobacilli and their host-interaction factors. However, the success of sequencing studies is affected by several choices. First, one needs to choose an appropriate sequencing method. At present, lactobacilli have been sequenced largely using the Roche 454 platform (Appendix Table 2), which provides a long read length and good accuracy (Liu *et al.*, 2012; Quail *et al.*, 2012). This sequencing approach is feasible, given that longer reads appear to result in assemblies that are more continuous than shorter reads. However, sequencing machines that yield comparable read lengths at a lower price per sample (Table 1). Second, if the assembly phase fails to assign reads into single contigs, the desired genome-finishing level needs to be chosen. Although draft

genomes are cheap and fast to produce, they miss data and can suffer from truncated genes (Klassen & Currie, 2012). The problems are mainly caused during the assembly of low-coverage areas, ribosomal gene clusters, and repetitive elements; however, there is no guarantee that important genes will not be missed in draft genomes. Nevertheless, the risk of missing an important gene is rather small, as was evident from a high level of sequence relatedness and colinearity that was in Study V found between the draft and high-quality *L. crispatus* genomes.

The third choice to make is to decide the level of manual curation of the genome data. In Studies III and IV, a semi-automated annotation strategy was used, whereby the gene start sites and functional annotations were subjected to manual curation. In this process, the manual curation of gene start sites was based on comparing the gene models among lactobacilli and affected only a small fraction of L. rhamnosus GG, L. rhamnosus LC705, and L. crispatus ST1 CDSs, suggesting that the current gene-calling systems are accurate (as shown in Table 8) or, at least, call genes consistently across lactobacilli. The manual curation of function calls involved comparing protein function calls, reviewing the protein functions of orthologous sequences, and choosing the best alternatives. This step made the data more interpretable and enabled the description of gene functions with nomenclature that was more consistent; however, this step affected only some hundreds of genes in each genome. Moreover, the refinements made were mostly cosmetic and rarely changed the predicted protein function completely, indicating that the manual curation did not provide any additional advantage for a large majority of L. rhamnosus GG, L. rhamnosus LC705, and L. crispatus ST1 proteins. Specifically, the bioinformatic tools that were used in Studies III-V (Table 6) performed remarkably well for enzymes, phage-related proteins, and transcription factors in the given bacteria. In contrast, the classification of genes that were implicated in EPS production, adhesion, and antimicrobial activity was less satisfactory (Studies III-V). Thus, it was necessary to generate annotation data for these proteins using specific tools and to manually refine the results provided by the automated methods. It was also noted that in various instances, DEs were more informative than the GO terms of the corresponding gene products. In particular, GO terms were of limited value in the quest for genes that are involved in immunomodulatory processes, bacterialhost interaction, and the production of bacteriocins. Unfortunately, the large vocabulary that is present in DEs precludes their use for the automated cataloguing of protein genes into categories. Surprisingly, the various gene callers and protein function prediction approaches examined provided results of almost equal quality, indicating that the choice of gene caller or automated protein function prediction method might not be as critical as stated previously (Bakke et al., 2009). It remains to be determined whether these findings are specific to Lactobacillus or can be generalised to a wider array of bacteria.

The power of comparative genomics and genome island prediction in identifying genes determining host-microbe interactions was also demonstrated. In Study III, comparative analyses suggested key roles for the SpaCBA pilins in the adhesion of *L. rhamnosus* GG to human tissues. In Study V, protein comparison data revealed a role for the LEA protein in the competitive exclusion of *G. vaginalis* from vaginal cells. Notably, both discoveries were confirmed experimentally. Based on the successful analyses, it can be concluded that comparative genomics provides an appealing starting point to call host-interaction factors

and gene-phenotype associations in *Lactobacillus*. However, the success of comparative approaches in resolving correlations between genotypes and phenotypes appears to depend more on the amount of commensurate biological information available for each organism (Study III) than on the number of genomes under study (Study V). Comparative analyses should therefore be based on organisms for which commensurate empirical information is available rather than on the genome assemblies of poorly characterised bacteria, such as those generated as a part of the HMP (Nelson et al., 2010). The HMP is however useful for studies aiming to find functions universally conserved in the given set of organisms, as in Study V, where comparative genomics was used to investigate the genetic mechanisms underlying the inverse association between L. crispatus and G. vaginalis colonisation in the human vagina (Fredricks et al., 2007; Srinivasan et al., 2012; Shipitsyna et al., 2013). Genomic island calling represents another useful computational method for the annotation of host-microbe interaction, as was exemplified in Study III by the finding of an island in L. rhamnosus GG that contains genes for 3 secreted pilins and a pilin-dedicated sortase. The approach was also useful in the annotation of putative prophage elements in vaginal L. crispatus strains. Based on Studies III-V, it can be argued that the use of genomic island predictors for *Lactobacillus* genomes is advisable and these predictors can reveal genomic regions that code for functions that differentiate the given strain from others. Importantly, this approach can even be fruitful for identifying genes that underlie strain-specific traits in the absence of evolutionary counterparts. Nevertheless, orthology grouping and genome island prediction are simply methods that allow finding patterns of sequence conservation. To understand the biological relevance of the conservation patterns and to filter patterns, sequences underlying these patterns need to be associated with function information, using tools such as LOCP (Study I) and BLANNOTATOR (Study II). For example, application of the LOCP tool to the Study III data helped to understand which of the 364 GG-specific proteins are relevant for adhesion and provided additional information for the support of pilus-encoding gene clusters in strains GG and LC705. BLANNOTATOR on the hand played a pivotal role in Study V and provided to classify proteins into consistent classes. It is clear that without this analysis, functions of many host-interaction factors would have remained undetected.

To conclude, this study has described two bioinformatics algorithms for cumbersome genome annotation tasks and has disclosed the genomes of two also human-associated *Lactobacillus* species: *L. rhamnosus* and *L. crispatus*. The algorithms yielded impressive accuracy and were of great value in improving our understanding of *Lactobacillus* host-interaction factors. Annotation of the *L. rhamnosus* and *L. crispatus* genomes has provided new insights into the physiological, genetic, biochemical, and fermentative properties of two biomedically important *Lactobacillus* species. Markedly, analyses revealed molecules involved in host-interaction and those that might protect the vagina from pathogen attack, thereby representing a major advance in understanding the host-interaction mechanisms of lactobacilli.

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Appendixes

Method	Description	Ref
Base calling in Sanger technology	r technology	
Phred	Base caller for Sanger sequence traces published in 1998. Introduced the logarithmically	Ewing & Green, 1998
	related base-calling error probabilities	
KB Basecaller	Base caller developed by ABI	1
Base calling in NGS technologies	ichnologies	
Rolexa	Parametric base caller for Illumina sequence data. IUPAC codes used to describe base	Rougemont <i>et al.</i> , 2008
	calling quality	
Alta-Cyclic	Mixed Parametric base caller for Illumina data that uses SVMs	Erlich & Mitra, 2008
BayesCall	Parametric base caller for Illumina sequence data	Kao <i>et al.</i> , 2009
All Your Base	Method that uses a completely empirical model for generalising both cross-talk and	Massingham & Goldman, 2012
	phasing	
BM-BC	Bayesian method of base calling for Illumina sequence data. Uses a hierarchical model	Ji <i>et al.</i> , 2012
	that accounts for three sources of noise in the data	
Ibis	Fully empirical SVM-based base caller for Illumina data	Kircher <i>et al.</i> , 2009
Pyrobayes	Base caller for 454 sequence data. Adapts an empirical prior on the homopolymer length	Quinlan <i>et al.</i> , 2008
HPCall	454 base calling method that make use of a weighted Hurdle Poisson model	Beuf et al., 2012
Rsolid	R package for normalizing intensity data from SOLiD platform. Should be applied before	Wu <i>et al.</i> , 2010
	calling colors	
Quality analysis and read manipulation	ead manipulation	
FastQC	Quality control tool for high throughput sequence data	1
PrinSeq	Bioinformatic tool for quality control and data preprocessing of genomic datasets.	Schmieder & Edwards, 2011
BIGpre	Quality assessment package for NGS data	Zhang <i>et al.</i> , 2011
NGS QC Toolkit	Standalone and open source application for quality check and filtering of high-quality	Patel & Jain, 2012
	data	

Appendix Table 1. Software and databases that can be used as sources of bacterial genome projects.

RAY	MaSuRCA	SPAdes	De Bruijn graph bas	MIRA	SGA	EDENA	Newbler	Overlap-based genome assembly	VCAKE	SSAKE	Greedy assemblers	Hybrid-SHREC	ECHO	HITEC		Reptile	Read correction		Staden package	fastx	Cutadapt	NARWHAL
Assembler for simultaneous assembly based on reads from a combination of sequencing	Algorithm that transforms paired-end reads into longer super-reads. Capable of handling reads from different sequencing platforms	De Bruijn graph assembly at multiple k -mer sizes. Ideal for genomes of varying coverage	De Bruijn graph based genome assembers	Whole-genome sequence assembler for Sanger, 454 and Illumina	Set of memory efficient assembly algorithms based on the FM-index	Exact de novo assembler dedicated to process very short reads	Whole-genome assembler designed for 454 sequence data	me assembly	Greedy short read data assembler with robust error correction	De novo short read DNA assembler. Searches progressively for perfect 3' read matches		Read correction tool designated to account substitutions, insertions and deletions	k-spectrum based error correction algorithm. Does not require user-specified	Error correction algorithm that uses suffix trees to identify and correct substitution	errors	k-spectrum based error correction algorithm. Detects and corrects substitution		analysis	Package including a number of tools for DNA sequence assembly, editing and sequence	Toolkit for NGS data preprocessing	Finds and removes adapter sequences from high-throughput sequencing data	Software pipeline to automate the primary analysis of Illumina data. Includes de- multiplexin component
Boisvert et al., 2010	Zimin <i>et al.</i> , 2013	Bankevich <i>et al.</i> , 2012		Chevreux et al., 2004	Simpson & Durbin, 2012	Hernandez <i>et al.</i> , 2008	Margulies et al., 2005		Jeck et al., 2007	Warren <i>et al.</i> , 2007		Salmela, 2010	Kao <i>et al.,</i> 2011	Ilie <i>et al.</i> , 2011		Yang <i>et al.</i> , 2010			Staden et al., 1999	I	Martin, 2011	Brouwer <i>et al.</i> , 2012

	derive model parameters. RBS information used to locate the correct start site	
Besemer <i>et al.</i> , 2001	Popular gene caller suite. Includes GeneMarkS that uses a self-training procedure to	GeneMark
Delcher <i>et al.</i> , 2007	Interpolated Markov model based CDS predictor. Uses long orfs to train CDS models	Glimmer
	lictors	Ab initio CDS predictors
Yao <i>et al.</i> , 2012	Graph accordance assembly program	GAA
Nijkamp, <i>et al.</i> , 2010	Graph-based algorithm for integration of several de novo and comparative assemblies	MAIA
Sommer <i>et al.</i> , 2007	Basic genome assembler for merging one or two sequence sets	Minimus2
	itors	Assembly integrators
	reference genome based scaffolding	
Swain <i>et al.</i> , 2012	Post-assembly genome-improvement toolkit that has modules for both mate pair and	PAGIT
	to the analysed genome at the protein level	
Bartels et al., 2005	Uses gene pairs from a related reference genome. Reference genome genes are mapped	BACCardI
	assembly and a reference assembly	
Richter et al., 2007	Scaffolder that order and orient contigs based on matching sequences in a target	OSLay
Nagarajan <i>et al.</i> , 2008	Scaffolding using optical restriction mapping	SOMA
Boetzer <i>et al.</i> , 2011	Stand-alone program for scaffolding pre-assembled contigs using paired-read data	SSPACE
Pop <i>et al.</i> , 2004a	General purpose scaffolder that can use mate-pair data or reference genome alignments	Bambus
		Scaffolders
Pop <i>et al.</i> , 2004b	Comparative assembler based on MUMmer	Amos-Cmp
Nusbaum <i>et al.</i> , 2008	Variant ascertainment algorithm. Assemble reads using reference genome for assistance	VAAL
	Reference-based genome assemblers	Reference-based g
	larger genomes and one of the best assemblers when used with default parameters	
Simpson et al., 2009	De novo sequence assembler that supports parallel computing. Capable of assembling	ABySS
	sequence libraries	
Zerbino & Birney, 2008	Sequence assembler dedicated to short read sequences. Can make use of multiple	Velvet
Li <i>et al.</i> , 2010	Short-read assembly method designed for Illuman reads	SOAPdenovo
Butler <i>et al.</i> , 2008	Assembler for large genomes. Requires at least two types of read libraries to work	ALL-PATHS
	platforms	

EasyGene	HMM-based gene finder. Extensions of similarities in Swiss-Prot are used to learn parameters	Larsen & Krogh, 2003
Prodigal	Prokaryotic gene caller that uses dynamic programming to find CDSs. Suitable also for	Hyatt <i>et al.</i> , 2010
	organisms with high GC genomes	
ZCURVE	Gene caller based on the Z-curve representation of the DNA sequences	Guo <i>et al.</i> , 2003
Evidence-base CDS predictors	dictors	
ORPHEUS	Gene caller that guide gene prediction based on database similarity search	Frishman <i>et al.</i> , 1998
CRITICA	Gene calling system that uses comparative analysis to derive statistical characteristics of	Badger & Olsen, 1999
	CDSs. Initial coding regions are detected based on non-synonymous mutations	
CDS model integrators		
Reganor	Combines evidence from Glimmer and CRITICA to produce a consensus gene model	McHardy et al., 2004
YACOP	Produces a consensus gene model by integrating gene models predictions of CRITICA,	Tech & Merkl, 2003
	Glimmer and ZCURVE	
CDS model refinement tools	tools	
GenePRIMP	Gene prediction improvement pipeline. Handle various types of gene calling anomalies	Pati <i>et al.</i> , 2010
Mugsy-Annotator	Identifies anomalies in annotated gene structures based on comparative genomics.	Angiuoli et al., 2011
	Useful for standardising annotations across closely related organisms	
ORFCor	Corrects annotation inconsistencies based on consensus start and stop positions derived	Klassen & Currie, 2013
	from sets of closely related orthologs	
MICheck	Microbial genome checker is a tool for finding missed or inaccurate gene annotations	Cruveiller et al., 2005
	and frameshifts	
ncRNA predictors		
RNAmotif	RNA secondary structure definition and search algorithm	Macke <i>et al.</i> , 2001
RNAmmer	HMM-based gene predictor to annotate 5s, 16s and 23s ribosomal RNA in full genome	Lagesen <i>et al.</i> , 2007
	sequences	
QRNA	Gene finding system that uses comparative genome sequence analysis to guide RNA	Rivas & Eddy, 2001
	prediction	

 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tnmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes Bioinformatic software that uses covariance models to find intrinsic terminators CR1SPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows Extension to the PILER family of repeat analysis algorithms for fast and accurate identification of CR1SPR repeats Web tool offering tools to detect CR1SPRs and for their comprative analysis Software for repeat analysis on a genomic scale Discovers repetitive substrings in DNA 	Wagner <i>et al.</i> , 2007	Package to identify known insertion sequences in bacterial genomes	IScan
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverlear Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses covariance models RNA motif finding algorithm that uses covariance models to find intrinsic terminators Bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows Extension to the PLER family of repeat analysis algorithms for fast and accurate identification of CRISPR repeats Web tool offering tools to detect CRISPRs and for their comprative analysis Software for repeat analysis on a genomic scale Discovers repetitive substrings in DNA 	Siguier <i>et al.</i> , 2006	Database of insertion sequences isolated from eubacteria and archae	ISfinder database
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models MA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows Extension to the PILER family of repeat analysis algorithms for fast and accurate identification of CRISPR repeats Web tool offering tools to detect CRISPRs and for their comprative analysis Software for repeat analysis on a genomic scale Discovers repetitive substrings in DNA 			Insertion sequences
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs Gene ral RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows Extension to the PILER family of repeat analysis algorithms for fast and accurate identification of CRISPR repeats Web tool offering tools to detect CRISPRs and for their comprative analysis 	Price <i>et al.</i> , 2005	Discovers repetitive substrings in DNA	RepeatScout
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows Extension to the PILER family of repeat analysis algorithms for fast and accurate identification of CRISPR repeats Web tool offering tools to detect CRISPRs and for their comprative analysis 	Kurtz & Schleiermacher, 1999	Software for repeat analysis on a genomic scale	REPuter
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses covariance models to furnination and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes Bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows Extension to the PILER family of repeat analysis algorithms for fast and accurate identification of CRISPR repeats Web tool offering tools to detect CRISPRs and for their comprative analysis 			Repeats
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes Bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows Extension to the PILER family of repeat analysis algorithms for fast and accurate identification of CRISPR repeats 	Grissa et al., 2007	Web tool offering tools to detect CRISPRs and for their comprative analysis	CRISPRFinder
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes Bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance by sliding windows 	Edgar, 2007	v or repeat analysis algorithms for fast and	FILER-CR
 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes Bioinformatic software that uses covariance models to find intrinsic terminators CRISPR recognition tool. Searches for series of k-mers separated by a similar distance 			
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 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes Bioinformatic software that uses covariance models to find intrinsic terminators 			CRISPR arrays
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 Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance models 	Kingsford et al., 2007	Annotates low-energy hairpins followed by a stretch of thymines in bacterial genomes	TransTermHP
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired scan-SE Gene caller that uses covariance models to detect tRNAs nal Gene ral RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases scan Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences k Detects RNase P RNA genes using pattern matching and covariance models nder RNA motif finding algorithm that uses both expectation maximization and covariance			Intrinsic terminators
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf scan-SE Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases scan Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences kt Detects RNase P RNA genes using pattern matching and covariance models RNA motif finding algorithm that uses both expectation maximization and covariance		models	
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases scan Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences k Detects RNase P RNA genes using pattern matching and covariance models	Yao <i>et al.</i> , 2006	RNA motif finding algorithm that uses both expectation maximization and covariance	CMFinder
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments prm Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf nal Gene caller that uses covariance models to detect tRNAs for searching DNA sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in genomic DNA sequences	Yusuf et al., 2010	Detects RNase P RNA genes using pattern matching and covariance models	Bcheck
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments prm Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases Gene finding system that uses covariance models to annotate SRP RNA genes in		genomic DNA sequences	
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models for searching DNA sequence databases	Regalia <i>et al.</i> , 2002	to annotate SRP RNA genes	SRP-scan
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments prn Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs General RNA structure and sequence similarity search tool that uses covariance models		for searching DNA sequence databases	
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments prm Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf Gene caller that uses covariance models to detect tRNAs	Nawrocki <i>et al.</i> , 2009		Infernal
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure on homology with known RNA consensus sequences and ability to form a base-paired cloverleaf	Lowe & Eddy, 1997	Gene caller that uses covariance models to detect tRNAs	tRNAscan-SE
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure		on nomology with known KNA consensus sequences and ability to form a base-paired cloverleaf	
Predicts structurally conserved and thermodynamically stable RNA secondary structures in multiple sequence alignments	Laslett & Canback, 2004	Employs heuristic algorithms to predict tRNA and tmRNAs based on secondary structure	Aragorn
	Wasnieti <i>et al., 2</i> 005	in multiple sequence alignments	KNAZ
		Department of the property and the property of	

ISsaga	Web application pipeline for insertion sequence annotation	Varani et al., 2011
Prophages		
ACLAME	Database dedicated to the collection and classification of mobile genetic elements.	Leplae <i>et al.</i> , 2004
	Provides access to a list of functional ontologies	
PhiPsy	Phage detection algorithm that calls prophage regions based on seven distinctive	Akhter et al., 2012
	characteristics of prophages including similarity of phage proteins as well as other	
	statistics	
Phage_Finder	Heuristic computer program to identify prophage regions in bacterial genomes. Uses	Fouts, 2006
	comparative genomics appraoches to locate prophage regions	
ProphageDB	Database of prophage elements and phage remnants	Srividhya et al., 2007
PHAST	Homology-based tool to annotate prophage sequences	Zhou et al., 2011
Prophinder	Detects genome regions with a significantly high density of known phage-like proteins	Lima-Mendez et al., 2008
	from ACLAME	
Genomic islands		
SIGI-HMM	Discriminative genomic island caller that uses HMMs to identify genome regions with	Waack et al., 2006
	abnormal sequence composition statistics	
IslandViewer	Web-server for predicting genomic islands. Provides access to three different tools	Langille & Brinkman, 2009
PAI-IDA	Combines dinucleotide frequency, G+C content and codon usage to predict the location	Tu & Ding, 2003
	of genomic islands in genomes	
Alien_Hunter	Discriminative genomic island prediction method that uses interpolated variable order	Vernikos & Parkhill, 2006
	motifs to identify genome islands	
IslandPick	One of the few comparative genomics-based genomic island identification tools	Langille et al., 2008
Plasmids, integrative a	Plasmids, integrative and conjugative elements, gene cassettes, integrons	
ICEberg	Web-based resource for integrative and conjugative elements found in bacteria	Bi <i>et al.</i> , 2012
INTEGRALL	Database and search engine for integrons, integrases and gene cassettes.	Moura et al., 2009
ACID	Community resource for annotation of cassette and integron data	Joss et al., 2009
cBar	Distinguishes plasmid-derived from chromosome-derived sequence fragments based on	Zhou <i>et al.</i> , 2010
	their sequence composition	

Origin of replication		
Ori-Finder	Finds origins of replication. Combines evidence from base composition asymmetry,	Gao & Zhang, 2008
	distribution of DnaA boxes and the occurrence of genes frequently close to oriCs to	
	produce predictions	
Sequence database search	rch	
BLAST and PSI-	Set of programs to find similarity between a query protein or DNA sequence and a	Altschul et al., 1997
BLAST	sequence database	
FASTA	Suite of sequence analysis tools for searching Protein or DNA sequence databases. More	Pearson & Lipman, 1988
	accurate than BLAST, but slower to run	
HMMER3	Sequence comparison toolikit implemeting profile hidden Markov models. Accurate and	Eddy, 2011
	fast to run	
Biological databases		
GenBank	One of the largest genetic sequence databases. Maintained by NCBI	Benson et al., 2013
UniProt	UniProt is a catalog of information on proteins. Includes sequence and annotation data	Bairoch et al., 2008
PATRIC	Information system that integrates bacterial genome information with rich data and analysis tools	Gillespie <i>et al.</i> , 2011
CharProtDB	Resource of expertly curated, experimentally characterised proteins described in	Madupu et al., 2012
	published literature	
Rfam	Collection of RNA families	Griffiths-Jones et al., 2003
COG	Database of clusters of orthologous groups of proteins. Orthologue groups build by	Tatusov et al., 1997, Tatusov et al.,
	merging triangles with a common side. Original versions classified sequences into 23	2003
	functional categories	
SEED	Subsystem-based approach and database to high-throughput genome annotation	Overbeek et al., 2005
Protein signature databases	ases	
PFAM	Collection of protein domain families represented by multiple sequence alignments and	Punta et al., 2012
	HMMs	
TigrFAM	Collection of protein families featuring curated multiple sequence alignments, HMMs and	Haft <i>et al.</i> , 2003
	associated information. Most signature models represent full-length proteins	

InterProScan	Tool that combines different protein signature recognition methods into one resource and allows to query different databases at one go	Zdobnov & Apweiler, 2001
Advanced function classification	assification	
BAGEL3	Web-based bacteriocin genome mining tool	van Heel <i>et al.</i> , 2013
RASTA-Bacteria	Bioinformatic tool for identifying toxin-antitoxin loci in bacteria	Sevin & Barloy-Hubler, 2007
TADB	Resource for type 2 toxin-antitoxin loci in prokaryotes	Shao <i>et al.</i> , 2011
ARGO	Database of βlactam and vancomycin resistance genes	Scaria <i>et al.</i> , 2005
MvirDB	Database of protein toxins, virulence factors and antibiotic resistance genes	Zhou <i>et al.</i> , 2007
ARDB	Antibiotic resistance gene database	Liu & Pop, 2009
DBD	Transcription factor database	Wilson et al., 2008
antiSMASH	Software pipeline for secondary metabolite biosynthesis gene cluster identification	Medema <i>et al.</i> , 2011
Mist2	Microbial signal transduction database	Ulrich & Zhulin, 2010
LOCP	Tool for locating pilus operons in gram-positive bacteria	Study I
Metabolism related genes	enes	
PRIAM	Method for automated enzyme detection. Relies on sets of position-specific score	Claudel-Renard et al., 2003
	matrices tailored for each enzyme class	
KAAS	Annotation server that provides functional annotation of CDSs by BLAST comparisons	Moriya <i>et al.</i> , 2007
	against the known KEGG genes. Constructs also KEGG pathway mappings	
TCDB	Database and classification system for membrane transport proteins	Saier <i>et al.</i> , 2006
CAZY	Database of enzymes that degrade, modify, or create glycosidic bonds	Cantarel et al., 2009
MEROPS	Database of peptidase and proteins that inhibit them. Proteins classified into families	Rawlings <i>et al.</i> , 2004
	and homology clans	
REBASE	Restriction enzyme database	Roberts et al., 2010
Context-based protein function prediction	n function prediction	
ContextMirror	Method to find co-evoluting genes. Builds upon family tree similarities	Juan <i>et al.</i> , 2008
String	Database of known and predicted protein-protein interactions. Makes use of homology	von Mering et al., 2005
	information and information on genomic context, high-throughput experiments, co-	
	expression and text-mining	

Prolinks	Database of predicted protein-protein interactions. Interactions dervied from phylogenetic profile, fusion gene, gene neighbor and gene cluster predictions	Bowers <i>et al.</i> , 2004
Ab initio protein function prediction	on prediction	
CSS-Palm	Predicts palmitoylation sites	Ren <i>et al.</i> , 2008
DISIS	Ab initio DNA-binding residue prediction tool that uses multiple classifying algorithms	Ofran <i>et al.</i> , 2007
MetalDetector	Annotates metal binding sites in proteins from sequence information alone	Lippi <i>et al.</i> , 2008
VirulentPred	Virulent protein prediction algorithm that uses SVM	Garg & Gupta, 2008
SPAAN	Neural network prediction of adhesins and adhesin-like proteins	Sachdeva et al., 2005
Subcellular location		
PSORTb v3.0	Subcellular localization prediction tool that uses both SVMs and Bayesian networks	Yu <i>et al.</i> , 2010
tatP	Ppredicts the presence and location of Twin-arginine signal peptide cleavage sites in	Bendtsen <i>et al.</i> , 2005
LipoP	Lipoprotein signal peptide predictor that uses HMMs	Rahman <i>et al.,</i> 2008
SignalP	Signal sequence prediction utility that uses a combination of several artificial neural	Petersen <i>et al.</i> , 2011
	networks to product signal peptide cleavage sites	
ТМНММ	HMM-based protein topology predictor. Identifies transmembrane helices in proteins	Krogh et al., 2001
LocateP	Genome-scale subcellular-location prediction suite. Distinguishes between seven cellular	Zhou <i>et al.</i> , 2008
	locations for gram-positive proteins	
EffectiveT3	Sequence-based prediction of type III secreted proteins	Arnold et al., 2009
CoBaltDB	Complete prokaryote protein subcellular localization database and associated resources.	Goudenège <i>et al.</i> , 2010
	The database integrates the results of 43 localization predictors for over 700 complete	
	prokaryotes proteomes	
Orthology prediction		
RBH	Reciprocal best hit approach. Find two-way best hits among genes in two organisms	Tatusov et al., 1996
RSD	Reciprocal smallest distance algorithm. Uses global sequence alignment and maximum	Wall et al., 2003
	likelihood estimation of evolutionary distances to predict orthologs	
RIO	Mehtod for automated phylogenomics using explicit phylogenetic inference	Zmasek & Eddy, 2002
OrthoStrapper	Program for predicting orthologs between two species. Calculates orthology support	Storm & Sonnhammer, 2002

IMG	Annotation pipelines	KEGG	UniPathway	MetaCyc	FMM	Pathway tools	Metabolic reconstruction	riiyiip	Phylin	BioNJ	PhyML	Phylogenetic trees	SiPhy	GERP	Prediction of constrained elements		BLASTatlas		Gepard		Mauve aligner	
System to support the annotation and analysis microbial genome datasets. Genomes are associated with rich annotation data. Various visualization tools		Database of metabolic pathways and enzymes	Manually curated resource of enzyme-catalyzed and spontaneous chemical reactions and pathways	Database of metabolic pathways and enzymes. Pathways in MetaCyc are more compact and contain on average fewer reactions than those of KEGG	Web-resource for reconstructing metabolic pathways form one metabolite to the other one by combining KEGG maps	Utility for capturing and integrating metabolic information. Uses MetaCyc pathways	SULT STATES AND	sequences	d evolutionary relationshins	Neighbor-joining algorithm to estimate phylogenies	Method to estimate maximum-likelihood phylogenies		Software package for detecing bases under selection from a multiple alignment data	Identifies constrained elements in multiple sequence alignments	ed elements	highlighting conserved and unconserved genome areas in the reference genome	Maps of genome homology of a list of sequences against a reference genome. Good in	and rerarrangements between two sequences	Rapid and sensitive dotplot tool. Dotplots are useful in detecting inversions, duplications	insertions. No support for duplications. Mauve contig mover is a popular scaffolder	Constructs multiple genome alignments in the presence of rearrangements and	require a reference genome, but needs a guide tree
Markowitz <i>et al.</i> , 2012		Kanehisa <i>et al.</i> , 2004	Morgat <i>et al.</i> , 2012	Krieger <i>et al.</i> , 2004	Chou et al., 2009	Karp <i>et al.</i> , 2002		ו פוזפווזרפווו, דסמס	Felsenstein, 1989	Gascuel, 1997	Guindon et al., 2003		Garber et al., 2009	Cooper et al., 2005			Wassenaar et al., 2010		Krumsiek <i>et al.</i> , 2007		Rissman <i>et al.</i> , 2009	

RAST	Server for high quality genome annotation of prokaryotes. Includes a component that	Aziz et al., 2008
	can be used to produce a draft metabolic mode	
DOE-JGI MAP	Microbial annotation pipeline	Mavromatis <i>et al.</i> , 2009
CG pipeline	Resource for assembling sequence data and running feature prediction and annotation	Kislyuk <i>et al.</i> , 2010
	tools on the assembly	
ERGO	Commercial microbial genome analysis and discovery system	Overbeek et al., 2003
PGAAP	NCBI prokaryotic genome annotation pipeline	
Broad Institute	Broad prokaryotic genome annotation pipeline	1
BCM	Baylor Prokaryotic Annotation Pipeline	
JCVI	The automated prokaryotic annotation pipeline of TIGR/JCVI institute	Tanenbaum <i>et al.</i> , 2010
Visualization		
Genome atlas	Circular plots of chromosomes or plasmids on which general properties of the DNA	Wassenaar <i>et al.</i> , 2010
	molecule are plotted as colors	
ACT	Genome browser for feature viewing and annotation. Can display pairwise comparisons	Carver et al., 2005
	between two or more DNA sequences	
Combo	Argo comparative genome viewer	Engels <i>et al.</i> , 2006
Circoletto	Tool for the generation of circularly composited renditions of genomic data and sequence	Darzentas, 2010
	similarity	

Nelson <i>et al.</i> , 2010		58	189	3042	Draft	2.9	20.02.2009	Dairy product	L. cas	L. paracasei ATCC 25302
Nelson <i>et al.</i> , 2010	0	49	140	3021	Draft	3.0	26.08.2008	Human GIT	L. cas	L. paracasei 8700:2
Zhang <i>et al.</i> , 2010	ч	74	64	2848	Complete	2.9	12.07.2010	Koumiss	L. cas	L. casei Zhang
								product		
Chen <i>et al.</i> , 2011	л	73	55	3164	Complete	3.1	01.04.2011	Mongolian dairy	L. cas	L. casei LC2W
Cai <i>et al.</i> , 2009	0	0	83	3044	Complete	3.1	19.06.2008	Laboratory strain	L. cas	L. casei BL23
Ai <i>et al.</i> , 2011	ы	74	42	3204	Complete	3.1	01.04.2011	Koumiss	L. cas	L. casei BD-II
Makarova <i>et al.</i> , 2006	ы	75	72	2771	Complete	2.9	13.10.2006	Swiss cheese	L. cas	L. casei ATCC 334
								cavity		
Nelson <i>et al.</i> , 2010	0	44	506	3183	Draft	2.9	16.12.2011	Human oral	L. buc	L. parafarraginis F0439
								cavity		
Nelson <i>et al.</i> , 2010	1	46	574	3325	Draft	3.0	10.01.2012	Human oral	L. buc	L. kisonensis F0435
Nelson <i>et al.</i> , 2010		60	190	2791	Draft	2.6	19.02.2009	Wine	L. buc	L. hilgardii ATCC 8290
								production plant		
Liu <i>et al.</i> , 2011	л	78	134	2392	Complete	2.6	18.04.2011	Ethanol	L. buc	L. buchneri NRRL B-30929
								cavity		
Nelson <i>et al.</i> , 2010	ц	60	228	3002	Draft	2.9	12.02.2009	Human oral	L. buc	L. buchneri ATCC 11577
Makarova <i>et al.</i> , 2006	ч	81	235	2218	Complete	2.3	13.10.2006	Silage	L. bre	L. brevis ATCC 367
<i>et al.</i> , 2010										
Nelson	1	61	254	3041	Draft	3.1	12.02.2009	Wine	L. bre	L. brevis ATCC 27305
Kim <i>et al.</i> , 2011a	2	55	290	2354	Draft	2.4	21.06.2011	Fermented salami	L. ali	L. versmoldensis KCTC 3814
Nam <i>et al.</i> , 2011b	н	57	349	2440	Draft	2.5	17.12.2010	Sausage	L. ali	L. farciminis KCTC 3681
Original reference	16S	genes	groups	CDSs	Status	(Mb)	Date	Source	Clade	Organism
		RNA	Ortholog			Length	Release			

Appendix Table 2. Characteristics of the sequenced *Lactobacillus* and their genomes.

van de Guchte <i>et al.</i> , 2006	9	122	67	1562	Complete	1.9	26.05.2006	Bulgarian yogurt	L. del	L. delbrueckii ATCC 11842
Zheng <i>et al.</i> , 2008	9	115	41	1792	Complete	1.9	03.03.2011	Dairy product	L. del	L. delbrueckii 2038
Study IV	4	76	72	2024	Draft	2.0	21.04.2010	Chicken crop	L. del	L. crispatus ST1
Nelson <i>et al.</i> , 2010	ω	71	146	2174	Draft	2.1	08.07.2010	Human vagina	L. del	L. crispatus SJ-3C-US
Nelson <i>et al.</i> , 2010	0	57	46	2330	Draft	2.4	01.10.2009	Human vagina	L. del	L. crispatus MV-3A-US
Nelson <i>et al.</i> , 2010	0	62	22	2151	Draft	2.3	24.08.2009	Human vagina	L. del	L. crispatus MV-1A-US
Nelson <i>et al.</i> , 2010	4	66	63	2209	Draft	2.1	27.04.2009	Human vagina	L. del	L. crispatus JV-V01
Nelson <i>et al.</i> , 2010	щ	51	115	2248	Draft	2.4	29.10.2010	Human vagina	L. del	L. crispatus CTV-05
Nelson <i>et al.</i> , 2010	щ	55	50	2163	Draft	2.1	11.03.2010	Human vagina	L. del	L. crispatus 214-1
Nelson <i>et al.</i> , 2010	0	57	24	2082	Draft	2.3	24.08.2009	Human vagina	L. del	L. crispatus 125-2-CHN
Kant <i>et al.</i> , 2011b	4	74	57	1920	Complete	2.0	29.03.2011	Porcine GIT	L. del	L. amylovorus GRL1118
Kant <i>et al.</i> , 2011c	4	72	121	2121	Complete	2.1	19.11.2010	Porcine feces	L. del	L. amylovorus GRL1112
								wort		
Nelson <i>et al.</i> , 2010	ц	58	182	1684	Draft	1.5	23.04.2010	Acidified beer	L. del	L. amylolyticus DSM 11664
Altermann <i>et al.</i> , 2005	4	74	29	1862	Complete	2.0	27.01.2005	Human faeces	L. del	L. acidophilus NCFM
Nelson <i>et al.</i> , 2010	щ	61	79	2020	Draft	2.0	10.03.2009	1	L. del	L. acidophilus ATCC 4796
Oh <i>et al.</i> , 2011	4	75	72	2059	Complete	2.1	07.03.2011	Swine GIT	L. del	L. acidophilus 30SC
Kim <i>et al.</i> , 2011b	ω	57	281	2958	Draft	3.1	21.06.2011	Corn steep liquor	L. cas	L. zeae KCTC 3804
								culture		
Tompkins <i>et al.</i> , 2012	ω	63	15	2719	Draft	2.9	18.11.2011	Dairy starter	L. cas	L. rhamnosus R0011
Prajapati <i>et al.</i> , 2012	ω	53	942	3255	Draft	2.5	07.04.2011	Human faecal	L. cas	L. rhamnosus MTCC 5462
Nelson <i>et al.</i> , 2010	щ	51	102	3155	Draft	3.1	16.03.2009	Human GIT	L. cas	L. rhamnosus LMS2-1
Study III	თ	76	44	2992	Complete	3.0	02.09.2009	Starter culture	L. cas	L. rhamnosus LC705
·	щ	53	69	2758	Draft	2.9	11.09.2008	Cheese	L. cas	L. rhamnosus HN001
Morita <i>et al.</i> , 2009	ы	71	20	2834	Complete	3.0	25.09.2009	Human faeces	L. cas	L. rhamnosus GG
Study III	ы	72	39	2944	Complete	3.0	02.09.2009	Human faeces	L. cas	L. rhamnosus GG
Pittet <i>et al.</i> , 2012	თ	75	65	2887	Complete	3.0	03.11.2011	,	L. cas	L. rhamnosus ATCC 8530
Nelson <i>et al.</i> , 2010	4	46	165	3014	Draft	2.9	18.11.2011	Human faeces	L. cas	L. rhamnosus ATCC 21052

Nelson <i>et al.</i> , 2010	1	51	26	1361	Draft	1.3	05.10.2010	Human vagina	L. del	L. iners LactinV 09V1-c
Nelson <i>et al.</i> , 2010	4	51	84	1459	Draft	1.3	05.10.2010	Human vagina	L. del	L. iners LactinV 03V1-b
Nelson <i>et al.</i> , 2010		50	108	1527	Draft	1.3	05.10.2010	Human vagina	L. del	L. iners LactinV 01V1-a
Nelson <i>et al.</i> , 2010		47	9	1214	Draft	1.3	27.04.2009	Human urine	L. del	L. iners DSM 13335
Nelson <i>et al.</i> , 2010	ч	49	11	1144	Draft	1.2	27.12.2010	Human	L. del	L. iners ATCC 55195
Macklaim <i>et al.</i> , 2011	ഗ	49	13	1209	Draft	1.3	02.02.2010	Human vagina	L. del	L. iners AB-1
Prajapati <i>et al.</i> , 2011	2	68	421	2239	Draft	2.0	07.04.2011	Human vagina	L. del	L. helveticus MTCC 5463
		:	ġ			i		fermented milk	[[[
Zhao et al 2011	4 4	74	л Т Т Т	1978	Complete	2 2 2 2	16 02 2011	Traditional		L helveticus H10
Nelson <i>et al</i> 2010	-	48	144	2078	Draft	-1 20	27.04.2009	Cheese	l del	1 helveticus DSM 20075
Callanan <i>et al.</i> , 2008	4	73	13	1610	Complete	2.1	15.11.2007	Cheese	L. del	L. helveticus DPC 4571
Nelson <i>et al.</i> , 2010	ч	74	28	1955	Draft	2.0	08.07.2010	Human vagina	L. del	L. gasseri SV-16A-US
Nelson <i>et al.</i> , 2010	2	66	ω	1688	Draft	1.8	08.07.2010	Human vagina	L. del	L. gasseri SJ-9E-US
Nelson <i>et al.</i> , 2010		39	67	1945	Draft	1.9	14.10.2008	Human vagina	L. del	L. gasseri MV-22
Nelson <i>et al.</i> , 2010	4	55	82	1977	Draft	2.0	19.02.2009	Human vagina	L. del	L. gasseri JV-V03
Makarova <i>et al.</i> , 2006	6	97	24	1755	Complete	1.9	13.10.2006	Human	L. del	L. gasseri ATCC 33323
Nelson <i>et al.</i> , 2010	ы	97	160	2252	Draft	2.0	30.12.2009	Human vagina	L. del	L. gasseri 224-1
Nelson <i>et al.</i> , 2010		45	21	1773	Draft	1.8	03.06.2009	Human vagina	L. del	L. gasseri 202-4
										PB2003/044-T3-4
Nelson <i>et al.</i> , 2010	2	69	77	1909	Draft	2.0	13.07.2010	Human vagina	L. del	L. delbrueckii
					-			milk		
Sun <i>et al.,</i> 2011	9	121	130	2018	Complete	2.1	19.11.2010	Fermented vak	L. del	L. delbrueckii ND02
Nelson <i>et al.</i> , 2010	щ	74	155	2006	Draft	1.9	10.03.2011	Cheese	L. del	L. delbrueckii DSM 20072
								culture		
McNulty et al., 2011	ц	69	54	1850	Draft	1.8	07.09.2011	Yogurt starter	L. del	L. delbrueckii CNCM I-1632
								culture		
McNulty et al., 2011	1	64	62	1893	Draft	1.8	23.09.2011	Yogurt starter	L. del	L. delbrueckii CNCM I-1519
Makarova <i>et al.</i> , 2006	9	126	87	1721	Complete	1.9	13.10.2006	Starter culture	L. del	L. delbrueckii ATCC BAA-365

Abriouel <i>et al.</i> , 2011	1	64	34	2755	Draft	3.8	23.05.2011	Fermented green	L. pla	L. pentosus MP-10
Kim <i>et al.</i> , 2011c	Ν	65	196	1968	Draft	2.0	22.06.2011	Beer	L. mal	L. malefermentans KCTC 3548
Vogel <i>et al.</i> , 2011	7	82	127	1284	Complete	1.4	06.09.2011	Sourdough	L. fru	<i>L. sanfranciscensis</i> TMW 1.1304
Nam <i>et al.</i> , 2012	1	39	163	1358	Draft	1.4	05.01.2011	Spoiled sake	L. fru	L. fructivorans KCTC 3543
Nelson <i>et al.</i> , 2010	ц	58	201	2210	Draft	2.2	19.02.2009	Human GIT	L. del	L. ultunensis DSM 16047
Wang <i>et al.</i> , 2011a	0	0	249	2162	Complete	2.4	25.05.2011	Kefir grain	L. del	L. kefiranofaciens ZW3
Lee <i>et al.</i> , 2011a	ω	43	113	1846	Draft	1.9	28.06.2011	Piglet feces	L. del	L. johnsonii PF01
Pridmore <i>et al.</i> , 2004	6	97	27	1821	Complete	2.0	02.02.2004	Human GIT	L. del	L. johnsonii NCC 533
Wegmann <i>et al.</i> , 2009	4	67	57	1735	Complete	1.8	04.11.2009	Poultry GIT	L. del	L. johnsonii F19785
Guinane <i>et al.</i> , 2011	4	89	17	1772	Complete	2.0	20.04.2011	Porcine GIT	L. del	L. johnsonii DPC 6026
Nelson <i>et al.</i> , 2010	ц	55	96	1838	Draft	1.8	19.02.2009	Human blood	L. del	L. johnsonii ATCC 33200
Nelson <i>et al.</i> , 2010	0	49	62	1630	Draft	1.7	01.10.2009	Human vagina	L. del	L. jensenii SJ-7A-US
Nelson <i>et al.</i> , 2010	Ν	63	37	1450	Draft	1.6	19.02.2009	Human vagina	L. del	L. jensenii JV-V16
Nelson <i>et al.</i> , 2010	0	50	11	1476	Draft	1.7	24.08.2009	Human vagina	L. del	L. jensenii 27-2-CHN
Nelson <i>et al.</i> , 2010	щ	46	21	1575	Draft	1.7	03.06.2009	Human vagina	L. del	L. jensenii 269-3
Nelson <i>et al.</i> , 2010	Ν	60	43	1347	Draft	1.7	14.10.2008	Human vagina	L. del	L. jensenii 1153
Nelson <i>et al.</i> , 2010	0	50	ω	1470	Draft	1.6	01.10.2009	Human vagina	L. del	L. jensenii 115-3-CHN
Nelson <i>et al.</i> , 2010	ц	51	23	1276	Draft	1.3	07.03.2011	Human vagina	L. del	L. iners UPII 60-B
Nelson <i>et al.</i> , 2010	ц	53	17	1186	Draft	1.3	07.03.2011	Human vagina	L. del	L. iners UPII 143-D
Nelson <i>et al.</i> , 2010	1	49	21	1273	Draft	1.3	05.10.2010	Human vagina	L. del	L. iners SPIN 2503V10-D
Nelson <i>et al.</i> , 2010	0	29	27	1238	Draft	1.3	15.04.2011	Human vagina	L. del	L. iners SPIN 1401G
Nelson <i>et al.</i> , 2010	ц	51	ഗ	1210	Draft	1.3	04.11.2010	Human vagina	L. del	L. iners LEAF 3008A-a
Nelson <i>et al.</i> , 2010	ц	49	13	1265	Draft	1.3	04.11.2010	Human vagina	L. del	L. iners LEAF 2062A-h1
Nelson <i>et al.</i> , 2010	4	52	45	1277	Draft	1.4	04.11.2010	Human vagina	L. del	L. iners LEAF 2053A-b
Nelson <i>et al.</i> , 2010	ч	52	24	1256	Draft	1.3	04.11.2010	Human vagina	L. del	L. iners LEAF 2052A-d
Nelson <i>et al.</i> , 2010	ц	51	27	1338	Draft	1.3	05.10.2010	Human vagina	L. del	L. iners LactinV 11V1-d

L. reuteri mlc3	L. reuteri lpuph	L. reuteri JCM 1112	L. reuteri DSM 20016	L. reuteri CF48-3A	L. reuteri ATCC 53608	L. reuteri 100-23	L. oris PB013-T2-3		L. oris F0423	L. mucosae LM1	L. gastricus PS3		L. fermentum IFO 3956	L. fermentum CECT 5716	L. fermentum ATCC 14931	L. fermentum 28-3-CHN	L. coryniformis KCTC 3535	L. coryniformis KCTC 3167	L. coleohominis 101-4-CHN	L. antri DSM 16041		L. plantarum WCFS1	L. plantarum ST-III	L. plantarum NC8	L. plantarum JDM1	L. plantarum ATCC 14917	
L. reu	L. reu	L. reu	L. reu	L. reu	L. reu	L. reu	L. reu		L. reu	L. reu	L. reu		L. reu	L. reu	L. reu	L. reu	L. reu	L. reu	L. reu	L. reu		L. pla	L. pla	L. pla	L. pla	L. pla	
Mouse	Mouse	Human faeces	Human faeces	Human faeces	Swine GIT	Rat GIT	Human vagina	cavity	Human oral		Human milk	material	Fermented plant	Human Milk	Fermented beets	Human vagina	Kimchi	Silage	Human vagina	Human GIT	cavity	Human oral	Kimchi	Grass silage		Pickled cabbage	olives
13.07.2010	13.07.2010	15.04.2008	01.06.2007	11.03.2009	24.03.2011	25.07.2008	04.11.2010		25.07.2011	21.02.2012	16.02.2012		15.04.2008	29.06.2010	12.02.2009	01.10.2009	17.12.2010	17.11.2010	24.08.2009	27.04.2009		05.02.2003	01.10.2010	16.02.2012	17.07.2009	20.02.2009	
2.0	2.1	2.0	2.0	2.0	2.0	2.3	2.1		2.2	2.2	1.9		2.1	2.1	1.8	2.0	2.8	2.7	1.7	2.2		3.3	ω .3	3.2	3.2	3.2	
Draft	Draft	Complete	Complete	Draft	Draft	Draft	Draft		Draft	Draft	Draft		Complete	Complete	Draft	Draft	Draft	Draft	Draft	Draft		Complete	Complete	Draft	Complete	Draft	
1962	2008	1820	1900	2164	1931	2181	2038		2050	2039	1269		1843	1051	1866	1880	2818	2678	1652	2224		3108	3038	2868	2948	3154	
71	95	ω	4	63	69	157	69		55	320	11		34	15	93	40	216	152	131	183		83	84	27	63	153	
77	82	81	98	55	76	88	62		84	58	43		69	74	60	53	54	35 5	57	59		85	79	75	78	62	
9	4	6	6	2	ω	6	щ		ы	щ	щ		თ	თ	щ	0	щ	щ	0	щ		ഗ	თ	ഗ	თ	щ	
Frese <i>et al.</i> , 2011	Frese <i>et al.</i> , 2011	Morita <i>et al.</i> , 2008	Frese <i>et al.</i> , 2011	Nelson <i>et al.</i> , 2010	Heavens <i>et al.</i> , 2011	Frese <i>et al.</i> , 2011	Nelson <i>et al.</i> , 2010		Nelson <i>et al.</i> , 2010	Lee <i>et al.</i> , 2012			Morita <i>et al.</i> , 2008	Jiménez <i>et al.</i> , 2010a	Nelson <i>et al.</i> , 2010	Nelson <i>et al.</i> , 2010		Nam <i>et al.</i> , 2011a	Nelson <i>et al.</i> , 2010	Nelson <i>et al.</i> , 2010		Kleerebezem <i>et al.</i> , 2003	Wang <i>et al.</i> , 2011b	Axelsson <i>et al.</i> 2012	Zhang <i>et al.</i> , 2009	Nelson <i>et al.</i> , 2010	

L. reuteri MM2-3	L. reu	Human Milk	20.04.2009	1.9	Draft	2045	54	56		Nelson <i>et al.</i> , 2010
L. reuteri MM4-1A	L. reu	Human Milk	19.02.2009	2.1	Draft	2095	18	111	6	Nelson <i>et al.</i> , 2010
L. reuteri SD2112	L. reu	Human Milk	20.06.2011	2.3	Complete	2300	43	88	6	Nelson <i>et al.</i> , 2010
L. vaginalis ATCC 49540	L. reu	Human vagina	19.02.2009	1.8	Draft	1870	196	62	ω	Nelson <i>et al.</i> , 2010
L. curvatus CRL 705	L. sak	Fermented	26.10.2011	1.8	Draft	1862	145	56	ц	Hebert <i>et al.</i> , 2012
		sausage								
L. sakei 23K	L. sak	French sausage	02.11.2005	1.9	Complete	1885	173	84	7	Chaillou <i>et al.</i> , 2005
L. acidipiscis KCTC 13900	L. sal	Cheese	21.06.2011	2.3	Draft	2262	386	55	ω	1
L. animalis KCTC 3501	L. sal	Kimchi	09.12.2010	1.9	Draft	1823	223	29	щ	Nam <i>et al.</i> , 2011c
L. mali KCTC 3596	L. sal	Apple juice	21.06.2011	2.7	Draft	2642	440	58		Kim <i>et al.</i> , 2011d
L. ruminis ATCC 25644	L. sal	Human faeces	19.02.2009	2.1	Draft	2251	292	59	щ	Forde <i>et al.</i> , 2011
L. ruminis SPM0211	L. sal	Human faeces	09.06.2011	2.2	Draft	2326	369	62	щ	Lee <i>et al.</i> , 2011b
L. salivarius	L. sal	Human vagina	19.07.2010	2.0	Draft	2121	108	58	щ	Nelson <i>et al.</i> , 2010
ACS-116-V-Col5a										
L. salivarius ATCC 11741	L. sal	Human oral	19.02.2009	2.0	Draft	1976	134	66	2	Nelson <i>et al.</i> , 2010
		cavity								
L. salivarius CECT 5713	L. sal	Human Milk	07.07.2010	2.1	Complete	1552	17	120	7	Jiménez <i>et al.</i> , 2010b
L. salivarius GJ-24	L. sal	Human faeces	09.06.2011	2.0	Draft	1876	84	83	4	Cho <i>et al.</i> , 2011
L. salivarius NIAS840	L. sal	Chicken faeces	31.05.2011	2.0	Draft	1869	136	103	7	Ham <i>et al.</i> , 2011
L. salivarius SMXD51	L. sal	Chicken GIT	14.03.2012	2.0	Draft	1771	57	102	∞	Kergourlay <i>et al.</i> , 2012
L. salivarius UCC118	L. sal	Human GIT	30.03.2006	2.1	Complete	2014	69	66	7	Claesson <i>et al.</i> , 2006
L. suebicus KCTC 3549	L. vac	Apple mash	21.06.2011	2.7	Draft	2534	368	58	щ	Nam <i>et al.</i> , 2011d
L. sp. 7_1_47FAA	ı	Human GIT	03.10.2011	1.3	Draft	1181	20	76	ч	Nelson <i>et al.</i> , 2010

L. rhamnosus ATCC 8530		L. rhamnosus ATCC 21052	L. paracasei ATCC 25302	L. paracasei 8700:2	<i>L. casei</i> Zhang	<i>L. casei</i> LC2W	L. casei BL23	<i>L. casei</i> BD-II	L. casei ATCC 334		L. parafarraginis F0439		L. kisonensis F0435	L. hilgardii ATCC 8290	L. buchneri NRRL B-30929	L. buchneri ATCC 11577	L. brevis ATCC 367	L. brevis ATCC 27305		L. versmoldensis KCTC 3814		L. farciminis KCTC 3681	Organism
454/Sanger		Illumina	454/Illumina/Sanger	454		454/Illumina/Sanger	454	454/Illumina/Sanger	Sanger		Illumina		Illumina	454/Illumina/Sanger	454/Illumina/Sanger	454/Illumina/Sanger	Sanger	454/Illumina/Sanger		454		454	Sequencing platform
Newbler		Velvet	Newbler		Phrap		Phrap	Newbler	Jazz		Velvet		Velvet	Newbler	Newbler	Newbler	Jazz	Newbler		Newbler		Newbler	Assembler
IGS Annotation Engine	tRNAscan-SE/RNAmmer/Rfam	GeneMark/Glimmer/BLAST/	BCM	Broad Institute			AGMIAL		GeneMarkS/tRNAscan-SE	tRNAscan-SE/RNAmmer/Rfam	GeneMark/Glimmer/BLAST/	tRNAscan-SE/RNAmmer/Rfam	GeneMark/Glimmer/BLAST/	BCM	IMG	BCM	GeneMarkS/tRNAscan-SE	BCM	RNAmmer	RAST/Glimmer/BLAST/	tRNAscan-SE	RAST/Glimmer/RNAmmer/	Gene calling
IGS Annotation Engine		BLAST/BER/KEGG	JCVI	JCVI			AGMIAL		COG/PSI-BLAST		BLAST/BER/KEGG		BLAST/BER/KEGG	JCVI	IMG	JCVI	COG/PSI-BLAST	JCVI		BLAST		RAST/BLAST/COG	Functional Annotation
CP003094		AFZY00000000	ACGY0100000	NZ_DS990485	CP001084	CP002616	FM177140	CP002618	CP000423		AGEY0100000		AGRJ0100000	ACGP01000000	CP002652	ACGH01000000	CP000416	ACGG01000000		BACR01000000		AEOT01000000	ACC

Appendix Table 3. Summary of sequencing status of *Lactobacillus* genome projects.

JCVI
RAST/GeneMark/Glimmer/ tRNAscan-SE
tRNAscan-SE
Gamola/ERGO/Glimmer/
GeneMark/Glimmer/Metagene
GeneMark/Glimmer/Metagene
Broad Institute
GeneMarkS/tRNAscan-SE
GeneMarkS/tRNAscan-SE
AGMIAL/ERGO
SE/RNAmmer
Glimmer/tRNA-scan-

RAST/Glimmer RAST
FrameD/BLAST/tRNAscan-SE
A/ GenDB/Glimmer/tRNAscan- SE
RBSfinder/REGANOR5/CRITIC
Gamola/Glimmer/RAST
BCM
Broad Institute
BCM
/ BLAST
GeneMark/Glimmer/Metagene
JCVI
Broad Institute
Broad Institute
JCVI

AICN00000000	·		Newbler	454	L. gastricus PS3
		tRNAscan-SE			
AP008937	BLAST/PFAM/COG	Glimmer/BLAST/EMBOSS/	Phrap	Sanger	L. fermentum IFO 3956
CP002033			Newbler	454	L. fermentum CECT 5716
ACGI0100000	JCVI	BCM	Newbler	454/Illumina/Sanger	L. fermentum ATCC 14931
NZ_GG704699	JCVI	Broad Institute	Newbler	454	L. fermentum 28-3-CHN
AEOS01000000	PGAAP	PGAAP	Newbler	454	L. coryniformis KCTC 3535
		tRNAscan-SE			
AELK0100000	RAST/COG/BLAST	RAST/Glimmer/RNAmmer/	Newbler	454	L. coryniformis KCTC 3167
NZ_GG698802	BLAST/PFAM	GeneMark/Glimmer/Metagene	Newbler	454	L. coleohominis 101-4-CHN
ACLL0100000	JCVI	BCM	Newbler	454/Illumina/Sanger	L. antri DSM 16041
AL935263				Sanger	L. plantarum WCFS1
CP002222			Newbler	454/Illumina/Sanger	L. plantarum ST-III
AGRI0000000	BLAST/InterPro	IGS	Newbler	454/Illumina	L. plantarum NC8
			bler		
CP001617			Phrap/New	454/SOLiD/Sanger	L. plantarum JDM1
ACGZ0200000	JCVI	BCM	Newbler	454	L. plantarum ATCC 14917
FR871817	BLAST/InterPro		Newbler	454	L. pentosus MP-10
		tRNAscan-SE			
BACN0100000	BLAST/COG	RAST/Glimmer/RNAmmer/	Newbler	454	L. malefermentans KCTC 3548
		CRISPRFinder			
CP002461	PEDANT/BLAST	PEDANT/GenMark/Glimmer/	·	454/Sanger	L. sanfranciscensis TMW 1.1304
		tRNAscan-SE			
AEQY0100000	RAST/BLAST/COG	RAST/Glimmer/RNAmmer/	Newbler	454	L. fructivorans KCTC 3543
ACGU0100000	JCVI	BCM	Newbler	454/Sanger	L. ultunensis DSM 16047
CP002764	I	Glimmer/GeneMark	Phrap/New bler	454/Illumina/Sanger	L. kefiranofaciens ZW3

AF0J01000000			CLCbio/Ne	454/Illumina/Sanger	L. ruminis SPM0211
ACGS02000000	JCVI	BCM	Newbler	454	L. ruminis ATCC 25644
		tRNAscan-SE			
BACP01000000	BLAST/COG	RAST/Glimmer/RNAmmer/	Newbler	454	L. mali KCTC 3596
		tRNAscan-SE			
AEOF01000000	BLAST	Glimmer/RNAmmer/RAST/	Newbler	454	L. animalis KCTC 3501
BACS01000000	PGAAP	PGAAP	Newbler	454	L. acidipiscis KCTC 13900
CR936503	AGMIAL/InterPro/PFAM	AGMIAL	Phrap	Sanger	L. sakei 23K
		tRNAscan-SE/RNAmmer			
AGBU01000000	ISGA/RAST	ISGA/RAST/RAST/Glimmer/	Newbler	454	L. curvatus CRL 705
ACGV01000000	JCVI	BCM	Newbler	454/Illumina/Sanger	L. vaginalis ATCC 49540
CP002844	JCVI	BCM	Newbler	454	L. reuteri SD2112
ACGX02000000	JCVI	BCM	Newbler	454	L. reuteri MM4-1A
ACLB0100000	JCVI	BCM	Newbler	454/Sanger	L. reuteri MM2-3
AEAW01000000		1	Newbler	454	L. reuteri mlc3
AEAX01000000			Newbler	454	L. reuteri lpuph
		tRNAscan-SE			
AP007281	BLAST/PFAM/COG	Glimmer/BLAST/Emboss/	Phrap	Sanger	L. reuteri JCM 1112
CP000705	IMG	IMG		454/Sanger	L. reuteri DSM 20016
ACHG01000000	JCVI	BCM	Newbler	454/Illumina/Sanger	L. reuteri CF48-3A
CACS02000000	BLAST/InterPro	Glimmer3/GeneMark	Newbler	454	L. reuteri ATCC 53608
AAPZ02000000	IMG	IMG	ı	454/Sanger	L. reuteri 100-23
AEKL01000000	JCVI	JCVI	Newbler	454	L. oris PB013-T2-3
			mbler		
AFTL0100000	JCVI	JCVI	CeleraAsse	454	L. oris F0423
			CodonCode		
			wbler/		
AICN00000000	RAST/BLAST	RAST	CLCbio/Ne	454/Illumina	L. mucosae LM1

ACWR0100000	JCVI	Prodigal	Newbler	454	L. sp. 7_1_47FAA
		SE/RNAmmerm BLAST			
BACO01000000	BLAST	RAST/Glimmer/tRNAscan-	Newbler	454	L. suebicus KCTC 3549
		/Orpheus/Critica/tRNAscan-SE			
CP000233	ERGO/COG/TCDB	ERGO/YACOP/glimmer/Zcurve	Phrap	Sanger	L. salivarius UCC118
AICL0000000	RAST	RAST/Glimmer	MIRA	454	L. salivarius SMXD51
			wbler		
AFMN01000000	RAST	RAST/BLAST	CLCbio/Ne	454/Illumina	L. salivarius NIAS840
			wbler		
AF0I0100000	RAST/KEGG/COG	RAST	CLCbio/Ne	454/Illumina	L. salivarius GJ-24
CP002034		•	Newbler	454	L. salivarius CECT 5713
ACGT0100000	JCVI	BCM	Newbler	454/Illumina/Sanger	L. salivarius ATCC 11741
AEBA01000000	JCVI	JCVI	Newbler	454	L. salivarius ACS-116-V-Col5a
			wbler		