

Reannotation of the genome sequence of Clostridium difficile strain 630.

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1 Re-annotation of the genome sequence of *Clostridium difficile* strain 630 2 Marc Monot¹, Caroline Boursaux-Eude², Marie Thibonnier¹, David Vallenet³, Ivan 3 Moszer², Claudine Medigue³, Isabelle Martin-Verstraete^{1,4} and Bruno Dupuy¹ 4 ¹ Institut Pasteur, Laboratoire Pathogenèse des Bactéries Anaérobies, 28 rue du Dr Roux, 75724 Paris Cedex 5 15, France 6 ² Institut Pasteur, Intégration et Analyse Génomiques, 28 rue du Dr Roux, 75724 Paris Cedex 15, France 7 ³ CNRS UMR 8030, Université d'Evry & CEA, IG, Genoscope - LABGeM - 2 rue Gaston Crémieux, CP5706, F-8 91057 Evry cedex, France 9 ⁴ Université Paris Diderot-Paris 7 – 7 boulevard Diderot, 75012 Paris, France 10 11 Correspondence 12 Marc Monot 13 mmonot@pasteur.fr 14 Tel. +331 45 68 83 16 / Fax. +331 40 61 31 83 15 A regular update of genome annotations is a prerequisite step to help maintain the 16 17 accuracy and relevance of the information they contain. Five years after the first 18 publication of the complete genome sequence of *C. difficile* strain 630, we manually reannotated each of the coding sequences (CDS), using a high-level annotation platform. 19 20 The function of more than 500 genes annotated previously with putative functions, were 21 re-annotated based on updated sequence similarities of proteins whose functions have 22 been recently identified by experimental data from the literature. We also modified 222 23 CDS starts, detected 127 new CDS and added the enzyme commission numbers, which 24 were not supplemented in the original annotation. In addition, an intensive project was 25 undertaken to standardise the names of genes and gene products and thus harmonising 26 as much as possible with the HAMAP project. The re-annotation is stored in a relational 27 database that will be available on the MicroScope web-based platform, 28 "http://www.genoscope.cns.fr/agc/microscope/ClostridioScope". The original 29 submission stored in the INSDC nucleotide sequence databases was also updated. 30 31 **Scope :** *Clostridium difficile* special issue of the *Journal of Medical Microbiology* 32 *The EMBL accession number for the re-annotation of C. difficile strain 630 is AM180355 33 34 and its plasmid pCD630 is AM180356.

36 INTRODUCTION

37

The re-annotation of several model genomes has been recently performed, among these there are *Escherichia coli* for the gram negative bacteria (Luo *et al.*, 2009) and *Bacillus subtilis* for the firmicutes (Barbe *et al.*, 2009). This provided new information about genomic structure and organisation as well as gene function and plays an essential role in defining reference knowledge. In addition the re-annotation of the *B. subtilis* genome also benefits the other firmicutes such as *Clostridium difficile*.

44

C. difficile is one of the major enteropathogenic clostridia and *C. difficile* associated diarrhea (CDAD) is currently the most frequently occurring nosocomial infection in many European hospitals. Although toxins are generally recognised as the main virulence factors, *C. difficile* pathogenesis remains poorly understood. The global genetic analysis of *C. difficile* appeared to be an useful approach to find potential mechanisms involved in the bacterial virulence for which an updated of the gene list and corresponding annotations is tremendously important.

52

53 The first complete genome sequence of a *C. difficile* strain (630) was sequenced in 54 2006 (Sebaihia *et al.*, 2006). It led to the development of high throughput projects such 55 as comparative genomic, transcriptomic and proteomic studies (Jain *et al.*, 2010; Janvilisri et al., 2010; Marsden et al., 2010), which were recently reinforced with an 56 57 increase of multiple genomic projects (Stabler *et al.*, 2009). However, the relevance of all these experiments greatly depends on the information available for the genes 58 59 particularly their functions experimentally identified or predicted *in silico*. Thus, it is 60 critical that the information is accurate, relevant and useful. This is why we undertook the re-annotation of the *C. difficile* strain 630 genome. 61

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The advances in second-generation sequencing technologies combined with their relative low cost has led to the increased need for a rapid genome annotation system (Petty, 2010). However the fastest way to obtain an accurate annotation remains to transfer annotation from a reference strain. This requires to have access to a closely related genome for each species annotated to a high standard and regularly updated. 69 We described in this paper the manual re-annotation of all CDS of the *C. difficile* 70 strain 630 genome. For this purpose we used improved methods in bioinformatics, 71 literature surveys and genome data from closely related species such as *Clostridium* 72 sticklandii, which has recently been sequenced (Fonknechten et al., 2010) or B. subtilis 73 whose genome has been re-sequenced and re-annotated (Barbe et al., 2009). The re-74 annotation resulted in the function precision of more than 500 genes and the addition of 75 new CDSs as well as the correction of the start sites of 222 CDSs. All information from 76 laboratory research publications could be continuously integrated though the 77 MicroScope platform to maintain this up-to-date annotation.

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79 **METHODS**

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81 <u>Identification of new or modified CDS in the *C. difficile* genome</u>

82 The sequence and the original annotation of the published *C. difficile* 630 genome (Sebaihia et al., 2006) was integrated into the Microscope platform (Vallenet et al., 83 84 2009). MicroScope is a web-based framework for the systematic and efficient revision of microbial genome annotation and comparative analysis. Its main features are (i) 85 integration of annotation data from bacterial genomes enhanced by a gene coding re-86 87 annotation process using accurate gene models, (ii) integration of results obtained with a wide range of bioinformatics methods, among which exploration of gene context by 88 89 searching for conserved synteny and reconstruction of metabolic pathways, (iii) an advanced web interface allowing multiple users to refine the automatic assignment of 90 91 gene product functions. MaGe is also linked to numerous well-known biological databases and systems. The original gene prediction was systematically checked using 92 93 the AMIGene software (Bocs et al., 2003) and the MICHeck strategy (Cruveiller et al., 94 2005). The initial identifier of genes 'CD0000(A)' used a prefix of two letters, 'CD', 95 followed by a four-digit number corresponding to the position of CDS in the genome. Whenever a new gene interleave, a capital letter was added in alphabetical order. Since 96 97 2006, the locus tag usage has evolved (Cochrane *et al.*, 2008). The prefix now has to 98 contain only alpha-numeric characters and it must be at least 3 characters long. In 99 addition the locus tag prefix must be separated from the tag value by an underscore

100 ending with a number. So we assigned for all CDS a new locus tag code: 'CD630_00000'. 101 The four-digit number after the underscore is still the original CDS position in the 102 genome. The capital letter of the original identifier was converted to a number which 103 has been added at the end of each gene : 1 to 9 for genes previously ended with capital 104 letter A to I, and 0 for all others e.g. CD0001 into CD630_00010 and CD0163B into 105 CD630_01632. Finally, because the genomic position of the non coding CDS was defined 106 with only three-digit numbers, we replaced the first number after the locus tag prefix 107 with a 't' or 'r' respectively for transfer RNA and ribosomal RNA respectively, e.g. CDt001 108 into CD630 t0010 and CDr001 into CD630 r0010. We used the same coding method for 109 the 11 CDSs encoded by the plasmid pCD630, adding the letter 'p', after the locus tag e.g. 110 CDP01 into CD630_p010.

111

During the re-annotation process using the AMIGene predictions, we identified new CDS and we assigned them the locus tag of the previous CDS with the last number incremented by 1 e.g. a new gene detected after CD630_02670 (previously named CD0267) was coded CD630_02671. The original locus tag will be kept in the EMBL file using the /Old_locus_tag identifier.

117

118 <u>Re-annotation of the complete *C. difficile* strain 630</u>

119 The predicted proteins were subjected to a wide range of bioinformatics tools, 120 which includes conserved synteny computations, alignments against TrEMBL and 121 SWISS-PROT databases (Apweiler R, 2011) and TMHMM (Sonnhammer et al., 1998), 122 SignalP (Bendtsen et al., 2004) and PsortB (Yu et al., 2010) software to predict 123 subcellular localization of proteins as well as INTERPROSCAN (Zdobnov & Apweiler, 124 2001) to identify possible functions of newly discovered proteins (Apweiler R, 2011). 125 This work flow led to an automatic functional annotation for each CDS as previously 126 described (Vallenet et al., 2006). Finally, these pre-computed results served as basis for 127 the manual re-annotation of each CDS proceeding by inference.

128

To normalise the process of manual annotation among multiple users, we set up several guidelines: (a) The product field is filled with the functional annotation for all genes identified with 'Hypothetical protein' or 'Conserved Hypothetical protein' when the gene was not identified. For all others we added 'Putative' prior to the product 133 annotation. Pseudogene and gene remnant have a specific nomenclature : "Fragment of" 134 + function + position (N-terminal, C-terminal or center of the encoding protein). (b) The 135 name of gene was completed by searching in the literature using PubMed data libraries 136 (http://www.ncbi.nlm.nih.gov/pubmed) and when we changed gene names, old names 137 were indicated in the synonymous field. (c) The start sites were modified according to 138 the combination of the graphical data such as codage probability curves deduced from 139 the AMIGene method (Bocs *et al.*, 2003), as well as alignments with orthologous genes 140 (Altschul *et al.*, 1990). Then, the label '/START=' was added in the comment field 141 followed by a capital letter associated to an informative code (M: modified, C: coding 142 curve, S: sequence similarity, O: overlap, R: RBS). (d) PubMed identifiers (PMIDs) of 143 each gene were classified from the specific references to the articles corresponding to 144 orthologuous genes and/or the global reviews concerning its function. (e) Protein 145 families were standardized using the same keywords, PMIDs and global classification, 146 such as CMR roles (<u>http://cmr.jcvi.org/cgi-bin/CMR/RoleIds.cgi</u>).

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148 **RESULTS & DISCUSSION**

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150 Evaluation of annotation improvement

151 The original annotation of the *C. difficile* strain 630, published in 2006 (Sebaihia 152 et al., 2006), identified 3776 predicted coding sequences (CDSs). We have updated 153 annotation of all CDSs and assigned or precised their functions. During the re-annotation 154 process we attributed a class of function to each gene re-annotated: (i) "known": when 155 function was experimentally demonstrated or when high level of similarities with 156 characterised genes were found (ii) "putative": based on conserved motif, structural 157 feature or limited similarities, (iii) "unknown": when genes were unidentified and (iv) 158 "pseudo": for pseudogenes or gene remnants. The same classification was applied 159 manually to the 2006 annotation to allow comparison of both annotations (Table 1A).

160

161 Thus, 518 and 18 genes whose encoding function was previously described as 162 putative and unknown respectively have now a functional annotation identified by the 163 experimental data from the literature (Table 1A). For example, CD630_26030 164 (previously named CD2603), recognised as a putative response regulator, is now 165 designated cdtR, since it was shown that it controls the binary toxin expression in C. 166 difficile (Carter et al., 2007). In addition, 117 genes of unknown function have now a 167 putative function. For instance, 12 conserved hypothetical proteins which contain a 168 CRISPR-associated domain (clustered regularly interspaced short palindromic repeats) 169 are annotated "Putative CRISPR-associated family protein". Furthermore, we showed 170 that the ATP synthase epsilon chain, CD630_34670 (CD3467), which was defined as a 171 gene remnant (pseudo class) because of a lack of amino-acid in the C-terminus relative 172 to database matches, actually belongs to the class of "known function". This enzyme 173 usually combines ATP synthesis and hydrolysis but the hydrolysis function is still active 174 in the truncated version (Ferguson *et al.*, 2006).

175

Following the re-annotation we included 127 new CDSs and defined 222 new CDS start sites. The majority of the new CDS are divided into putative (25), unknown (86) or pseudogene (15) classes (Table 1B). Only one gene, *CD630_15951* has an orthologue, whose function was experimentally demonstrated. This gene, detected during the proteomic analysis recently performed in *C. difficile* (Lawley *et al.*, 2009), is highly homologous (~60%) to a ferredoxin gene of *Clostridium thermoaceticum (Elliott et al.*, *1982*).

183

184 We were looking for papers corresponding to each gene, and particularly those 185 published after the original annotation. We added at least one PMID reference number 186 to 64% of the C. difficile genes. Like many other genome-wide updates, several 187 specificities were added to the original product function. When possible, we attached 188 new motifs and enzymatic domains identified by INTERPROSCAN, allowing a more 189 accurate description of the original function. For example, putative peptidase enzymes 190 have now family information according to the classification scheme of the MEROPS 191 database (<u>http://merops.sanger.ac.uk</u>). The revised nomenclature of the pathogenicity 192 locus region (Rupnik *et al.*, 2005) has been introduced during re-annotation process as 193 well as genes involved in the *C. difficile* motility and flagellar glycosylation since they 194 were recently published (Twine *et al.*, 2009). A locus tag, product annotation and class 195 comparison between the two annotations performed in 2006 and 2010 were 196 summarised in the Table S1. All information of the CDS re-annotated (Fig. S1), are 197 currently available the MicroScope platform: on

http://www.genoscope.cns.fr/agc/microscope/ClostridioScope. We also updated the *C. difficile* 630 genomic entry in genomic databases: EMBL, GenBank and DDBJ.

200

201 <u>Deciphering the annotation origin</u>

202 Several pieces of information appeared when we evaluated the source used 203 during the functional annotation of known, putative or unknown genes (Fig. 1). In the 204 known category, 1% of the gene function came from *C. difficile* strain 630 publications, 205 1,5% from other C. difficile strains, 4% from other clostridia and 93,5% from other 206 species (Fig. 1A.). The putative category was defined according to the enzymatic domain 207 (40%), homology to mobile elements (20%) or cell localisation (15%) (Fig. 1B). As an 208 example a gene will be annotated "Putative membrane protein" when 3 or more 209 transmembrane helix was detected by TMHMM (Sonnhammer et al., 1998). Finally, we 210 classified the unknown genes from the alignement results with TREMBL (Boeckmann et 211 al., 2003). Although 45% were orphan of the *C. difficile* strains, 20% were also found in 212 the genus Clostridium, 15% in the firmicutes phylum and 20% in other bacteria (Fig. 213 1C).

214

215 Concerning genes annotated as known, we noted that only few of them came 216 from a published clostridial experiments (Fig. 1A). This was mainly due to the lack of 217 effective tools to mutate clostridial genes. However gene inactivation method and 218 random mutagenesis system recently developed in *C. difficile* (Cartman & Minton, 2010), 219 should greatly improve the number of publications on *C. difficile* gene functions. Half of 220 the genes with an unidentified function, orphan, are found only in *C. difficile* 630 (Fig. 221 1C.). However, most orphans are present in the *C. difficile* strains already sequenced 222 such as strains 027, CD196 and R20291 (Stabler et al., 2009). This may constitute a 223 source of gene targets that could be used both in research, diagnosis or treatment of the 224 CDAD.

225

226 <u>Miscellaneous improvements</u>

To re-annotate the *C. difficile* genome of strain 630 we used the MaGe interface, which contains classic database fields (type, position, name, product, EC numbers) and several specific fields such as gene synonymous (synonyms), authors notes (comments), pubmed identifiers (PMID), product type, localisation and functional classification (Fig. S1). All information found during the re-annotation process that did not fit in the classic
fields were added in the specific MaGe field or in the comments. For example, the novel
virulence factor called Srl for « Sensitivity regulation of *C. difficile* toxins », (Miura *et al.*,
2010) was presented during the third international clostridium difficile symposium.
This information was only indicated in the comment field of the CD630_22980 (CD2298)
gene until further validation.

237

238 The names of gene products were harmonizing as much as possible with the 239 HAMAP project (Lima *et al.*, 2009). On the other hand, all gene products have now been 240 named with a specific keyword related to their functional family (Fig. S1). Thus, 241 CD630_05310 (CD0531), previously annotated «DeoR-like regulator of transcription» (a 242 regulator of sugar and nucleoside metabolic systems) was re-annotated « Transcriptional regulator (keyword), DeoR family ». We also normalised the 243 244 annotation of genes that share the same characteristics. As an example, proteins that 245 were only determined according to their membrane localization were annotated: 246 « Putative membrane proteins ». The annotation standardization we used will facilitate 247 the mining of the data using bioinformatics as well as manual search (Fig. S1).

248

249 <u>Membrane Transport</u>

250 The *C. difficile* genome contains a lot of proteins encoding several membrane 251 transport systems: ATP-binding cassette (ABC) transporters, phosphoenolpyruvatedependent phosphotransferase systems (PTS), charged substrate transporters 252 253 (antiporters, symporters) and facilitators. The general function of the genes encoding 254 such proteins can be easily determined from bioinformatic approaches, like those used 255 for the protein domain analysis in InterProScan (Zdobnov & Apweiler, 2001). However 256 it is quite difficult to distinguish the exact metabolite they transport, especially when the 257 transport systems have a wide specificity. We reannotated most of the transporter 258 systems by inference including clues about targets using specialized databases such as 259 TransportDB (<u>http://www.membranetransport.org/</u>)(Ren *et al.*, 2007) which compile 260 all information on cytoplasmic membrane transporters. We added a suffix in the 261 classification which indicate, from a global trend to the expected target, the motif 262 (family), the high sequence homology (like) and the evidence of a target metabolite

(specific). However, this classification should be taken with caution since it was mainlydeduced from *in silico* analysis rather than from experimental data.

265

266 The table 2 showed annotation of 19 PTS systems with a specific 267 metabolite suggestion. The targeted metabolite was deduced from the INTERPROSCAN 268 motif search but could also be defined by the presence in the same locus of gene 269 encoding enzyme involved in specific sugar assimilation (associated enzyme). As an 270 example CD630_22690 (CD2269) is now annotated as "PTS system, fructose-specific 271 IIABC component". This is due to the detection of three motif signatures the mannitol 272 family PTS EII component A, B and C, as well as the presence of the neighbouring gene, 273 the CD630_22700 (CD2270), which encodes an enzyme involved in the utilization of fructose: "Fructose 1-phosphate kinase" as indicated in the gene annotation (Table S1). 274

275

276 <u>Metabolism update</u>

277 Updating the genome annotation of *C. difficile* led to many changes within the 278 metabolism pathways. The gene cluster involved in the anaerobic oxidative degradation 279 of L-ornithine has been identified in *C. sticklandii* (Fonknechten *et al.*, 2009). From this 280 publication we reannotated genes CD630_04420 (CD0442) to CD630_04480 (CD0448) 281 whose encoding proteins share high similarities to the ornithine catabolism compounds of *C. sticklandii* e.g. Ord, OrtA, OrtB, OraS, OraE, Or-4 and Orr, respectively (Table S1). 282 283 This suggested that *C. difficile* could produce acetyl-CoA from the ornithine 284 fermentation. The ability to use a variety of carbohydrates is an important feature for *C*. 285 difficile to colonize the host gut. Enterococcus faecalis found in the same niche as C. 286 *difficile*, provided hints to explore the consistency of a specific pathway required for 287 ethanolamine utilisation, a constituent of an abundant class of phospholipids present in 288 the eucaryotic cell membranes and the host's dietary intake (Del Papa & Perego, 2008) 289 (Fox et al., 2009). Using the E. faecalis gene synteny and protein similarities, we were 290 able to reconstruct the whole ethanolamine pathway in *C. difficile*, a cluster of 19 genes, 291 from CD630_19060 (CD1906) to CD630_19260 (CD1926) encoding the ethanolamine 292 ammonia-lyase, an alcohol dehydrogenase, a carboxysome associated proteins, the 293 transporter EutH and the two-component system EutV, EutW. (Table S1).

Interestingly, in *B. subtilis* several enzymes involved in RNA degradation were recently identified (Even *et al.*, 2005) (Shahbabian *et al.*, 2009). In *C. difficile*, a unique Rnase J protein *CD630_12890 (CD1289)* was detected as well as an ortholog of *ymdA CD630_13290 (CD1329)*, encoding the Rnase Y protein.

299

300 **CONCLUSION**

301

302 Finally, nearly half of the genes of the *C. difficile* strain 630 encode proteins with 303 known function, whereas one-third of the gene products have a putative function and 304 only fifteen percent of proteins with unknown function are encoded by C. difficile 305 genome (Table 1A). In addition, 127 new CDSs were discovered (Table 1B) and 222 CDS 306 starts were modified. The re-annotation was performed using a high standard 307 annotation MicroScope platform, which significantly increased the amount of 308 information available for the majority of the CDS, such as literature references, product 309 types, localisation and gene synonymous (Fig. S1).

310

Nevertheless, there is still great deal of work to be completed since only 116 annotated genes came from a published clostridial experiments. The EMBL entries are now resubmitted and to maintain the annotation up-to-date, all new information would be addressed directly to marc.monot@pasteur.

315

316 **ACKNOWLEDGEMENTS**

317

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325 AUTHORS' CONTRIBUTION

326

BD, MM, CB-E and IM designed the study. CB-E and MM carried out the major part of the
manual re-annotation of the genome together with MT and IM-V. DV and CM were
involved in automatic re-annotation and administration of the MicroScope platform.
MM, IM-V and BD wrote the manuscript.

331

TABLE & FIGURE LEGENDS

333

334 Table 1 : Review of the 2006's annotation update. A) CDSs were identified and separated 335 according to the four major annotation classes in both 2006 and 2010 annotations: 336 (Known) when function were experimentally demonstrated, (Putative) based on 337 conserved motif, structural feature or limited homology, (**Unknown**) when function are 338 unidentified and (**Pseudo**) for pseudogenes. Dark grey padding numbers indicated no change and "±" and "-" correspond to a change betwenn the classes of annotation 339 340 between 2006 annotation and 2010 re-annotation. B) Annotation of the new CDS 341 detected and referenced as known, putative, unknown and pseudo classes.

342

Table 2 : Re-annotation of the PTS systems according to the metabolite specificity. List of
 locus tags corresponding to 19 PTS re-annotated. The PTS metabolite was deduced from
 the motif class detection and/or the presence of associated enzymes involved in a
 specific sugar metabolism.

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Figure 1: Distribution of the functional re-annotation origin. A) Known functions 348 349 were identified from the literature references of: Clostridium difficile strain 630 (C. diff 350 630), other *Clostridium difficile* strains (C. difficile), *Clostridium* species (Clostridia) and 351 others species (Others). B) **Putative** functions were defined from: enzymatic domains 352 (Enzyme), homology with mobile elements (Mobile), localisation in the cell 353 (Localization) and the remaining origin (Others). C) Unknown functions which were 354 found only in: *Clostridium difficile* (Orphan), in the Clostridium species (Clostridia), in the firmicutes plylum (Firmicutes) or in diverse bacteria (Others). 355

356 SUPPLEMENTARY DATA

357

- 358 <u>Table S1 : Comparison between 2006 and 2010 annotations.</u> For each CDS, the locus
- tags, annotation function and classes are compared for both 2006 and 2010 annotations.
- 360

361 <u>Table S2 : Standardization of family product names.</u> The product names were
 362 constructed around a keyword specific to the gene's functional family.

363

364 Figure S1 : *C. difficile* CDS re-annotation by MaGe. MaGe annotation window for *tcdR*

365 gene. Bold and grey outline focus on information added specifically within this process :

366 mutation, synonyms, comments, PMID, product type, localization, MaGe classification

- 367 and standard classification (Bioprocess and Roles).
- 368

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A

2010	Known 47%	Putative 37%	Unknown 14%	Pseudo 2%
Known 34%	1222	-63	-0	-10
Putative 47%	+518	1163	-115	-12
Unknown 17%	+18	+177	397	-7
Pseudo 2%	+1	+2	+1	63

B

New CDS	Known	Putative	Unknown	Pseudo
127	1	25	86	15

<u>Table 2</u>

Locus tag	Motif class	Associated enzyme	Propposed PTS metabolites
CD630_04690	Glucose	CD630_04680	Sucrose
CD630_03880	Glucose	CD630_03890	b-glucoside
CD630_30970	Glucose	CD630_30950 / CD630_30960	b-glucoside
CD630_31160	Glucose	CD630_31150	b-glucoside
CD630_31250	Glucose	CD630_31240	b-glucoside
CD630_31370	Glucose	CD630_31360	b-glucoside
CD630_26660 / CD630_26670	Glucose	-	Glucose
CD630_30580 / CD630_30610	Glucose	CD630_30600	a-glucoside
CD630_22690	Mannitol	CD630_22700	Fructose
CD630_30750	Mannitol	CD630_30740	Tagatose
CD630_30860	Mannitol	CD630_30850	2-0-a-mannosyl-D-glycerate
CD630_23320 / CD630_23330	Mannitol	CD630_23310	Mannitol
CD630_00410 / CD630_00420 / CD630_00430	Mannitol	-	Galactitol
CD630_36450 / CD630_36470 / CD630_36480	Lactose	-	Lichenan
CD630_28800 / CD630_28830 / CD630_28840	Lactose	CD630_28820	Cellobiose
CD630_30130 / CD630_30140 / CD630_3015	Mannose	CD630_30120	Mannose
CD630_25660 / CD630_25670 / CD630_25680	Mannose	CD630_25690	Mannose
CD630_30670 / CD630_30680 / CD630_30690 / CD630_30700	Mannose	CD630_30710	Xyloside
CD630_07640 / CD630_07650 / CD630_07660 / CD630_07670	Sorbitol	CD630_07680	Sorbitol