

Pentjuss, A., Stalidzans, E., Liepins, J., Kokina, A., Martynova, J., Zikmanis, P., Mozga, I., Scherbaka, R., Hartman, H., Poolman, M., Fell, D. and Vigants, A. (2017) 'Model based biotechnological potential analysis of *Kluyveromyces marxianus* central metabolism', *Journal of Industrial Microbiology and Biotechnology*

DOI: <https://doi.org/10.1007/s10295-017-1946-8>

This document is the authors' Accepted Manuscript.

License: <https://creativecommons.org/licenses/by-nc-nd/4.0>

Available from RADAR: <https://radar.brookes.ac.uk/radar/items/73372cc8-727a-4e4c-b9b5-0523edf3ba59/1/>

Copyright © and Moral Rights are retained by the author(s) and/ or other copyright owners unless otherwise waved in a license stated or linked to above. A copy can be downloaded for personal non-commercial research or study, without prior permission or charge. This item cannot be reproduced or quoted extensively from without first obtaining permission in writing from the copyright holder(s). The content must not be changed in any way or sold commercially in any format or medium without the formal permission of the copyright holders.

1 Model based biotechnological potential analysis of *Kluyveromyces marxianus* central metabolism

2
3
4 Pentjuss A.¹, Stalidzans E.¹, Liepins J.¹, Kokina A.¹, Martynova J.¹, Zikmanis P.¹, Mozga I.¹, Scherbaka
5 R.¹, Hartman H.², Poolman M. G.², Fell D. A.², Vigants A.¹

6
7 ¹ Institute of Microbiology and Biotechnology, University of Latvia, Jelgavas str. 1, Riga, LV-1004,
8 Latvia

9 ² Department of Biological and Medical Sciences, Oxford Brookes University, Headington, Oxford
10 OX3 0BP, UK

11
12 Corresponding author: Egils Stalidzans (mail. stalidz@lu.lv, phone: +371 29575510)

13 14 **Abstract**

15
16 The non-conventional yeast *Kluyveromyces marxianus* is an emerging industrial producer for many
17 biotechnological processes. Here we show the application of a biomass-linked stoichiometric model of
18 central metabolism that is experimentally validated, and mass and charge balanced for assessing the
19 carbon conversion efficiency of wild type and modified *K. marxianus*. Pairs of substrates (lactose,
20 glucose, inulin, xylose) and products (ethanol, acetate, lactate, glycerol, ethyl acetate, succinate,
21 glutamate, phenylethanol and phenylalanine) are examined by various modeling and optimisation
22 methods.

23 Our model reveals the organism's potential for industrial application and metabolic engineering.
24 Modeling results imply that the aeration regime can be used as a tool to optimise product yield and flux
25 distribution in *K. marxianus*. Also rebalancing NADH and NADPH utilisation can be used to improve
26 the efficiency of substrate conversion. Xylose is identified as a biotechnologically promising substrate
27 for *K. marxianus*.

28
29 **Keywords** *Kluyveromyces marxianus*. Modelling. Central metabolism. Metabolic engineering.
30 Essentiality analysis.

31 **Introduction**

32

33 *Kluyveromyces marxianus* is an ascomycotous yeast with enormous biotechnological potential for
34 multiple industrial applications. There are a number of characteristics of *K. marxianus* that are
35 industrially useful, including: fast growth, broad substrate spectrum, thermotolerance, limited
36 fermentation at sugar excess, and secretion of extracellular glycolytic enzymes. In addition, *K.*
37 *marxianus* enjoys GRAS (Generally Regarded as Safe) status and therefore is useful in food or pharma
38 related applications [30, 46].

39

40 *K. marxianus* can grow on glucose, fructose, xylose, galactose, lactose and inulin as the sole carbon
41 sources [24]. Many of these carbon sources are of particular interest since they are waste products of
42 forestry (xylose) or dairy (lactose) industries. Xylose is a pentose and the main sugar of plant
43 hemicellulose; its content in hard wood wastes can be up to 30 % [50]. *K. marxianus* has been
44 engineered for xylitol production from xylose [42]. Cheese whey is a lactose rich by-product of the
45 dairy industry produced in an approximate 10 to 1 (v/w) ratio to cheese. Currently whey is considered
46 as a potential substrate for future microbial fermentations [25, 65]. Inulin is one of the widely available
47 plant polysaccharides common in many taxonomic groups (*Asteracea* family, wheat, onion, banana,
48 etc.). Some of those (e.g. Jerusalem artichoke, chicory) accumulate inulin in their underground tubers
49 in vast amounts [11, 18]. These plants might serve as “niche” substrates for fermentations by yeasts
50 including *K. marxianus*, [11] if not deprecated on account of competition with food use.

51

52 *K. marxianus* is a prospective producer for a range of important food additives and chemicals:
53 phenylethanol, phenylalanine [60], hexanoic acid [10], xylitol [107, 108] and ethylacetate [52]. Due to
54 its protein excretion, *K. marxianus* is suitable for extracellular protein production (galactosidase,
55 inulinase, etc.) [26, 98].

56

57 Stoichiometric models and reconstructions significantly facilitate analysis of metabolic effects and
58 limitations of microorganism metabolism, as well as predicting the phenotype of recombinant strains
59 [39, 40]. Modeling attempts on *K. marxianus* to date have been concentrated on particular problems:

60 e.g. kinetic models of ethanol batch fermentation [77], and of growth on cheese whey [51]. The first
61 attempt at a genome scale metabolic reconstruction [47] is patented in unreadable form and cannot be
62 used for metabolic flux calculations. A genome-scale metabolic model for the related species *K. lactis*
63 has been published [16]. Analysing medium scale stoichiometric models of central metabolism, where
64 the most significant metabolic fluxes are, has been successful for biotechnological applications.
65 Examples include assessment and selection of productive routes in *Escherichia coli* [88–90] and
66 *Zymomonas mobilis* [68]. Medium scale modelling also proved to be a successful strategy for
67 describing the uncharacterised central metabolism of the non-conventional yeast *Pichia pastoris* on the
68 basis of limited wet experimentation [87]. Recent extensive attempts at *K. marxianus* metabolic
69 engineering [33, 42, 43, 105, 109] underline the immediate need for modelling of limits of its
70 metabolic potential.

71

72 The aim of this study was to assess the biotechnological potential of *K. marxianus* by a constraint
73 based stoichiometric [79] modeling approach. A biomass-coupled model of central metabolism was
74 developed to be a basis for design of metabolic engineering and to assess *in silico* the production of
75 ethanol, acetate, lactate, glycerol, ethylacetate, succinate, glutamate phenylethanol, phenylalanine. As
76 well as being useful products in their own right, they are also representatives of other products that
77 could be derived from the same precursor metabolites.

78

79 **Materials and methods**

80

81 **Modeling methodology and software**

82

83 Two major strands of stoichiometric modelling are the constraint-based Flux Balance Analysis (FBA)
84 [63, 94] and elementary modes analysis [80]. A constraint based model of central metabolism including
85 biomass production of *K. marxianus* was created adapting and combining the high-quality genome-
86 scale metabolic reconstructions protocol [79] and structural modelling approach for development of
87 medium scale reconstruction and models [39].

88

89 Our medium scale *K. marxianus* central carbon metabolism model is based on the general mass balance
90 equation:

$$91 \quad dX/dt = r_{\text{met}} - \mu X_{\text{met}}$$

92 With respect to intermediate metabolite accumulation, a cell's metabolism is in pseudosteady state and
93 can be described by following equation:

$$94 \quad 0 = r_{\text{met}} - \mu X_{\text{met}} \text{ [85].}$$

95

96 We also assume the following:

97 - the specific growth rate (μ , h^{-1}) during the exponential growth phase is constant,

98 - the cells are at pseudosteady state: substrate uptake, metabolite and product fluxes are constant when
99 μ is constant.

100

101 For constraint based and structural analysis, the ScrumPy modelling package [71] was used. Flux
102 balance analysis (FBA) was carried out by setting a constant rate of substrate uptake to $10 \text{ mM g}^{-1} \text{ DW}$
103 h^{-1} , and searching for the maximum yield of one of the following products: ethanol, acetate, lactate,
104 glycerol, ethylacetate, succinate, glutamate, phenylethanol or phenylalanine. Solutions were further
105 examined using Flux Variability Analysis (FVA) [55] to determine the ranges of internal fluxes that are
106 consistent with the maximum if there were multiple equivalent FBA solutions. Inconsistencies in the
107 model formulation were additionally detected through null space analysis [21] combined with
108 determination of inconsistent enzyme subsets [69] using ScrumPy. The essentiality of genes and
109 reactions was analysed using FBA to check whether biomass production was feasible after deleting the
110 relevant reaction(s) from the model. The gene essentiality test took into account the gene – protein –
111 reaction (GPR) associations [86] that were determined for the model (next subsection). FVA was also
112 used to calculate the potential range in product production taking into account minimal and maximal
113 oxygen respiration levels at a fixed substrate uptake value.

114

115 **Reactions**

116

117 The *K. lactis* genome scale reconstruction [16] was used as a starting point given the high degree of
118 similarity between its metabolic networks and that of *K. marxianus*. The amino acid sequences of *K.*

119 *lactis* genes from the NIH genetic sequence database GenBank [3] were compared against fungal
120 species using NCBI BLAST [38]. The corresponding *K. marxianus* genes were also checked for
121 presence in the Uniprot database [54]. For each reaction, its Enzyme Commission number (E.C. number)
122 and reaction directionality was checked and validated. The IntEnz [22] (available at
123 <http://www.ebi.ac.uk/intenz/>) database was the main reference source for mass and charge balance
124 validation. To represent the *K. marxianus* biomass growth reaction, we used the *S. cerevisiae* biomass
125 composition as described by Gombert et al. [31].

126

127 **Metabolites**

128

129 Metabolite names, their neutral and charged formulas and InChI (International Chemical Identifier)
130 strings [22] were taken from the CheBi database [13] (available at <http://www.ebi.ac.uk/chebi/>), and
131 the yeast-specific Metacyc [7]. The PubChem database [96] (available at
132 <http://pubchem.ncbi.nlm.nih.gov/>) was used to get additional information about metabolites [22].

133

134 ***K. marxianus* strains and cultivation conditions**

135

136 The results of original experiments carried out by us to provide data for model development are marked
137 in Table 1 as “this study”. *K. marxianus* strain DSM 5422 was cultivated in semi synthetic medium
138 containing (g/ l) KHPO₄ (1.0), CaCl₂ (0.1), MgSO₄*7H₂O (0.5), NaCl (0.5), (NH₄)₂SO₄ (5.0) KH₂PO₄
139 (0.1) yeast extract (*Biolife*) (0.5). Different carbon sources (lactose or inulin) were added at
140 concentrations of 5 or 10 % w/v. All fermentations were carried out in 1 litre *Infors* 2HT or 0.4 litre
141 *Sartorius Biostat Qplus* 6-fold system fermenters at 35°C and 400 rpm.

142

143 **Metabolite and biomass analyses**

144 Extracellular lactose, ethanol, acetate and glycerol content were measured simultaneously using an
145 *Agilent 1100 HPLC* system with a *Shodex Asahipak SH1011* column. Metabolites were quantitated
146 with a refractive index detector (RI detector *RID G1362A*). The flow of the mobile phase (0.01 N
147 H₂SO₄) was 0.6 ml min⁻¹, the sample injection volume was 5 µL.

148 Biomass growth was estimated by absorbance measurements at 600 nm (OD600). The conversion
149 coefficient of *K. marxianus* DSM 5422 strain OD600 to culture dry weight was determined
150 gravimetrically: OD600 1.0 was equivalent to 0.3 g dw.L⁻¹.

151

152 **Results and Discussion**

153

154 **Model construction and properties**

155

156 The model is shown diagrammatically in Fig.1 and is supplied in SBML [34] (Online Resource 1)
157 format and in the form of a COBRA [79] MS Excel input file (Online Resource 2). Our *K. marxianus*
158 metabolism model contains 113 reactions and 101 metabolites organized in 3 compartments:
159 extracellular, cytoplasm and mitochondria. There are 72 cytosolic reactions (central metabolism
160 pathways), 28 transmembrane transport reactions, 11 mitochondrial reactions, one extracellular and one
161 biomass reaction (24 components).

162

163 Specific assumptions for our *K. marxianus* FBA model included:

- 164 • ammonium sulfate was the sole nitrogen and sulfur source and was available in excess;
- 165 • extracellular product accumulation had no effect on intracellular reactions;
- 166 • inorganic phosphate was available in excess;
- 167 • NADH and NADPH were assumed not to freely exchange between mitochondria and
168 cytoplasm. Instead, redox equivalents could be translocated across the mitochondrial
169 membrane by specific transport systems (shuttles). A malate – aspartate shuttle [17] and a 2-
170 oxo glutarate – citrate carrier [8] were included to model NAD and NADP dependent redox
171 exchange between cytosol and mitochondria.

172

173 To allow for succinate exchange across the mitochondrial membrane, a succinate - malate carrier was
174 introduced [1, 64]. An electron transport chain was included in the model as a lumped reaction with the
175 P/O ratio set to 1.2 [32].

176

177 AcetylCoA transport across the inner mitochondrial membrane occurs via a carnitine shuttle that is
178 related to fatty acid metabolism [102]. Since the main fluxes for many biotechnologically important
179 products stem directly from short chain carbon metabolites, we decided not to include a representation of
180 fatty acid metabolism and described AcetylCoA transport across the mitochondrial membrane as a
181 simple transport reaction (model reaction ACCOA_DIFF).

182

183 *K. marxianus* is an example of Crabtree negative yeast. Its physiology is believed to be closely related
184 to its sister species *K. lactis* [95]. It is reported that ethanol production in *K. lactis* coincides with
185 decreased oxygen supply [41]. It is assumed, that flux regulation around pyruvate bypass is the reason
186 for Crabtree negative yeasts to choose between fermentation or oxidative growth. The cytoplasmic
187 pyruvate bypass in *K. lactis* consists of pyruvate decarboxylase, NADP-dependent acetaldehyde
188 dehydrogenase and acetyl-CoA synthetase. The first step of the pyruvate bypass in *K. lactis* is strongly
189 upregulated during fermentative growth thus increasing the cytoplasmic production of AcetylCoA [41].
190 In the case of disturbed functioning of the mitochondrial pyruvate dehydrogenase complex, oxygen
191 limitation or blockage of respiration chain, this bypass can supply enough cytoplasmic acetylCoA to
192 support growth [103, 104].

193

194 Acetyl-CoA production in *K. marxianus* mitochondria occurs via the pyruvate dehydrogenase complex
195 (model reaction ed5). In the model, cytoplasmic AcetylCoA synthesis was catalysed by acetyl-CoA
196 synthetase (model reaction ACS). To model anaerobic or semi anaerobic fermentations, an AcetylCoA
197 (reaction ACCOA_DIFF) transport reaction from cytoplasm to mitochondria was included.

198

199

200 **Model validation**

201

202 **Data sources for validation/calibration.**

203

204 The main carbon fluxes for model validation in *K. marxianus* were: substrate uptake, CO₂, ethanol,
205 glycerol, acetate and biomass production. For a *K. marxianus* batch cultivation with limited oxygen
206 supply these fluxes can account for up to 100 % of total carbon [77]. Therefore this set of fluxes is
207 sufficient to validate this medium scale model. A similar set of fluxes has been successfully applied to
208 validate the medium scale carbon metabolism model of *Pichia pastoris* [87].

209

210 Here the model outputs were compared with previously published and original experimental data.
211 Metabolite and biomass data from the exponential growth phase were extracted from numerous
212 published studies involving *K. marxianus* batch cultivations on various substrates (Table 1).

213

214 **Model validation on lactose as substrate**

215

216 Lactose is the main carbohydrate in cheese whey. In *K. marxianus*, lactose is split by the enzyme β -
217 galactosidase into glucose and galactose, then each of these monosaccharides enters glycolysis at
218 different levels: glucose is converted to glucose-6P, but galactose is converted to glucose-1 phosphate
219 by the Leloir pathway. Strains of *K. marxianus* differ with respect to the first steps of lactose
220 metabolism – some strains have intracellular and some - extracellular β -galactosidase [6]. We modelled
221 *K. marxianus* lactose uptake with transport reaction lactD (lactose permease) and breakdown by
222 reaction GALSID (β -galactosidase). The model was able to achieve a steady state solution for all the
223 experimentally measured flux distributions (Table 1).

224

225 In addition to the published studies, we performed aerobic fermentations (semisynthetic broth with 7 %
226 or 10 % lactose, aeration 0.2 or 1 vol vol⁻¹ min⁻¹ of fermentation volume). The measured fluxes of the
227 extracellular metabolites are presented in Table 1 as "this study"; ethanol was the major product with
228 glycerol and acetate as the main byproducts.

229

230 Depending on the oxygen supply, *K. marxianus* lactose fermentation is biomass (aerobic) or ethanol
231 (anaerobic) orientated. Sansonetti et al [77] demonstrated results for *K. marxianus* DSM 5422 strain

232 lactose fermentation in “self anaerobic” mode reaching 3.33 units of ethanol per unit of lactose. In this
233 case biomass growth was slow ($\mu = 0.07 \text{ h}^{-1}$) and glycerol was produced as the main byproduct. On the
234 other hand, rapid biomass production by *K. marxianus* strain CBS 6556 from lactose has been
235 described under fully aerobic mode with comparatively low ethanol flux [51].

236

237 Interestingly, none of above mentioned cases reported acetate accumulation, which seems to be related
238 either to slow cytoplasmic consumption of AcetylCoA (as in the case of low μ), or sufficient
239 AcetylCoA supply by mitochondria (in the aerobic case). Longhi et al [51] reported possible
240 accumulation of acetate during fermentation, albeit they did not report exact concentrations.

241

242 **Model validation on glucose, sucrose and inulin as substrates**

243

244 Glucose, fructose and their derived glucose and fructose oligo- and polysaccharides form an important
245 group of substrates for industrial applications. Sugarcane or sugar beet molasses, starch, sucrose and
246 inulin are typical examples [66].

247 *K. marxianus* is able to hydrolyze inulin directly due to its extracellular inulinase activity [75]. We
248 performed fermentations with strain DSM 5422 in semisynthetic broth with inulin as a sole carbon
249 source; results are depicted in table 1.

250 For strain DSM 5422, extracellular inulinase activity by far exceeded the uptake of released
251 monosaccharides (data not shown). An ample amount of free fructose in the media due to extracellular
252 inulinase activity was also demonstrated by other authors [101]. In the model we assumed, that only
253 fructose is produced after inulin hydrolysis, glucose is released in negligible amount and has no effect
254 on fructose uptake. Similarly, when simulating the data of Etchmann et al. [20], we assumed that
255 sucrose is split outside the cell and invertase activity exceeds the rate of monosaccharide uptake [75].
256 Subsequent simultaneous consumption of glucose and fructose happens when sucrose is hydrolyzed by
257 invertase [23]. Fructose uptake (model reaction inulin_t) followed by fructose kinase (model reaction
258 onoHLK) was considered as a starting point for inulin consumption. All results from inulin, glucose
259 and sucrose fermentations described in table 1 were replicated by the model.

260

261 **Model validation on xylose as substrate**

262

263 *K. marxianus* is able to ferment xylose. As for many yeasts and fungi, in *K. marxianus* xylose is taken
264 up and converted to xylulose-5 phosphate (pentose phosphate pathway intermediate) via three
265 sequential reactions: xylose reductase (reaction XYL1) xylitol dehydrogenase (reaction XDH) and
266 xylulose kinase (reaction pengluc3). Moreover, xylose reductase in *K. marxianus* is exclusively
267 NADPH dependent [106]. Xylose reductase reaction in our model was represented as exclusively
268 NADPH dependent.

269

270 There are many reports of xylose fermentation by *K. marxianus*. We chose three example
271 fermentations [14, 56, 82] to extract data for model validation. All three xylose fermentations yielded
272 slow biomass growth with μ varying from 0.007 to 0.08 h⁻¹. Interestingly enough, the experimental μ
273 values correlated with oxygen supply: increased oxygen supply led to increased μ [82].

274

275 **Reaction and gene essentiality**

276 In this study we linked gene (or reaction) essentiality to the inability to form biomass (maximal
277 biomass flux = 0) on deletion of all reactions catalysed by that gene product. Reaction deletion was
278 performed by setting a zero flux for the reaction in model. Mostly there were one-to one relations
279 between genes and reactions but in some cases there were 1) redundant genes when each of alternative
280 genes encoded enzyme (OR relationship), 2) two or more genes encoded polypeptides that form
281 functional enzyme (AND relationship) and 3) one gene encoded more than one reaction. According to
282 the analysis results (Online Resource 3), the model contained 38 essential reactions. (Fig. 2). The 26
283 reactions (23%) essential for all analysed substrates belonged to central carbon metabolism. Due to the
284 small model size (113 reactions), there were not many redundant or parallel pathways included. Large
285 scale experimental deletion studies with *S. cerevisiae* report 17% [99] and 19% [29] essential proteins
286 for viability in rich medium which is close to our medium scale model based prediction. For
287 comparison, 2 - 66% of reactions are essential across different eukaryotes [9].

288

289 **Model optimisation**

290

291 The model was optimised by FBA for production of ethanol, acetate, lactate, glycerol, ethylacetate,
292 succinate, glutamate phenylethanol and phenylalanine at fixed substrate uptake rate (glucose/inulin,
293 lactose and xylose) at $10 \text{ mM g}^{-1} \text{ DW h}^{-1}$ and μ at 0.4 h^{-1} as a compromise between different substrate
294 consumption and growth rates. The substrate uptake flux was set high to make flux distribution and
295 yield calculations more practical. The maximal percentage of substrate carbon atoms converted into
296 product (Fig.3 A) always was below the maximal theoretical yield (Fig. 3B).

297

298 *K. marxianus* metabolism is sensitive to the oxygen consumption rate. To assess the opportunities of
299 metabolic control by variable oxygen supply, optimisation (FBA) and variability analysis (FVA) were
300 performed for two extreme respiration cases – low (necessary for biomass production) (Fig. 3 C) and
301 high (Fig. 3 D) oxygen consumption rates at fixed $\mu = 0.4 \text{ h}^{-1}$ in the following steps:

302 1) maximal and minimal oxygen consumption was determined minimised/maximised by FVA at $\mu =$
303 0.4 h^{-1} for each substrate;

304 2) 90 and 100% of maximum oxygen consumption rate (determined in step 1) were set as lower and
305 upper oxygen consumption rate bounds FBA analysis at high oxygen consumption;

306 3) minimal and three minimal oxygen consumption rates (determined in step 1) were set as lower and
307 upper oxygen consumption rate bounds FBA analysis at low oxygen consumption;

308 3) maximal product rate at low (Fig. 3C)/high (Fig. 3D) oxygen consumption was determined by FBA;

309

310 In the case of no constraints on oxygen consumption, high values of carbon flux to product (Fig. 3A)
311 were predicted for lactate, glutamate and phenylalanine. Ethanol, acetate and ethyl acetate yields were
312 identical and close to their theoretical maxima (Fig. 3B). The lowest fractions of carbon flux to product
313 were in the cases of succinate, phenylethanol and glycerol. All other cases attained at least 75% of their
314 theoretical yields. Succinate was the only product that had higher yields on xylose as substrate
315 compared to other substrates (Fig. 3A). At minimal oxygen consumption (Fig. 3C) for most products
316 the carbon flux to products was lower in the case of xylose as substrate. With lactose, inulin and
317 glucose as substrates, ethanol and lactate yields were very close to the theoretical maxima. Carbon flux
318 to products at maximal oxygen (Fig. 3D) consumption still was relatively close to maximum in case of
319 ethanol and acetate with lactose, inulin and glucose as substrates while lactate production had become
320 very low.

321

322 **Ethanol production**

323 *K. marxianus* is able to convert lactose, inulin (fructose), glucose and xylose into ethanol. Maximum
324 theoretical ethanol to substrate ratios are: for lactose 4, for inulin/glucose 2, but for xylose 1.6. The
325 model predicted the maximum ratio of lactose conversion into ethanol to be 3.93, inulin/ glucose 1.91
326 and xylose 0.96. While the model predictions for lactose and glucose/ inulin were close to the
327 theoretical maximum, the low ethanol: xylose ratio came as a surprise. This most probably relates to
328 the carbon flux re-routing through the glucose-6P dehydrogenase reaction to generate the NADPH
329 needed for xylose reduction. Additionally, according to the model, a notable portion (0.5 units per unit
330 of xylose) was directed to acetate production.

331

332 Many authors have noted the need for an oxygen supply for *K. marxianus* growth on respiration when
333 fermenting xylose as a sole carbon source [82]. *K. marxianus* respiration mutants are not able to ferment
334 xylose [49]. This dependence might be related to different reasons/ factors.

335

336 Firstly, to metabolise xylose, a *K. marxianus* cell needs enough resources of cytoplasmic NAD⁺ which
337 is consumed by xylitol dehydrogenase (reaction XDH). The cytoplasmic demand for NAD⁺ is fulfilled
338 by the alcohol dehydrogenase reaction (reaction Alcohol4) producing ethanol and NAD⁺ [95] or
339 through the activity of the mitochondrial malate – aspartate shuttle [17, 36]. The model predicted that
340 the latter is not active in the case of a poor oxygen supply.

341

342 Secondly, the model predicted acetate accumulation, up to 50 % of the ethanol flux, if oxygen
343 consumption was kept low. A similar effect was demonstrated in xylose fermentation by *S. cerevisiae*
344 with engineered XYLT, XDH and XK (reaction pengluc3) reactions [44, 70]. According to our model,
345 a decrease in acetate production was observed if the xylose reductase (XYLT) cofactor specificity was
346 changed from NADPH to NADH; in this case the ethanol to xylose ratio reaches 1.6. This *in silico*
347 result complements *in vivo* results from various authors, who engineered the xylose reductase cofactor
348 (enzyme preference to use NADH or NADPH) specificity in *S. cerevisiae* [44] or explored cofactor
349 specificity of wild type xylose reductases of various yeast species [5].

350

351 A third reason why respiration activity is crucial for xylose utilisation, is because of increased ATP
352 consumption to maintain cytoplasmic pH. Stambuk et al. [83] found that xylose uptake in *K. marxianus*
353 is symport. Each xylose is imported together with a proton. In this process cytoplasmic pH drops. To
354 maintain cytoplasmic pH unchanged, cell membrane ATPases exports protons at the expense of ATP
355 hydrolysis.

356

357 The model predicted the maximum ratio of fructose conversion into ethanol to be 1.9 units of ethanol
358 per unit fructose or glucose. Flux variability analyses revealed that if the maximum ethanol flux
359 changes by 10 %, then 4 – 17 % of incoming carbon flux is always routed to glycerol production.
360 Within the limits of error, our *in vivo* data on inulin fermentation were in good correlation with this
361 model prediction. We observed ethanol to monosaccharide flux ratio to be 1.72 +/- 10 % (comprising
362 90 % of maximum) while the rest of the carbon was distributed between glycerol and acetate.

363

364 **Acetate**

365 Acetic acid (ethanoic acid) is among the first chemicals to have been industrially produced by
366 microorganisms. Traditionally bacterial producers (*Clostridium* sp. or *Acetobacter* sp.) are used.
367 Potentially *K. marxianus* can produce acetic acid, though its commercial value is low. Acetic acid
368 accumulation during exponential growth phase is typical for *K. marxianus* fermentations. This has been
369 demonstrated by us as well as other authors [53, 82]. Acetate, as with glycerol, is perceived as an
370 unwanted fermentation side product. In our model, acetate and acetylCoA reactions in our model were
371 cofactor “entangled”, so when there was need for acetylCoA, extra acetate was produced; in parallel,
372 there was a risk of cytoplasmic redox imbalance since acetate is produced along with NADH.

373

374 Maximum theoretical acetate/substrate ratios were calculated: for lactose 4, for inulin or glucose to
375 acetate 2, but for xylose 1.6. The model predicted the maximum ratio of lactose conversion into acetate
376 to be 3.94, inulin or glucose 1.94 and the maximum ratio of xylose conversion into ethanol to be 1.44
377 units of acetate per unit of xylose.

378

379 Acetate production can occur without formation of significant byproducts, but strong aeration must be
380 supplied (optimal oxygen consumption up to 2 units of O₂ per unit of substrate). Based on our model

381 results, cytoplasmic acetate accumulation occurred in two situations – if there was need for extra
382 NADH (as in succinate production) or need for cytoplasmic acetylCoA (in the case of ethyl acetate or
383 biomass production).

384

385 **Lactate**

386 Lactic acid is widely used acid in the food industry and has potential application as a monomer for
387 biodegradable plastics. In both of those applications L-lactic acid is used. Microbial (bacterial, yeast or
388 fungi) fermentation is one of the options for L-lactic acid isomer synthesis in industrial amounts [35].
389 *Kluyveromyces sp.* has been proposed as a prospective lactic acid producer due to its fast production
390 rates and GRAS status [15]. Yeasts do not have lactate dehydrogenase, therefore for lactate production
391 recombinant strains harbouring LDH of eukaryotic origin (mammals, moulds) are used.

392

393 The theoretical molar yield of L-lactate from mole of glucose was 2, from lactose 4, but from xylose
394 1.6. Our model predicted L-lactate formation with the following molar ratios: 3.9 from lactose, 1.9 from
395 glucose and 1.6 from xylose.

396

397 Introduction of heterologous lactate dehydrogenase alone does not lead to maximal L-lactate
398 production *in vivo*. Carbon flow towards lactate or ethanol was divided at the level of pyruvate by
399 pyruvate decarboxylase (reaction in the model PDC) or the pyruvate dehydrogenase complex in
400 mitochondria. If the pyruvate gets decarboxylated, direct lactate production from pyruvate was not
401 possible, instead carbon was routed to acetaldehyde and ethanol or acetate formation. Flux variability
402 analyses revealed that this was the case – when simulating a decrease in lactate flux, an equimolar
403 increase in CO₂ and ethanol fluxes was observed.

404

405 *Kluyveromyces sp.*, unlike *Saccharomyces*, contain just one PDC gene, therefore preparation of *pdc*
406 functional knockouts is comparatively easy. Lactate dehydrogenase overexpression in a *K. lactis pdc*
407 strain has proven to be an efficient strategy yielding a lactic acid: consumed glucose ratio up to 0.5
408 [72]. Lactate production close to the theoretical maximum was achieved when both pyruvate
409 consuming branches (pyruvate decarboxylase and dehydrogenase) were inactivated. A molar lactate/
410 glucose ratio close to 2 in *K. lactis pdc pdh* knockouts was obtained by Bianchi and colleagues [4].

411 Alternatively, additional heterologous expression of lactate dehydrogenase by increasing gene copy
412 numbers can be a strategy to increase lactate production [67].

413

414 **Glycerol production**

415

416 Glycerol is a typical byproduct of yeast ethanol fermentation that forms in response to the need to
417 balance cytoplasmic NADH oxidation. Glycerol formation as an NADH sink becomes crucial when
418 NADH oxidation via the electron transport chain is not possible (limited oxygen supply). Even though
419 glycerol synthesis by microbial producers *per se* has no applications in biotechnology, we included this
420 metabolite in our analyses since this is one of the major carbon and redox sinks in the *K. marxianus*
421 metabolism.

422 Theoretical maximal glycerol production from different substrates in molar ratios were: from lactose 4,
423 from glucose 2, from xylose 1.66. Our medium scale *K. marxianus* metabolic model predicted
424 maximum molar yields from lactose 2.4 from glucose 1.2 and from xylose 1.2. The model predicted the
425 need for a certain respiratory activity (up to 0.5 units of O₂ per unit of substrate) for glycerol
426 production to reach a maximum, hence CO₂ was the only byproduct in the case of optimal glycerol
427 production.

428 We and other researchers have observed similar effects *in vivo* in inulin and lactose fermentations with
429 *K. marxianus* – higher aeration leads to smaller ethanol and glycerol flux and *vice versa* [82]. Severe
430 fermentation dependence on oxygen supply has also been demonstrated in the physiology of *K.*
431 *marxianus*' sister species *K. lactis* [58].

432

433 **Ethyl acetate production**

434

435 Ethyl acetate is a volatile, slightly polar molecule, used as an organic solvent. Nowadays it has many
436 applications in cosmetics (nail polish remover), electronics (cleaning circuit boards, etc.), and has a
437 potential future application as an environmentally friendly acyl acceptor in biodiesel production instead
438 of methanol. Currently ethyl acetate is produced from petrochemical sources, but it can be produced
439 through biotechnological synthesis by many yeasts. Currently *K. marxianus* is regarded as the most
440 productive ethyl acetate producer [52].

441

442 For ethyl acetate, the theoretical molar product/substrate yield when considering pyruvate
443 decarboxylation, was 2 for lactose, 1 for glucose and 1 for xylose. Our model predicted the maximum
444 ethyl acetate to substrate ratio from lactose to be 1.97, 0.72 from xylose, and 0.97 for inulin or glucose.
445 FVA results revealed strong effects of aeration on ethyl acetate formation. Most ethyl acetate was
446 produced at increased aeration. However, the most effective ethyl acetate formation was not during
447 growth with maximal respiration (Fig 3C). Additionally, FVA revealed a notable increase in glycerol
448 production during oxygen limitation, which indicated the necessity of cytoplasmic NADH reoxidation
449 to support acetate production. In the case of respiration, cytoplasmic NADH could be reoxidised
450 through the electron transport chain and mitochondrial shuttle activity.

451

452 Careful fine-tuning of oxygen consumption might be a strategy for maximum ethyl acetate production.
453 A similar strategy was applied when limiting *K. marxianus* access to metal ions [91, 92]. Metal ion (Fe,
454 Cu, Zn) limitation was found to affect ethyl acetate production. Amongst them, Fe limitation had the
455 most effect. *K. marxianus* culture starving for Fe produced ethyl acetate at close to 50% of theoretical
456 maximum. Fe limitation lowered the activities of Fe-dependent mitochondrial aconitase and succinate
457 dehydrogenase; this subsequently led to accumulation of acetylCoA, which was used to increase ethyl
458 acetate production [92].

459

460 **Succinate production**

461

462 Succinate is one of the 12 most recognised sugar-derived chemical precursors. There is
463 biotechnological potential for succinate due to its wide application spectrum, since it can serve as a
464 precursor for tetrahydrofuran, butanediol, succinonitrile etc. Cheap microbial production of succinate
465 has huge market potential [97]. There are already several examples of microbial succinate production at
466 industrial scale (Reverdia, Myriant, BioAmber, BASFPurac, etc.). At least one of the processes is
467 yeast-based (Reverdia, *S. cerevisiae*) [12]. Even though succinate yields close to the theoretical
468 stoichiometric maximum are reached by bacterial cells, yeast offer several advantages over bacteria:
469 they are not obligately anaerobic, they are robust, acid and osmotically tolerant, and non pathogenic
470 organisms [73].

471

472 The theoretical maximal molar ratio for succinate production from lactose was 3, from glucose 1.5 and
473 xylose 1.25. Our *K. marxianus* carbon metabolism model predicted the maximum succinate production
474 ratio from glucose to be 0.55, from lactose 1.1 but from xylose 0.78. From here, it seems, that xylose
475 might be the most potent substrate for succinate production; however, there are not many *in vivo* results
476 on succinate production by *Kluyveromyces sp.* from xylose. Interestingly, a xylose / ethanol mixture is
477 suggested as a prospective substrate for glyoxylate production along with succinate (isocitrate lyase
478 reaction) in *S. cerevisiae* and *K. lactis* isocitrate lyase overexpressed strains [45].

479

480 Succinate can be produced via the tricarboxylic acid cycle or the glyoxylate shunt. It is not a redox
481 neutral product with respect to carbohydrate substrates – theoretically, 2 NAD⁺ are consumed per each
482 molecule of succinate. Reduced cofactors can be oxidised in the electron transport chain or by
483 production of glycerol or ethanol – NAD regenerating pathways. The model predicted accumulation of
484 at least one byproduct when optimised for succinate production. FVA results demonstrated that,
485 depending on oxygen supply, many byproducts were formed. Interestingly, the model predicted
486 glycerol formation in the case of poor aeration, independent of substrate. The compensatory NADH
487 reoxidation through increased glycerol production in *S. cerevisiae* strains, optimised for succinic acid
488 production, was demonstrated *in vivo* [73].

489

490 Based on our medium scale model, phenylalanine can also be formed as a byproduct in rather large
491 amounts (0.3 to 0.6 units of phenylalanine per unit of lactose) if oxygen is supplied in surplus (3.7 units
492 of oxygen per unit of lactose). In this case, production of a relatively large amount of phenylalanine is
493 possible, since our medium scale model is not nitrogen (ammonia) restricted (see model assumptions).
494 In real applications, however, nitrogen bioavailability might prevent such high levels of phenylalanine
495 production being reached.

496

497 Deletion of the genes for succinate dehydrogenase subunits is a popular strategy for yeast-based
498 succinate production [73]. Succinate accumulation in the case of KISDH1 (succinate dehydrogenase
499 subunit) deletion was observed in the case of *Kluyveromyces lactis* [76]. Our model predicted that
500 inactivation of the aspartate malate shuttle in combination with increased oxygen consumption (up to

501 1.3 per unit of glucose) would give maximum succinate yield, while inactivation of succinate
502 dehydrogenase together with inactivated glyoxylate shunt would be preferable in the case of
503 fermentation of xylose.

504

505 **Glutamate production**

506

507 In our central metabolism model, the *K. marxianus* biomass reaction consisted of 24 metabolites,
508 excluding the amino acids, although phenylalanine and glutamate are included in the model as desired
509 products. The amino acid content of yeast biomass is of particular industrial interest, since some of
510 them (like glutamic acid) are responsible for developing of umami taste [37]. Random mutations is a
511 typical method for generating yeast strain with increased glutamic acid content [61]. Here we provide
512 model-based theoretical analysis of possible scenarios for increasing glutamic acid yield from substrate
513 in *K. marxianus*.

514

515 The theoretical maximal glutamate production from different substrates in molar ratios would be: from
516 lactose 2.4, from glucose 1.2, from xylose 1. Our medium scale *K. marxianus* metabolic model
517 predicted maximum molar yields from lactose 1.97, from glucose 0.97, and from xylose 0.8. Glutamate
518 in *K. lactis* and, most probably also in *K. marxianus*, can be produced by either of two reactions:
519 NADP dependent glutamate dehydrogenase (EC 1.4.1.4 reaction GLUDE_nadp) or by GOGAT (EC
520 1.4.1.13, reaction Glude_NAD) [74]. Our model predicted the larger carbon flux to be routed through
521 NADPH-dependent glutamate dehydrogenase. To recover enough cytoplasmic oxoglutarate an NADP
522 – oxoglutarate – citrate shuttle was used (see Fig. 1). At the same time, a high respiration rate was
523 needed to reach maximum glutamate production if glucose or lactose were used as substrates (approx.
524 1.5 O₂ / glucose). Interestingly, a higher fractional of molar yield of glutamate was achieved by xylose
525 fermentation (80 % from theoretical) and less oxygen needed to be supplied per substrate moiety (1.2).
526 In addition, the main glutamate synthesis flux in *K. marxianus* consuming xylose was predominantly,
527 through the GOGAT reaction, unlike when lactose or glucose were consumed (see above).

528

529 In *K. marxianus* glutamate synthesis is tightly product-regulated by feedback inhibition. As in *S.*
530 *cerevisiae*, glutamate synthesis via NADPH dependent glutamate dehydrogenase is subject to nitrogen

531 catabolite repression [59]. To analyse these type of product – substrate interactions, kinetic modelling
532 would need to be applied.

533

534 **Phenylethanol and phenylalanine as products**

535

536 Phenylalanine is the precursor metabolite for many industrial fragrances as well as an ingredient of the
537 artificial sweetener aspartame. Phenylethanol is a rose flavour used in food and pharmaceutical
538 industries. It is usually extracted from rose petals. The global demand for phenylethanol continues to
539 increase, and it cannot be fulfilled by traditional extraction methods. Bacterial synthesis might be an
540 sustainable alternative for phenylethanol production [84]. Traditionally the yeast *S. cerevisiae* is
541 considered as a vehicle for phenylethanol production, though there have been attempts to use other
542 yeasts, including *K. marxianus* [19].

543

544 Central *K. marxianus* carbon metabolism was extended to phenylethanol production. Phenylethanol
545 production in the model was introduced as a linear, 11 reaction chain from PEP and E4P. This chain
546 consumed additional NADPH, NADH, ATP and PEP.

547

548 The final steps of phenylethanol production were introduced according to Uzunov et al. [93]. The
549 model supported two instances of phenylethanol production by *K. marxianus*: from sucrose and
550 glucose. It predicted the maximum phenylethanol to substrate ratio from lactose to be 1.1 and around
551 0.5 from xylose, inulin, sucrose or glucose. That is far above *in vivo* experiments where phenylethanol
552 to substrate ratio ranges around 0.02-0.1 were typically observed [28, 100].

553

554 **Conclusions**

555

556 We developed a mass and charge balanced stoichiometric model of central *Kluyveromyces marxianus*
557 carbon metabolism including biomass production. The model is published in forms ready for
558 simulation (SBML and COBRA toolbox formats). Most of the model information was based on recent
559 *K. marxianus* genome annotations [48]. Mitochondrial shuttles were included to describe proton and
560 molecule allocation between mitochondria and cytoplasm. The model is able to reproduce the

561 experimentally observed mix of industrially valuable products, as well as explaining formation of
562 unwanted side products (acetate and glycerol).

563

564 Our modeling results imply that oxygen control can be used to influence product yield and flux
565 distributions. Also cofactor swapping (NADPH to NADH and/or vice versa) can significantly improve
566 xylose conversion to products. Interestingly, xylose turned out to be a biotechnologically promising
567 substrate for *K. marxianus* with yet unused potential linked to fine-tuned redox engineering. Succinate
568 is a commercially appealing product that could potentially be produced from xylose in a more efficient
569 way compared to other substrates analysed.

570

571 The model predicted that ethanol, acetate, L-lactate and ethyl acetate can be produced at close to their
572 theoretical yield and without the need for genetic engineering in the cases of lactose, glucose and inulin
573 as substrates. At the same time, the model predicted that a high fraction of the phenylethanol and
574 succinate theoretical yields could not be achieved without metabolic engineering, for which the
575 proposed model is a powerful tool.

576

577 Our model can be used for analysis of *K. marxianus* and its metabolic engineering as well as basis of
578 larger scale models. It would be valuable to extend the model to include precise characteristics of
579 transport reactions (proton symport and antiport), together with the plasma membrane H⁺-ATPase
580 system.

581

582 **Acknowledgements**

583

584 The research was supported by ERDF project Nr. 2DP/2.1.1.1.0/14/APIA/VIAA/043.

585

586

587

588 **References**

589

- 590 1. Aliverdieva D, Mamaev DM, Lagutina LS, Bondarenko DI (2010) Transport of
591 dicarboxylates in *Saccharomyces cerevisiae*. *Curr Res Technol Educ Top Appl Microbiol*
592 *Microb Biotechnol* 1611–1620.
- 593 2. Behera S, Sharma NK, Arora R, Kumar S (2016) Effect of Evolutionary Adaption on
594 Xylosidase Activity in Thermotolerant Yeast Isolates *Kluyveromyces marxianus* NIRE-K1
595 and NIRE-K3. *Appl Biochem Biotechnol* 179:1143–1154. doi: 10.1007/s12010-016-2055-2
- 596 3. Benson D a., Cavanaugh M, Clark K, et al (2013) GenBank. *Nucleic Acids Res* 41:36–42. doi:
597 10.1093/nar/gks1195
- 598 4. Bianchi MM, Brambilla L, Protani F, et al (2001) Efficient Homolactic Fermentation by
599 *Kluyveromyces lactis* Strains Defective in Pyruvate Utilization and Transformed with the
600 Heterologous LDH Gene. *Appl Environ Microbiol* 67:5621–5625. doi:
601 10.1128/AEM.67.12.5621-5625.2001
- 602 5. Bruinenberg PM, de Bot PHM, van Dijken JP, Scheffers WA (1984) NADH-linked aldose
603 reductase: the key to anaerobic alcoholic fermentation of xylose by yeasts. *Appl Microbiol*
604 *Biotechnol* 19:256–260. doi: 10.1007/BF00251847
- 605 6. Carvalho-Silva M, Spencer-Martins I (1990) Modes of lactose uptake in the yeast species
606 *Kluyveromyces marxianus*. *Antonie Van Leeuwenhoek* 57:77–81.
- 607 7. Caspi R, Altman T, Dreher K, et al (2012) The MetaCyc database of metabolic pathways and
608 enzymes and the BioCyc collection of pathway/genome databases. *Nucleic Acids Res*
609 40:D742-53. doi: 10.1093/nar/gkr1014
- 610 8. Castegna A, Scarcia P, Agrimi G, et al (2010) Identification and functional characterization of
611 a novel mitochondrial carrier for citrate and oxoglutarate in *Saccharomyces cerevisiae*. *J Biol*
612 *Chem* 285:17359–17370. doi: 10.1074/jbc.M109.097188

- 613 9. Chen WH, Minguéz P, Lercher MJ, Bork P (2012) OGEE: An online gene essentiality
614 database. *Nucleic Acids Res* 40:901–906. doi: 10.1093/nar/gkr986
- 615 10. Cheon Y, Kim J-S, Park J-B, et al (2014) A biosynthetic pathway for hexanoic acid
616 production in *Kluyveromyces marxianus*. *J Biotechnol* 182–183:30–6. doi:
617 10.1016/j.jbiotec.2014.04.010
- 618 11. Chi ZM, Zhang T, Cao TS, et al (2011) Biotechnological potential of inulin for bioprocesses.
619 *Bioresour Technol* 102:4295–4303. doi: 10.1016/j.biortech.2010.12.086
- 620 12. Cok B, Tsiropoulos I, Roes AL, Patel MK (2014) Succinic acid production derived from
621 carbohydrates: An energy and greenhouse gas assessment of a platform chemical toward a
622 bio-based economy. *Biofuels, Bioprod Biorefining* 8:16–29. doi: 10.1002/bbb.1427
- 623 13. Degtyarenko K, de Matos P, Ennis M, et al (2008) ChEBI: a database and ontology for
624 chemical entities of biological interest. *Nucleic Acids Res* 36:D344–50. doi:
625 10.1093/nar/gkm791
- 626 14. Delgenes J, Moletta R, Navarro J (1986) The effect of aeration on D-xylose fermentation by
627 *Pachysolen tannophilus*, *Pichia stipitis*, *Kluyveromyces marxianus* and *Candida shehatae*.
628 *Biotechnol Lett* 8:7–14.
- 629 15. Dequin S, Barre P (1994) Mixed Lactic Acid–Alcoholic Fermentation by *Saccharomyes*
630 *cerevisiae* Expressing the *Lactobacillus casei* L(+)-LDH. *Bio/Technology* 12:173–177. doi:
631 10.1038/nbt0294-173
- 632 16. Dias O, Pereira R, Gombert AK, et al (2014) iOD907, the first genome-scale metabolic model
633 for the milk yeast *Kluyveromyces lactis*. *Biotechnol J* 9:776–90. doi: 10.1002/biot.201300242
- 634 17. Easlon E, Tsang F, Skinner C, et al (2008) Erin Easlon, Felicia Tsang, Craig Skinner, Chen
635 Wang, and Su-Ju Lin 1. *Genes Dev* 931–944. doi: 10.1101/gad.1648308.a
- 636 18. Ende van den W (2013) Multifunctional fructans and raffinose family oligosaccharides. *Front*
637 *Plant Sci* 4:1–11. doi: 10.3389/fpls.2013.00247
- 638 19. Etschmann M, Bluemke W, Sell D, Schrader J (2002) Biotechnological production of 2-
639 phenylethanol. *Appl Microbiol Biotechnol* 59:1–8. doi: 10.1007/s00253-002-0992-x

- 640 20. Etschmann MMW, Sell D, Schrader J (2003) Screening of yeasts for the production of the
641 aroma compound 2-phenylethanol in a molasses-based medium. *Biotechnol Lett* 25:531–536.
642 doi: 10.1023/A:1022890119847
- 643 21. Fell DA, Poolman MG, Gevorgyan A (2010) Building and analysing genome-scale metabolic
644 models. *Biochem Soc Trans* 38:1197–201. doi: 10.1042/BST0381197
- 645 22. Fleischmann A, Darsow M, Degtyarenko K, et al (2004) IntEnz, the integrated relational
646 enzyme database. *Nucleic Acids Res* 32:D434-7. doi: 10.1093/nar/gkh119
- 647 23. Fonseca GG, de Carvalho NMB, Gombert AK (2013) Growth of the yeast *Kluyveromyces*
648 *marxianus* CBS 6556 on different sugar combinations as sole carbon and energy source. *Appl*
649 *Microbiol Biotechnol* 97:5055–67. doi: 10.1007/s00253-013-4748-6
- 650 24. Fonseca GG, Heinzle E, Wittmann C, Gombert AK (2008) The yeast *Kluyveromyces*
651 *marxianus* and its biotechnological potential. *Appl Microbiol Biotechnol* 79:339–54. doi:
652 10.1007/s00253-008-1458-6
- 653 25. Gabardo S, Pereira GF, Rech R, Ayub MAZ (2015) The modeling of ethanol production by
654 *Kluyveromyces marxianus* using whey as substrate in continuous A-Stat bioreactors. *J Ind*
655 *Microbiol Biotechnol* 42:1243–1253. doi: 10.1007/s10295-015-1661-2
- 656 26. Galindo-Leva LÁ, Hughes SR, López-Núñez JC, et al (2016) Growth, ethanol production, and
657 inulinase activity on various inulin substrates by mutant *Kluyveromyces marxianus* strains
658 NRRL Y-50798 and NRRL Y-50799. *J Ind Microbiol Biotechnol* 43:927–939. doi:
659 10.1007/s10295-016-1771-5
- 660 27. Gao J, Yuan W, Li Y, et al (2015) Transcriptional analysis of *Kluyveromyces marxianus* for
661 ethanol production from inulin using consolidated bioprocessing technology. *Biotechnol*
662 *Biofuels*. doi: 10.1186/s13068-015-0295-y
- 663 28. Garavaglia J, Flôres SH, Pizzolato TM, et al (2007) Bioconversion of L-phenylalanine into 2-
664 phenylethanol by *Kluyveromyces marxianus* in grape must cultures. *World J Microbiol*
665 *Biotechnol* 23:1273–1279. doi: 10.1007/s11274-007-9361-3
- 666 29. Giaever G, Chu AM, Ni L, et al (2002) Functional profiling of the *Saccharomyces cerevisiae*
667 genome. *Nature* 418:387–391. doi: 10.1038/nature00935

- 668 30. Gombert AK, Madeira JV, Cerdán M-E, González-Siso M-I (2016) *Kluyveromyces*
669 *marxianus* as a host for heterologous protein synthesis. *Appl Microbiol Biotechnol*. doi:
670 10.1007/s00253-016-7645-y
- 671 31. Gombert AK, Moreira dos Santos M, Christensen B, Nielsen J (2001) Network Identification
672 and Flux Quantification in the Central Metabolism of *Saccharomyces cerevisiae* under
673 Different Conditions of Glucose Repression. *J Bacteriol*. doi: 10.1128/JB.183.4.1441-
674 1451.2001
- 675 32. Groeneveld P (1999) Control of specific growth rate and physiology of the yeast
676 *Kluyveromyces marxianus*: a BioThermoKinetic approach. Free University of Amsterdam
- 677 33. Hong SJ, Kim HJ, Kim JW, et al (2015) Optimizing promoters and secretory signal sequences
678 for producing ethanol from inulin by recombinant *Saccharomyces cerevisiae* carrying
679 *Kluyveromyces marxianus* inulinase. *Bioprocess Biosyst Eng* 38:263–272. doi:
680 10.1007/s00449-014-1265-7
- 681 34. Hucka M, Finney A, Sauro HM, et al (2003) The systems biology markup language (SBML):
682 a medium for representation and exchange of biochemical network models. *Bioinformatics*
683 19:524–531. doi: 10.1093/bioinformatics/btg015
- 684 35. Inkinen S, Hakkarainen M, Albertsson AC, Södergård A (2011) From lactic acid to poly(lactic
685 acid) (PLA): Characterization and analysis of PLA and Its precursors. *Biomacromolecules*
686 12:523–532. doi: 10.1021/bm101302t
- 687 36. Inokuma K, Ishii J, Hara KY, et al (2015) Complete Genome Sequence of *Kluyveromyces*
688 *marxianus* NBRC1777, a Nonconventional Thermotolerant Yeast. *Genome Announc* 3:1–2.
689 doi: 10.1128/genomeA.00389-15.Copyright
- 690 37. Jinap S, Hajeb P (2010) Glutamate. Its applications in food and contribution to health.
691 *Appetite* 55:1–10. doi: 10.1016/j.appet.2010.05.002
- 692 38. Johnson M, Zaretskaya I, Raytselis Y, et al (2008) NCBI BLAST: a better web interface.
693 *Nucleic Acids Res* 36:5–9. doi: 10.1093/nar/gkn201
- 694 39. Kalnenieks U, Pentjuss A, Rutkis R, et al (2014) Modeling of *Zymomonas mobilis* central
695 metabolism for novel metabolic engineering strategies. *Front. Microbiol.* 5:42.

- 696 40. Kerkhoven EJ, Lahtvee P-J, Nielsen J (2014) Applications of computational modeling in
697 metabolic engineering of yeast. *FEMS Yeast Res.* doi: 10.1111/1567-1364.12199
- 698 41. Kiers J, Zeeman AM, Luttik M, et al (1998) Regulation of Alcoholic Fermentation in Batch
699 and Chemostat Cultures of *Kluyveromyces lactis* CBS 2359. *Yeast* 14:459–469.
- 700 42. Kim JS, Park JB, Jang SW, Ha SJ (2015) Enhanced Xylitol Production by Mutant
701 *Kluyveromyces marxianus* 36907-FMEL1 Due to Improved Xylose Reductase Activity. *Appl*
702 *Biochem Biotechnol* 176:1975–1984. doi: 10.1007/s12010-015-1694-z
- 703 43. Kim T-Y, Lee S-W, Oh M-K (2014) Biosynthesis of 2-phenylethanol from glucose with
704 genetically engineered *Kluyveromyces marxianus*. *Enzyme Microb Technol* 61–62:44–7. doi:
705 10.1016/j.enzmictec.2014.04.011
- 706 44. Klimacek M, Krahulec S, Sauer U, Nidetzky B (2010) Limitations in xylose-fermenting
707 *saccharomyces cerevisiae*, made evident through comprehensive metabolite profiling and
708 thermodynamic analysis. *Appl Environ Microbiol* 76:7566–7574. doi: 10.1128/AEM.01787-
709 10
- 710 45. Koivistoinen OM, Kuivanen J, Barth D, et al (2013) Glycolic acid production in the
711 engineered yeasts *Saccharomyces cerevisiae* and *Kluyveromyces lactis*. *Microb Cell Fact*
712 12:82. doi: 10.1186/1475-2859-12-82
- 713 46. Lane MM, Morrissey JP (2010) *Kluyveromyces marxianus*: A yeast emerging from its sister's
714 shadow. *Fungal Biol Rev* 24:17–26. doi: 10.1016/j.fbr.2010.01.001
- 715 47. Lee K, Kim T, Sohn S, et al (2014) Genome-scale metabolic network model reconstruction of
716 *Kluyveromyces marxianus* and strategies for engineering non-native pathways for 3-
717 hydropropionate production in *Kluyveromyces marxianus*. Patent, Pub. No.: US
718 2014/0093901 A1.
- 719 48. Lertwattanasakul N, Kosaka T, Hosoyama A, et al (2015) Genetic basis of the highly efficient
720 yeast *Kluyveromyces marxianus*: complete genome sequence and transcriptome analyses.
721 *Biotechnol Biofuels*. doi: 10.1186/s13068-015-0227-x
- 722 49. Lertwattanasakul N, Suprayogi, Murata M, et al (2013) Essentiality of respiratory activity for
723 pentose utilization in thermotolerant yeast *Kluyveromyces marxianus* DMKU 3-1042.

- 724 Antonie van Leeuwenhoek, *Int J Gen Mol Microbiol* 103:933–945. doi: 10.1007/s10482-012-
725 9874-0
- 726 50. Liu S, Lu H, Hu R, et al (2012) A sustainable woody biomass biorefinery. *Biotechnol Adv*
727 30:785–810. doi: 10.1016/j.biotechadv.2012.01.013
- 728 51. Longhi LGS, Luvizetto DJ, Ferreira LS, et al (2004) A growth kinetic model of
729 *Kluyveromyces marxianus* cultures on cheese whey as substrate. *J Ind Microbiol Biotechnol*
730 31:35–40. doi: 10.1007/s10295-004-0110-4
- 731 52. Löser C, Urit T, Keil P, Bley T (2014) Studies on the mechanism of synthesis of ethyl acetate
732 in *Kluyveromyces marxianus* DSM 5422. *Appl Microbiol Biotechnol* 99:1131–1144. doi:
733 10.1007/s00253-014-6098-4
- 734 53. Löser C, Urit T, Stukert A, Bley T (2013) Formation of ethyl acetate from whey by
735 *Kluyveromyces marxianus* on a pilot scale. *J Biotechnol* 163:17–23. doi:
736 10.1016/j.jbiotec.2012.10.009
- 737 54. Magrane M, Consortium U (2011) UniProt Knowledgebase: a hub of integrated protein data.
738 Database (Oxford) 2011:bar009. doi: 10.1093/database/bar009
- 739 55. Mahadevan R, Schilling CH (2003) The effects of alternate optimal solutions in constraint-
740 based genome-scale metabolic models. *Metab Eng* 5:264–76.
- 741 56. Margaritis A, Bajpai P (1982) Direct fermentation of D-xylose to ethanol by *Kluyveromyces*
742 *marxianus* strains. *Appl Environ Microbiol* 44:1039–1041.
- 743 57. Martynova J, Kokina A, Kibilds J, et al (2016) Effects of acetate on *Kluyveromyces*
744 *marxianus* DSM 5422 growth and metabolism. *Appl Microbiol Biotechnol*. doi:
745 10.1007/s00253-016-7392-0
- 746 58. Merico A, Galafassi S, PiÅ;rkur J, Compagno C (2009) The oxygen level determines the
747 fermentation pattern in *Kluyveromyces lactis*. *FEMS Yeast Res* 9:749–756. doi:
748 10.1111/j.1567-1364.2009.00528.x
- 749 59. Morais-Júnior MA de (2003) The NADP⁺-dependent glutamate dehydrogenase of the yeast
750 *Kluyveromyces marxianus* responds to nitrogen repression similarly to *Saccharomyces*

- 751 cerevisiae. *Brazilian J Microbiol* 34:334–338. doi: 10.1590/S1517-83822003000400009
- 752 60. Morrissey JP, Etschmann MMW, Schrader J, de Billerbeck GM (2015) Cell factory
753 applications of the yeast *Kluyveromyces marxianus* for the biotechnological production of
754 natural flavour and fragrance molecules. *Yeast* 32:3–16. doi: 10.1002/yea.3054
- 755 61. Nakajo Y, Sano H (1998) Yeast extract composition, yeast for obtaining the same, and process
756 for producing yeast extract composition. Patent, Pub. No.: US6344231 B1
- 757 62. Nitiyon S, Keo-oudone C, Murata M, et al (2016) Efficient conversion of xylose to ethanol by
758 stress-tolerant *Kluyveromyces marxianus* BUNL-21. *Springerplus* 5:185. doi:
759 10.1186/s40064-016-1881-6
- 760 63. Orth JD, Thiele I, Palsson BO (2010) What is flux balance analysis? *Nat Biotechnol* 28:245–
761 8. doi: 10.1038/nbt.1614
- 762 64. Pallotta ML, Fratianni A, Passarella S (1999) Metabolite transport in isolated yeast
763 mitochondria: Fumarate/malate and succinate/malate antiports. *FEBS Lett* 462:313–316. doi:
764 10.1016/S0014-5793(99)01535-5
- 765 65. Panesar PS, Kennedy JF (2012) Biotechnological approaches for the value addition of whey.
766 *Crit Rev Biotechnol* 32:327–348. doi: 10.3109/07388551.2011.640624
- 767 66. Patelski P, Berłowska J, Dziugan P, et al (2015) Utilisation of sugar beet bagasse for the
768 biosynthesis of yeast SCP. *J Food Eng.* doi: 10.1016/j.jfoodeng.2015.03.031
- 769 67. Pecota DC, Rajgarhia V, Da Silva N a (2007) Sequential gene integration for the engineering
770 of *Kluyveromyces marxianus*. *J Biotechnol* 127:408–16. doi: 10.1016/j.jbiotec.2006.07.031
- 771 68. Pentjuss A, Odzina I, Kostromins A, et al (2013) Biotechnological potential of respiring
772 *Zymomonas mobilis*: a stoichiometric analysis of its central metabolism. *J Biotechnol* 165:1–
773 10. doi: 10.1016/j.jbiotec.2013.02.014
- 774 69. Pfeiffer T, Sanchez-Valdenebro I, Nuno J, et al (1999) METATOOL: for studying metabolic
775 networks. *Bioinformatics* 15:251–257. doi: 10.1093/bioinformatics/15.3.251
- 776 70. Pitkänen J-P, Aristidou A, Salusjärvi L, et al (2003) Metabolic flux analysis of xylose
777 metabolism in recombinant *Saccharomyces cerevisiae* using continuous culture. *Metab Eng*

- 778 5:16–31. doi: 10.1016/S1096-7176(02)00012-5
- 779 71. Poolman MG (2006) ScrumPy: metabolic modelling with Python. *IEE Proc - Syst Biol*
780 153:375. doi: 10.1049/ip-syb:20060010
- 781 72. Porro D, Bianchi MM, Brambilla L, et al (1999) Replacement of a metabolic pathway for
782 large-scale production of lactic acid from engineered yeasts. *Appl Environ Microbiol*
783 65:4211–4215.
- 784 73. Raab AM, Gebhardt G, Bolotina N, et al (2010) Metabolic engineering of *Saccharomyces*
785 *cerevisiae* for the biotechnological production of succinic acid. *Metab Eng* 12:518–525. doi:
786 10.1016/j.ymben.2010.08.005
- 787 74. Romero M, Guzmán-León S, Aranda C, et al (2000) Pathways for glutamate biosynthesis in
788 the yeast *Kluyveromyces lactis*. *Microbiology* 146:239–245.
- 789 75. Rouwenhorst RJ, Ritmeester WS, Scheffers WA, Dijken JP VAN (1990) Localization of
790 Inulinase and Invertase in *Kluyveromyces* Species. *Appl Environ Microbiol* 56:3329–3336.
- 791 76. Saliola M, Bartoccioni PC, De Maria I, et al (2004) The Deletion of the Succinate
792 Dehydrogenase Gene *K1SDH1* in *Kluyveromyces lactis* Does Not Lead to Respiratory
793 Deficiency. *Eukaryot Cell* 3:589–597. doi: 10.1128/EC.3.3.589-597.2004
- 794 77. Sansonetti S, Hobley TJ, Calabrò V, et al (2011) A biochemically structured model for ethanol
795 fermentation by *Kluyveromyces marxianus*: A batch fermentation and kinetic study. *Bioresour*
796 *Technol* 102:7513–20. doi: 10.1016/j.biortech.2011.05.014
- 797 78. Santharam L, Samuthirapandi AB, Easwaran SN, Mahadevan S (2016) Modeling of exo-
798 inulinase biosynthesis by *Kluyveromyces marxianus* in fed-batch mode: correlating
799 production kinetics and metabolic heat fluxes. *Appl Microbiol Biotechnol* 1–11. doi:
800 10.1007/s00253-016-7971-0
- 801 79. Schellenberger J, Que R, Fleming RMT, et al (2011) Quantitative prediction of cellular
802 metabolism with constraint-based models: the COBRA Toolbox v2.0. *Nat Protoc* 6:1290–
803 1307. doi: 10.1038/nprot.2011.308
- 804 80. Schuster S, Fell DA, Dandekar T (2000) A general definition of metabolic pathways useful for

- 805 systematic organization and analysis of complex metabolic networks. *Nat Biotechnol* 18:326–
806 32. doi: 10.1038/73786
- 807 81. Sharma NK, Behera S, Arora R, Kumar S (2016) Enhancement in xylose utilization using
808 *Kluyveromyces marxianus* NIRE-K1 through evolutionary adaptation approach. *Bioprocess*
809 *Biosyst Eng* 39:835–843. doi: 10.1007/s00449-016-1563-3
- 810 82. Signori L, Passolunghi S, Ruohonen L, et al (2014) Effect of oxygenation and temperature on
811 glucose-xylose fermentation in *Kluyveromyces marxianus* CBS712 strain. *Microb Cell Fact*
812 13:51. doi: 10.1186/1475-2859-13-51
- 813 83. Stambuk BU, Franden MA, Singh A, Zhang M (2003) d-Xylose Transport by *Candida*
814 *succiphila* and *Kluyveromyces marxianus*. In: *Biotechnology for Fuels and Chemicals*.
815 Humana Press, Totowa, NJ, pp 255–263
- 816 84. Stark D, Zala D, Münch T, et al (2003) Inhibition aspects of the bioconversion of L-
817 phenylalanine to 2-phenylethanol by *Saccharomyces cerevisiae*. *Enzyme Microb Technol*
818 32:212–223. doi: 10.1016/S0141-0229(02)00237-5
- 819 85. Stephanopoulos G, Arisitidou A, Nielsen J (1998) *Metabolic engineering: Principles and*
820 *Methodologies*. Academic Press, San Diego
- 821 86. Thiele I, Palsson BØ (2010) A protocol for generating a high-quality genome-scale metabolic
822 reconstruction. *Nat Protoc* 5:93–121. doi: 10.1038/nprot.2009.203
- 823 87. Tortajada M, Llaneras F, Picó J (2010) Validation of a constraint-based model of *Pichia*
824 *pastoris* metabolism under data scarcity. *BMC Syst Biol* 4:115. doi: 10.1186/1752-0509-4-115
- 825 88. Trinh CT, Sreenc F (2009) Metabolic engineering of *Escherichia coli* for efficient conversion
826 of glycerol to ethanol. *Appl Environ Microbiol* 75:6696–705. doi: 10.1128/AEM.00670-09
- 827 89. Trinh CT, Unrean P, Sreenc F (2008) Minimal *Escherichia coli* cell for the most efficient
828 production of ethanol from hexoses and pentoses. *Appl Environ Microbiol* 74:3634–43. doi:
829 10.1128/AEM.02708-07
- 830 90. Unrean P, Trinh CT, Sreenc F (2010) Rational design and construction of an efficient *E. coli*
831 for production of diapolycopendioic acid. *Metab Eng* 12:112–122. doi:

- 832 10.1016/j.ymben.2009.11.002
- 833 91. Urit T, Manthey R, Bley T, Löser C (2013) Formation of ethyl acetate by *Kluyveromyces*
834 *marxianus* on whey: Influence of aeration and inhibition of yeast growth by ethyl acetate. *Eng*
835 *Life Sci* 13:247–260. doi: 10.1002/elsc.201200077
- 836 92. Urit T, Stukert A, Bley T, Löser C (2012) Formation of ethyl acetate by *Kluyveromyces*
837 *marxianus* on whey during aerobic batch cultivation at specific trace element limitation. *Appl*
838 *Microbiol Biotechnol* 96:1313–1323. doi: 10.1007/s00253-012-4107-z
- 839 93. Uzunov ZG, Petrova VY, Ivanov SL, Kujumdzieva AV (2014) In Silico Study of Aro
840 Genes Involved in the Ehrlich Pathway: Comparison between *Saccharomyces*
841 *Cerevisiae* and *Kluyveromyces Lactis*. *Biotechnol Biotechnol Equip* 25:133–
842 137. doi: 10.5504/BBEQ.2011.0128
- 843 94. Varma A, Palsson BO (1994) Metabolic Flux Balancing: Basic Concepts, Scientific and
844 Practical Use. *Bio/Technology* 12:994–998. doi: 10.1038/nbt1094-994
- 845 95. Walker GM (1998) *Yeast Physiology and Biotechnology*. Wiley
- 846 96. Wang Y, Xiao J, Suzek TO, et al (2012) PubChem’s BioAssay Database. *Nucleic Acids Res*
847 40:D400-12. doi: 10.1093/nar/gkr1132
- 848 97. Werpy T, Petersen G (2004) *Top Value Added Chemicals from Biomass*.
- 849 98. Wilkowska A, Kregiel D, Guneser O, Karagul Yuceer Y (2014) Growth and by-product
850 profiles of *Kluyveromyces marxianus* cells immobilized in foamed alginate. *Yeast*. doi:
851 10.1002/yea.3044
- 852 99. Winzeler EA, Shoemaker DD, Astromoff A, et al (1999) Functional characterization of the *S.*
853 *cerevisiae* genome by gene deletion and parallel analysis. *Science* 285:901–906. doi:
854 10.1126/science.285.5429.901
- 855 100. Wittmann C, Hans M, Bluemke W (2002) Metabolic physiology of aroma-producing
856 *Kluyveromyces marxianus*. *Yeast* 19:1351–1363. doi: 10.1002/yea.920
- 857 101. Yuan W, Zhao X, Chen L, Bai F (2013) Improved ethanol production in Jerusalem artichoke
858 tubers by overexpression of inulinase gene in *Kluyveromyces marxianus*. *Biotechnol*

- 859 Bioprocess Eng 18:721–727. doi: 10.1007/s12257-013-0026-9
- 860 102. Zeeman AM, Luttik M a H, Pronk JT, et al (1999) Impaired growth on glucose of a pyruvate
861 dehydrogenase-negative mutant of *Kluyveromyces lactis* is due to a limitation in
862 mitochondrial acetyl-coenzyme A uptake. *FEMS Microbiol Lett* 177:23–28. doi:
863 10.1016/S0378-1097(99)00283-9
- 864 103. Zeeman AM, Luttik MAH, Thiele C, et al (1998) Inactivation of the *Kluyveromyces lactic*
865 *KIPDA1* gene leads to loss of pyruvate dehydrogenase activity, impairs growth on glucose
866 and triggers aerobic alcoholic fermentation. *Microbiology* 144:3437–3446. doi:
867 10.1099/00221287-144-12-3437
- 868 104. Zeeman AM, Steensma HY (2003) The acetyl co-enzyme A synthetase genes of
869 *Kluyveromyces lactis*. *Yeast* 20:13–23. doi: 10.1002/yea.936
- 870 105. Zhang B, Zhang J, Wang D, et al (2016) Simultaneous fermentation of glucose and xylose at
871 elevated temperatures co-produces ethanol and xylitol through overexpression of a xylose-
872 specific transporter in engineered *Kluyveromyces marxianus*. *Bioresour Technol* 216:227–
873 237. doi: 10.1016/j.biortech.2016.05.068
- 874 106. Zhang B, Zhang L, Wang D, et al (2011) Identification of a xylose reductase gene in the
875 xylose metabolic pathway of *Kluyveromyces marxianus* NBRC1777. *J Ind Microbiol*
876 *Biotechnol* 38:2001–2010. doi: 10.1007/s10295-011-0990-z
- 877 107. Zhang J, Zhang B, Wang D, et al (2014) Xylitol production at high temperature by engineered
878 *Kluyveromyces marxianus*. *Bioresour Technol* 152:192–201. doi:
879 10.1016/j.biortech.2013.10.109
- 880 108. Zhang J, Zhang B, Wang D, et al (2015a) Improving xylitol production at elevated
881 temperature with engineered *Kluyveromyces marxianus* through over-expressing transporters.
882 *Bioresour Technol* 175:642–645. doi: 10.1016/j.biortech.2014.10.150
- 883 109. Zhang J, Zhang B, Wang D, et al (2015b) Rapid ethanol production at elevated temperatures
884 by engineered thermotolerant *Kluyveromyces marxianus* via the NADP(H)-preferring xylose
885 reductase-xylitol dehydrogenase pathway. *Metab Eng* 31:140–152. doi:
886 10.1016/j.ymben.2015.07.008

887

888

889 Fig. 1 The scheme of *Kluyveromyces marxianus* central carbon metabolism model

890

891 Fig 2. Model reaction essentiality for biomass production depending on substrate.

892 Analyses revealed 26 essential reactions for all substrates, 5 reactions exclusively essential for
893 xylose, and 4 for lactose. Glucose and inulin are shown as one substrate. Fructose
894 kinase and inulinase were essential reactions for inulin consumption, while hexokinase
895 was essential for both lactose and glucose consumption.

896

897 Fig 3 A. Maximal percentage of substrate carbon atoms converted into product at biomass

898 growth $\mu=0.4 \text{ h}^{-1}$

899

900

901 Fig 3 B. Difference between theoretical yield and maximal carbon flux to product at biomass

902 growth $\mu=0.4 \text{ h}^{-1}$

903

904

905 Fig. 3 C. Maximal percentage of substrate carbon atoms converted into product at low oxygen

906 consumption and biomass growth fixed at $\mu=0.4 \text{ h}^{-1}$

907

908

909 Fig. 3 D Maximal percentage of substrate carbon atoms converted into product at high oxygen

910 consumption and biomass growth fixed at $\mu=0.4 \text{ h}^{-1}$

911

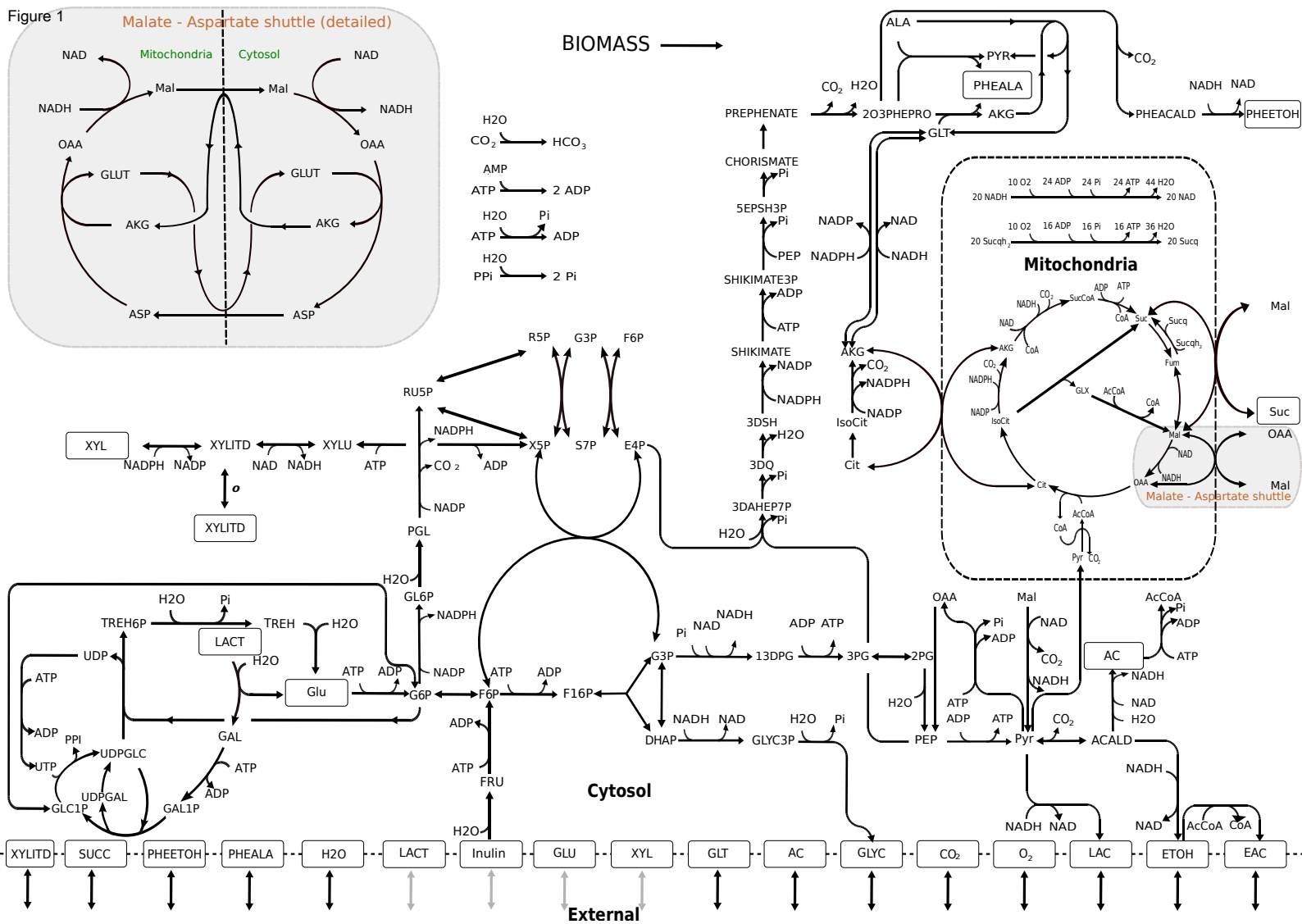
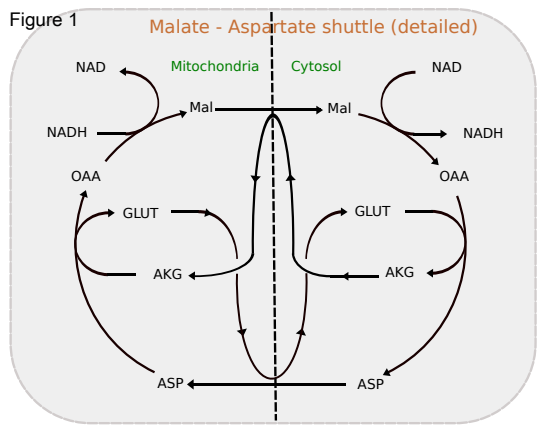


Figure 2

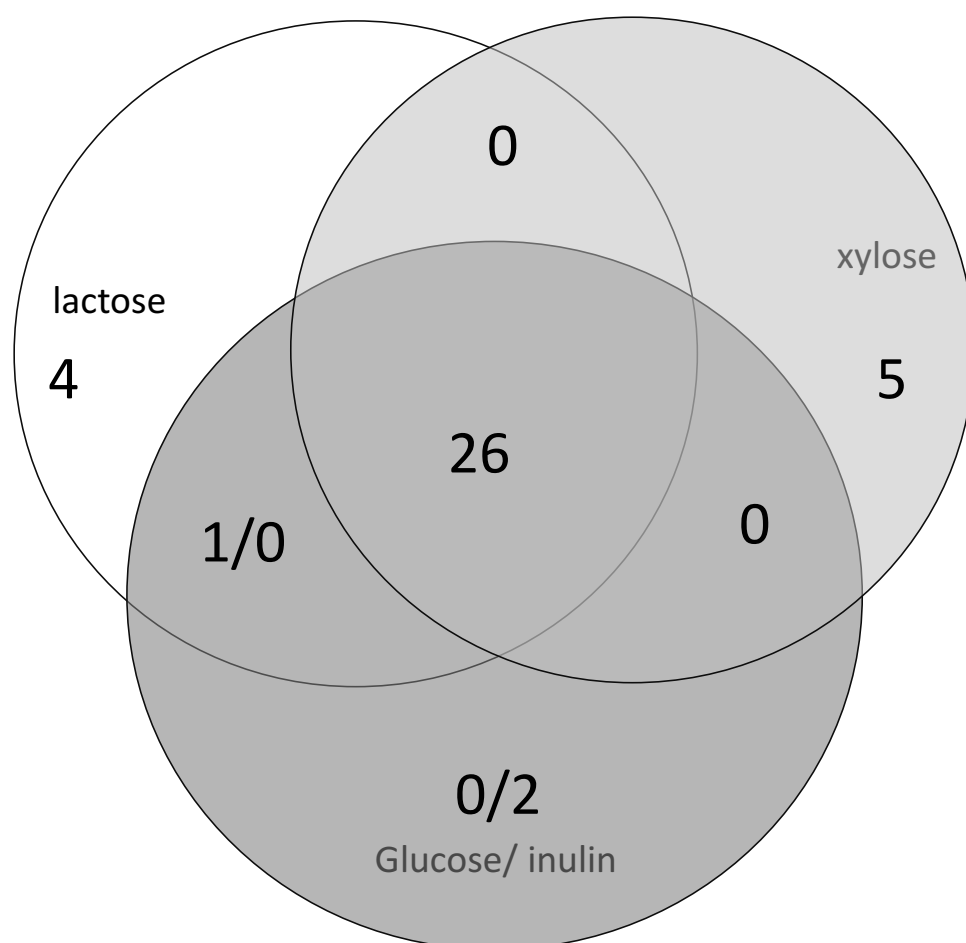


Figure 3

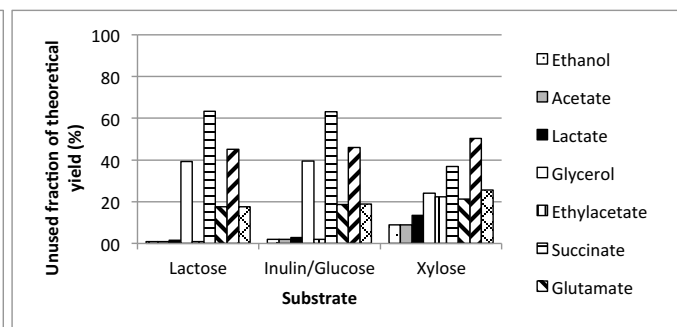
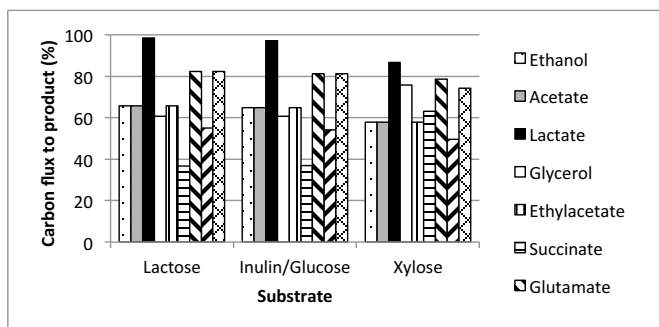


Fig 3 A. Maximal percentage of substrate carbon atoms converted into product at biomass growth $\mu=0,4 \text{ h}^{-1}$

Fig 3 B. Difference between theoretical yield and maximal carbon flux to product at biomass growth $\mu=0,4 \text{ h}^{-1}$

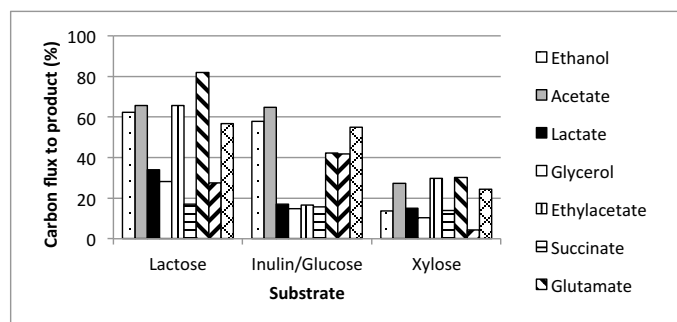
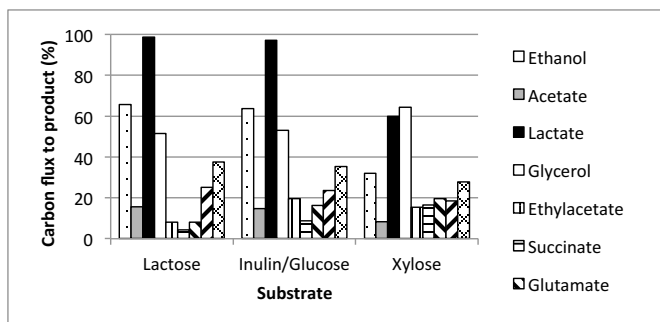


Fig. 3 C. Maximal percentage of substrate carbon atoms converted into product at low oxygen consumption and biomass growth fixed at $\mu=0,4 \text{ h}^{-1}$

Fig. 3 D Maximal percentage of substrate carbon atoms converted into product at high oxygen consumption and biomass growth fixed at $\mu=0,4 \text{ h}^{-1}$

1 **Table 1.** Model validation data. Substrate uptake, biomass growth and product formation fluxes (mM
 2 g DW⁻¹ h⁻¹) were calculated from exponential phase of batch fermentations. Data was extracted from
 3 other author publications or obtained from our fermentations (denoted as “this study”).
 4 Means and standard deviation are calculated from 3 technical replicates where applicable.
 5

Lactose as substrate	Substrate consumption	Biomass μ_{max} h ⁻¹	Ethanol	Glycerol	Acetate	Ethylacetate	Aeration Vol/vol *min
This study, lactose	4.4 +/- 0.4	0.39 +/- 0.06	19 +/- 2	0.72 +/- 0.2	0.24 +/- 0.05	ND	0.2
This study, lactose	5.0 +/- 0.4	0.30 +/- 0.05	9.0 +/- 0.5	0.57 +/- 0.16	0.17 +/- 0.017	ND	1
[57]	13.04	0.31	30.04	0.86	0.19	ND	1
[77]	3.6	0.07	12	0.34	ND	ND	Self anaerobic
[51]	2.3	0.48	2.73	ND	ND	ND	3
[51]	2.1	0.40	7.2	ND	ND	ND	3
[53]	2.4	3.6	1.0	ND	3.0	0.97	1.32
Inulin, glucose and sucrose as substrates	Summary sugar / glucose consumption	Biomass μ_{max} h ⁻¹	Ethanol	Glycerol	Acetate	Phenyl ethanol	Aeration Vol/vol *min
this study, glucose,	15.20	0.48	26.69	1.33	0.013	ND	0.25
this study, glucose,	13.71	0.54	24.25	0.46	0.008	ND	2.5
this study, inulin	5.8 +/- 0.5	0.25 +/- 0.2	10 +/- 1.0	0.16 +/- 0.02	0.04 +/- 0.002	ND	1.5

[43], substrate glucose	0.47	0.02	ND	ND	ND	0.7	ND
[78], substrate inulin	2.60	0.2	ND	ND	ND	ND	Shake flasks
[27], substrate inulin	42.6	0.26	72.4	ND	ND	ND	Self anaerobic
[101], substrate inulin	16	0.14	18	ND	ND	ND	Shake flasks
[28], substrate grape must	2.3	0.40	1.1	ND	ND	0.068	1 and 2
[20], substrate sucrose	4.8	0.26	10	ND	ND	0.33	Shake flasks
[100], substrate glucose	5.0	0.081	1.7	0.34	ND	0.14	1
Xylose as substrate	Substrate consumption	Biomass $u_{max} h^{-1}$	Ethanol	Acetate	Xylitol		Aeration Vol/vol *min
This study,	1.06	0.11	0.18	0.04	ND	ND	2.5
This study,	1.38	0.089	0.085	0.00007	ND	ND	0.25
[82]	0.55	0.014	0	0.4	ND	ND	1
[14]	0.43	0.007	0.28	ND	ND	ND	1
[42]	0.84	0.023	ND	ND	0.66	ND	Shake flasks
[62]	5.78	0.09	3.02	ND	0.04	ND	Shake flasks, +30
[62]	7.54	0.10	2.55	ND	2.44	ND	Shake flasks +37
[81]	1.20	0.13	ND	ND	ND	ND	Shake flasks
[56]	1.83	0.08	1.54	ND	ND	ND	Shake flasks

[2]	0.69	0.12	ND	ND	ND	ND	Shake flasks
Simultaneous uptake of xylose and glucose	Glucose/xylose consumption	Biomass $\mu_{max} h^{-1}$	Ethanol	Acetate	Glycerol	Xylitol	Aeration Vol/vol *min
[105]	4.94/ 6.67	0.17	4.83	ND	ND	4.02	anaerobic

6