1	Maximization of non-nitrogenous metabolite production in E. coli
2	using population systems biology
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15	Abstract
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17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 26	The <i>E. coli</i> metabolome is an interconnected set of enzymes that has measurable kinetic parameters ascribed for the production of most of its metabolites. Flux Balance Analysis (FBA) or Ordinary Differential Equation (ODE) models are used to simulate and increase product yield using defined media. However, such simulations either give a range (min-max) of the metabolite yield for FBA model or tries to enumerate the exact production in the ODE models. The transcriptome expression diversity of the individual cells are not taken into account. We formulate the metabolic behaviour of individual cells by using a robust POpulation SYstem Biology ALgorithm (POSYBAL). This allowed for predicted multiple gene knockouts for increasing homologous metabolite like shikimate and heterologous metabolite like isobutanol. We demonstrate the performance one such triple knockout prediction viz. adhE, ackA and ldhA for isobutanol and aroK, aroA and aroL triple knockout for shikimate. The isobutanol yield increased by 40 times in the knockouts (>2000 ppm) compared to 50ppm produced in the wild-type (BL21) strain, and the shikimate yield was increased to 42 times, i.e. from 58 to ~2100ppm. This formulation was also based on the additional concept of "Nitrogen Swapping" where cells were grown in standard multi-component media during the growth phase and then swapped in low Nitrogen media in the production phase. This swap redistributed the flux distribution such that it flowed primarily through non-nitrogenous pathway such as to maximize the metabolites like shikimate and isobutanol that lack the element Nitrogen in its constituent. Further analysis of various feature of the prediction of the POSYBAL model indicates that even under normal glucose uptake the bacterial cell population diverges into rapidly growing and nearly non-growing cells thus increasing its diversity and hence robustness under any antibacterial attack. This feature is discussed from an evolutionary standpoint.
37 38	Keywords: Isobutanol, Shikimate, <i>Escherichia coli</i> , Knockouts, Flux balance analysis, Nutrient modulation, Minimal media, Matrices

40 Introduction

41

42 Naturally growing bacteria have evolved to survive in the "occasionally famine and rarely feast" conditions for 43 biomass. In contrast, the nutritional factors are made optimal in laboratories to return higher biomass, 44 subsequently increasing the product yield. Inherent in this optimizing process is often an opposite pull of flux of 45 either maximizing the biomass which is a near precise stoichiometric summation of multiple essential 46 metabolites or the specific maximization of any particular metabolite. If these 'nutritional conditions' can be 47 exploited, we can have a 'minimalistic' system to carry out reactions of commercially valuable metabolites with 48 low-cost inputs. To this end, we need to construct an in-silico model and validate its formulation by precise 49 experimental operations.

50

51 One of the significant drivers for constructing in-silico models for different micro-organisms in the field of 52 metabolic engineering is to forecast the genetic changes (gene deletions and over expressions) required in 53 enhancing the levels of a specific metabolite of the interest. There are two types of platform that are commonly 54 used to simulate a bacterial cell. It is either based on linear Flux Balanced analysis (FBA) platform [1-4] or 55 dependent on ordinary differential equations (ODE) where kinetic parameters (Km and Vmax) of enzymes 56 participating in the particular reaction type (single substrate, multi-substrate, ping pong, ternary complex etc.) 57 are used [5-8]. The popular FBA platform generally incorporates the development of a stoichiometric model 58 with genome-level annotations of pathways that maps conversion of xi moles of substrates to xj moles of

59 with genome-rever annotations of partways that maps conversion of x1 moles of substrates to x1 moles of 59 product. The simulations thereof are used to predict knockouts of non-essential genes, in specific pathways

60 (identified during the simulation studies) which would help in blocking the formation of particular metabolites

and allow the cells in redirecting the carbon feed into the production of the metabolite of interest.

62

63 All types of in-silico model, whether based on the mathematical framework of FBA or ODE format tacitly 64 assume that all the cells are in an identical metabolic state. However, bacterial population are generally 65 asynchronous in their growth and their reaction to a stimulus (positive or negative) at the individual cellular 66 level are not unique and can only be represented by a diverse array of responses. In a population of bacterial 67 cells under a standard nutrient condition, some proliferate while some are stunted. Similarly, upon exposure to 68 an antibacterial compound over a period of time, say about 90-99 % of the population are affected, while the remaining 1-10% remain unaltered due to the robustness in the system. For eg. the heterogenous expression of 69 70 araBAD promoter in the presence of limited arabinose quantities in *Mycobacterium tuberculosis* shows variation 71 in individuals of a population [9]. Additionally, as summarized in a recent review [10] it is a truism that in a 72 population of bacterial cells, the gene expression levels are highly variable in lower nutrient concentrations.

73

74 Traditional modelling assumes every bacterium to have equal exposure to the participating nutrients. Such a 75 'socialistic' approach is often invalidated in a natural environment where intra-species competition persists 76 leading to crony capitalism, among bacteria where nutrient availability is the 'currency' for driving a reliable 77 binary of 'haves' and 'have-nots'. In essence, the collection of cells in a group with the same genetic makeup 78 (isogenic) have different expression profile. Hence, there is an urgent need to develop a population model where 79 each cell has a unique metabolic signature.

80

For understanding the optimal flux of pathways in a population, we can use a similar approach. Several
industrially relevant metabolites are devoid of nitrogen. With glucose as the sole carbon source, the anabolism
of nitrogen-containing metabolites such as nucleotides and amino acids, are further downstream compared to the
C, H, O based metabolites. Hence, it is easier to use the POSYBAL platform owing to the requirement of lesser
dimension of matrices and hence lower computing power. We describe the mathematical foundations of a
population model extending the FBA architecture and validate its predictive power by constructing strains of *E. coli* with minimal genetic modifications that produce non-nitrogenous metabolites such as Isobutanol and

88 Shikimate at multigram levels.

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- 89 Isobutanol is produced by altering the pathway from the formation of branched-chain amino acids (BCAA),
- 90 whereas shikimate is an intermediate metabolite of the aromatic amino acid pathway (AAA) respectively.
- 91 Isobutanol is an important industrial solvent which has higher calorific value than ethanol making it an ideal fuel
- 92 substitute. It requires no infrastructure modifications for transport, and unlike ethanol, it is not hygroscopic and
- 93 is non-corrosive to motor engines. The generation of CO2 instead of SO2 or CO makes it a clean fuel compared
- by to the fuels derived from petroleum. On the other hand, shikimate is a high-value industrial precursor for
- producing herbicide such as Glyphosate and the antiviral compound Tamiflu. [11,12].
- 96

97 Both Isobutanol and shikimate are produced by designed alteration of the BCAA and AAA pathways

respectively. Figure 2 illustrates the metabolic pathways required for their production. The unique feature of all
 the intermediates/metabolites that are formed from glucose to isobutanol and shikimate (Figure 2) is the absence

- 100 of nitrogen in the metabolite composition. Hence, after producing the biomass in a rich media we generated the
- 101 metabolite by using media containing either zero or limited amount of nitrogen. This was done to force the flux 102 towards the production of isobutanol and shikimate. We term this shift of media as 'Nitrogen Swap' (N-swap)
- since it is literally the swapping of nitrogen to obtain more product yield. Of course, nitrogen is an essential
- element for cellular growth and is a part of proteins and DNA/RNA. Hence under nitrogen depleted condition,
 the growth of the cell is halted. However, since carbon, hydrogen and oxygen are present, the metabolite flux
- 106 does proceed through Nitrogen independent pathway.

Additionally, we use BL21 strain, a B-Strain of *E. coli* the strain has a valine-feedback independent acetolactate
synthase (ilvG). This, in addition to the heterologously expressed ketoisovalerate decarboxylase (KivD), is
better suited compared to the K12 (BW25113) strain for isobutanol production. Similarly, BL21 gave better

110 Shikimate yield than BW25113, albeit for unknown reasons (Supplementary figure S1).

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An additional advantage of N-swap in case of Isobutanol production is that under anaerobic growth conditions nitrate formation inactivates the enzyme Dihydroxy-acid dehydratase (IIvD). Dinitrosyl iron complex bound

114 IIvD is an inactive enzyme complex making the bacteria a BCAA auxotroph effectively halting isobutanol

production. However, under limiting or no nitrogen in N-swap condition, the NO formation is effectively

116 stunted, which helps in keeping the flux active through the BCAA pathway. This complex is activated under the

aerobic condition without the formation of a new enzyme [13]. We demonstrate a scheme which considers the

118 minimal amount of input with nominal gene manipulations necessary for the production of isobutanol and

shikimate in the BL21 E. coli strain.

120

122 Methods and protocols

123 Population system biology algorithm (POSYBAL):

124 In a traditional FBA model, the metabolites and their reactions are represented in a matrix (S) of size $a \times b$. 125 Every metabolite is represented by a row and reactions are represented by every column. The balancing 126 stoichiometric coefficients are entered into each column that enables computation of the compounds 127 participating in a reaction. As most, if not all reactions are reversible, a negative coefficient shows a catabolic 128 reaction and a positive coefficient portrays a metabolite anabolism whereas, the coefficient is entered as 0 when 129 a metabolite does not participate in the reaction. Individually, most enzymes intrinsically, are reversible but the 130 forward reaction is significantly higher than the reverse reaction. For our model, we have made the it 131 unidirectional for computational convenience. Mostly the matrix is sparse as biochemical reactions involve only 132 a few different metabolites. In general, the number of compounds/metabolites is lesser compared to the number 133 of reactions and hence more unknown variables are present than the number of equations. Consequently, making 134 the system underdetermined, and allowing the existence of multiple solutions rather than a unique one (a>b). 135 Constraining the reaction fluxes limits the range of the solutions for the model, which was manually curated to 136 make them unidirectional based on literature survey. Figure 1D shows the reduction of solution space (red) 137 where eventually an optimal solution (Orange) is obtained in traditional FBA by maximizing or minimizing the 138 target reactions [14]. Instead, in POSYBAL platform, the samples are fetched from the entire constrained 139 solution space randomly that encompasses the behaviour of an underdetermined system (population).

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141 At the systemic level each reaction is mediated by specific gene expression in the organism which is a 142 dimension or variable. The flux of a reaction is the quantitative solution sought by ensuring the conservation of 143 the mass at the systemic level. This essentially translates into a relationship between the reaction fluxes such 144 that, sum of reaction fluxes that drive up metabolites is on the right-hand side and this equals the sum of the 145 reaction fluxes in the left-hand side. We have used the standard convention that left hand side metabolites are 146 substrates and the right-hand side are products and allowing for reversibility of the reactions in which case the 147 fluxes are found to be negative. While the figure 1D shows three dimensions as an illustration, the number of 148 dimensions of the system is equal to the number of reactions that have been considered to govern the system. In 149 the context of using FBA in systems biology the solution most sought is either the maximization of biomass or 150 the maximization of the target metabolite reaction. We can identify the essential genes and the pathways in the 151 given system by maximizing the biomass reaction in contrast when the target metabolite reaction is maximized 152 the biomass has a tendency to run minimum.

153

154 We have created the shikimate model from IJO1366 [15] with the addition of a reaction corresponding to the 155 export of the shikimate metabolite. In the absence of a transport flux the system considers the shikimate 156 metabolite to be unused when reactions contributing to the utilization of the shikimate are knocked out, bringing 157 down the shikimate flux to 0. This prevent us from arriving at knockouts that can maximize shikimate. In a 158 different approach one need not optimize the system of linear equation for a given objective function but try and 159 see the solution distribution over a very large set of solutions within some boundary. This method tries to work 160 around the optimization problem where it is inherently assumed that the system has some definitive but 161 unknown intelligence to work towards (turn on genes expressions suitably) to attain maximization or 162 minimization of the unique function. The modified approach that is computationally very intensive looks to 163 setup a bounding set of minimal constraints so that all solutions that are possible to lie within a bound and then 164 try and generate sample solutions from the entire solution space obtaining a matrix of multiple solutions 165 (POSYBAL) using an algorithm based on Markov chain [16] and this data comprises of all the possible solution 166 (ways) in which the organism can act (population behaviour) corresponding to a given condition. The 167 POSYBAL data comprising of 100000 samples was developed (See supplementary methods 2 and 3 for detailed 168 protocol, scripting and output information).

169

Once the population results are obtained (i.e. multiple matrices), the system is filtered for the solutions with
 maximum production of the target metabolite. The maximum flux is identified and all the iterations with ~90%

and above of the maximum flux are filtered out. The next step is to identify the fluxes that run less than $\sim 10\%$ of

the maximum flux of respective reactions. These indicate knockouts for validating the platform in vitro. A

wrapper function using R programming [R development core team, 2010] is built to filter the possible knockouts

175 from a population.

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177 Generation of predicted *E. coli* knockouts:

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179 Both the triple knockouts i.e. $\Delta ackA:\Delta adhE:\Delta ldhA$ and $\Delta aroA:\Delta aroK:\Delta aroL$ were generated using P1 180 transduction method (See supplementary methods). Δ ackA (BW strain) was used the donor and the BL21 strain 181 was used as the recipient. Subsequently, $\Delta adh E$ and $\Delta ldh A$ were used to knockout the respective genes in BL21. 182 Similarly, BW dackA strain form in-house KIEO library of E. coli knockouts which have a kanamycin cassette 183 in place of the knocked-out gene as a marker. After successful transduction, BL21 AackA strain is made with 184 Kanamycin cassette replacing the knocked-out gene. This is confirmed by colony PCR. For shikimate 185 production Δ aroA from KIEO collection was used as the recipient strain. To further knockout the genes, the 186 kanamycin marker is first flipped out by using the λ -red recombinase method. pCP20 plasmid is first 187 transformed into the desired knockout and plated on LB-amp (30ug/ml) plate and incubated overnight. The 188 colonies are then grown in Luria broth until they reached 0.6 OD_{600} . Then, the culture is incubated at 37°C for 189 one hour followed by incubation of 43°C for four hours. It is then, plated on plain LB media, media containing 190 ampicillin and media containing kanamycin respectively. Growth of the culture on LB plate and no growth on 191 media having either ampicillin or kanamycin confirmed the absence of kanamycin cassette.

192

193 Protocol for isobutanol and shikimate production with various knock-outs and media swap:

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Initially, the conformation of in-silico simulations for nitrogen modulation was done in shake flasks before proceeding towards biotransformation in bioreactors. The desired knockouts and wild-type strains are then transformed with pUC57a kivD plasmid and plated on Luria agar plates with ampicillin (100ug/ml). A single transformant is then inoculated in 5ml of LB media with ampicillin (100ug/ml) and grown overnight. The pUC57-kivD plasmid, synthesized by Genscript, is engineered without an operator site and with a constitutive promoter therefore making the induction step void. For shikimate the cells (or knockouts) are grown overnight in LB media.

202

203 Shake-flask bioconversion experiments:

To conduct the shake-flask experiments for both shikimate and isobutanol growing the starter cultures in LB media, they are transferred to fresh Lysogeny broth and grown until 2.0 OD 600 (Secondary culture). Then the cells are centrifuged at 4000g and transferred into nitrogen deficient media. In case of Isobutanol, the knockouts are transferred to M9 media with varying percentage of Ammonium Sulphate (Nitrogen source) composition with 3.6% glucose (Carbon source) and ampicillin (100ug/ml) for sustaining the pUC57-kiVD plasmid. For shikimate production, the cultures are transferred into M9 media with varying percentage of LB with 1.6%

210 glucose.

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213

To get higher product yield the cells were grown in 500ml Bioreactor (Applikon miniBio). For producing

isobutanol and shikimate the cells were grown up to ~ $6.0-6.5 \text{ OD}_{600}$ in 20% dissolved oxygen (DO) at 200 rpm

216 impeller speed. PEG400 was added as an anti-foaming agent. To create microaerophilic conditions the DO is

reduced to 2.5% and impeller speed is reduced to 50 rpm to produce either isobutanol or shikimate.

²¹² Protocol for bioreactor

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219 Estimation of cell viability

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Cell viability was estimated by plating the spent culture in LB and LB with Ampicillin (100ug/ml) plates. This
 is done to give a projection of the number of cells alive and the plasmid loss after various time points. Samples
 are sent for GC analysis/HPLC performed with the corresponding media standards. We estimate isobutanol and

- ethanol produced along with the remaining glucose concentration, as well as other metabolites like formate,
- 225 lactate, acetate, succinate and pyruvate.

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HPLC analysis

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- 229 For the detection of both Shikimate and Isobutanol,1ml of the culture was spun at 4000 g for 5 minutes,
- followed by further spinning of the supernatant at 14.8 g for 5 minutes. About 50ul of supernatant was analysed
- 231 in HPLC. To perform the HPLC Aminex 87H column (Biorad) was used as stationary phase and 5mM H2SO4
- was used as the mobile phase at 0.750ml/min flowrate with RID detector which is useful for detecting
- 233 monosaccharides and organic acids at the same time.

235 Results

236 Looking at a metabolic map it is might be a non-controversial conjecture, for the carbon flux to proceed towards 237 the production of higher isobutanol or shikimate, lesser amount of nitrogen input is required. In fact, figure 2 238 shows the number of steps from glucose to the production of isobutanol (15 steps) and shikimate (12 steps) 239 which don't have nitrogen. Also, BL21 strain was used for the production of isobutanol due to valine-feedback 240 independent acetolactate synthase (ilvG) but the product yield for shikimate was also higher using the same 241 strain and its knockouts (Supplementary figure S1) for unknown reasons. Figure 3 shows the essential 242 requirement of aromatic amino acids in order to maintain cell viability for shikimate production. Whereas, for 243 isobutanol production this is not essential since there are no biochemical repercussions affecting cell viability. 244 Hence, isobutanol production is an ideal candidate for testing the effect of limited nitrogen source on product 245 yield. Figure 3 shows that the consumption of glucose remains same in both nitrogen and nitrogen depleted 246 conditions however, the production of isobutanol is increased by almost 30% in media devoid of nitrogen (i.e. 247 0% N). But this cannot be directly applied for increasing shikimate production as aromatic amino acid synthesis 248 is downstream to shikimate unlike valine.

249 The POSYBAL platform was used to identify the knockouts required for increasing the shikimate/isobutanol 250 yield. Knockouts were derived by observing the flux through metabolite pathways. A negligible flux essentially 251 represents a knockout. Hence, as depicted in figure 1C a matrix of multiple solutions was used for finding the 252 best fluxes for isobutanol and shikimate production. A set of 10⁵ iterations (Figure 4A) was used for simulating 253 isobutanol and 10⁴ iterations (Figure 4B) were used for simulating shikimate production. Each dot represents 254 one solution among the solution space. The optimal knockouts (encircled in Figure 4) for increased biomass 255 against product yield for isobutanol is Δ ackA: Δ ldhA: Δ adhE triple knockout and shikimate yield is 256 $\Delta aroA:\Delta aroK:\Delta aroL$ knockout. Further, to prove the significance of POSYBAL platform, about 10⁴ iterations 257 were done for Δ ackA, Δ ldhA, Δ adhE in various combinations. This produces a scatter plot of bacterial 258 population producing isobutanol. These 'knockouts' are produced by introducing negligible flux through the 259 respective genes. Figure 5A shows the progression of population towards the 'peak' of the population 260 distribution as the flux reduces from Wildtype (BW25113) to single, double and eventually triple knockout of 261 Δ ackA, Δ ldhA, Δ adhE. Hence, it is easier to understand a population behaviour using this platform unlike ODE 262 and FBA simulations. Figure 5B shows the normalized HPLC data for the wild type (BL21) and 263 ΔackA:ΔldhA:ΔadhE triple knockout. There is an increased production of isobutanol in nitrogen depleted 264 conditions especially at 3% nitrogen and a decreased production of acetate and no production of lactate in the 265 triple knockout.

266 For shikimate production the minimal requirements are indirect as the aromatic amino acid (essential) 267 production is downstream. It is observed that despite intuitively choosing ackA based on Figure 2 as a knockout 268 for producing more shikimate, the in vitro output proves otherwise (Figure 6B). POSYBAL simulations were 269 performed for some of the knockouts derived from Figure 4B. The individual scatter plots (Figure 6A) show 270 inverse triangle relationship for aroL, pykA/F and ptsG. Hence, in these cases knockouts are important for 271 increasing product yield. Whereas, gaussian plots are seen for ackA and partially for poxB. The scatter plots for 272 knockouts which showed higher shikimate production. The shake flask experiments were performed with the 273 previously predicted $\Delta aroK:\Delta aroA:\Delta aroL$ knockout along with $\Delta poxB:\Delta aroK:\Delta aroA:\Delta aroL$ and 274 $\Delta ptsG:\Delta aroA:\Delta aroA:\Delta aroL$ quintuple knockouts (Figure 6B). The experiments were done with various nitrogen 275 source (LB) concentrations of which the 20% LB (LB/5) was found to be the optimal concentration for 276 maximum product yield. It is also seen that there is negligible flux of glucose without the nitrogen source. This 277 shows that a certain amount of nitrogen is required for driving the carbon flux towards shikimate production. 278 Figure 6B also shows the limited acetate required for higher shikimate yield in $\Delta aroK:\Delta aroA:\Delta aroL$. The 279 addition of poxB and ptsG knockouts reduce the acetate production by 68% (see supplementary table 2). This in 280 turn increases the shikimate production in 20% LB (Figure 6B). Although the quadruple knockouts show that 281 POSYBAL simulations help in understanding the population behaviour and flux of carbon, we sought the best 282 triple knockouts which gave higher shikimate yield with enzymes such as tkaA and pntAB which showed direct 283 correlation with shikimate production. These enzymes were taken from the ASKA collection and expressed in 284 BL21 \(\Delta aroK:\Delta aroA:\Delta aroL knockouts.) The experiment was performed in bioreactor with 20% LB with 08 and 285 1.6% glucose. The production of shikimate after 24 hours in $\Delta aroK:\Delta aroA:\Delta aroL$ was 1634 ppm which was 286 doubled by tktA overexpression to 3022 ppm and whereas, pntAB overexpression gave initial higher yields i.e. 287 1130 ppm (at 12 hours) but they remained in similar concentration (1245 ppm) after 24 hours as well (Figure 288 7A). For isobutanol production a fairly straightforward correlation was observed in 3% nitrogen containing

289 complete minimal M9 media where the isobutanol concentration produced was 2235 ppm after 24 hours (Figure 290 7B).

291

292 Discussion

293

294 Traditionally, the production of butanol was done by ABE (Acetone, n-butanol and Ethanol) fermentation 295 method. The carbon source used during this anaerobic fermentation procedure was starch with Clostridium 296 acetobutylicum (or C. beijerinkii) as a whole-cell catalyst [17.]. The solvents were produced in a ratio of 3 parts 297 Acetone, 6 parts Butanol and 1-part Ethanol. In the post genomic era, the use of systems biology and 298 heterologous gene expression have ushered a revival in "bio-butanol". Atsumi et al. [18] showed that the 299 integration of the Ehrlich pathway into the branched chain amino acid pathway was sufficient to generate 300 isobutanol in E. coli under non-fermentative conditions. Also, the addition of the heterologous gene kivD 301 (ketoisovalerate decarboxylase) from L. lactis was required to produce isobutanal which is converted to 302 isobutanol by multiple native isobutyraldehyde dehydrogenases such as YqhD, AdhP, FucO, EutG, YaiY, BetA, 303 EutE and YjbB [19] (Figure 2). Historically, Shikimate is produced by using plant sources, since they contain 304 similar biosynthetic pathways. Star anise (*Illicium anisatum*) is used for extracting shikimate (1.5% w/v) [20]. A 305 better substitute is sweetgum (Liquidambar styraciflua) which has a product yield of 2.4-3.7% w/v [21]. 306 Engineering microorganisms is imperative for higher yields or easier extraction processes. In E. coli It is 307 obtained by knocking out genes further downstream which produce aromatic amino acids such as aroK, aroL [22].

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310 However, static biochemical maps often don't portray the nuances of complex dynamic systems such as a cell. 311 Even, traditional FBA or ODE models would give a single solution which may or may not work in vitro. 312 However, POSYBAL simulations are different from traditional FBA models as they consider the overall 313 'presence' of a metabolite 'through' a population of cells. Hence, multiple combinations entail this concept and 314 result in a normal distribution of substrate vs metabolite. Although, this stochastic combination requires 315 intensive computational power, it is useful for determining combinations which may not be always 'intuitive'. 316 The platform shows different phenotypes of a single strain in its population. Notice, that the 'cells' in the scatter 317 plot (Randomly picked points from the solution space) portrays a specific behaviour of a particular phenotype's 318 reaction flux and its corresponding effect on the target metabolite of interest. In general, three types of patterns 319 are observed namely, inverse triangle correlation where a knockout may result in the production of a target 320 metabolite. A direct triangle shows that a gene overexpression is required for metabolite production. A random 321 scatter means there is no correlation involved. In the current study we demonstrate this with two examples, the 322 production of isobutanol is straightforward since, valine biosynthesis can continue with limited nitrogen and the 323 carbon flux can be diverted towards the production of acetolactate (and subsequently isobutanol) through 324 microaerophilic conditions. However, in case of shikimate this approach fails as a stoichiometric deficit is 325 observed. An example of this is demonstrated here for the role of ackA in shikimate synthesis. One may assume 326 that knocking out acetate production (ackA) in Figure 2 would divert the flux towards shikimate production. 327 However, this is counter-productive as seen in figure 9.

328

329 Intuitively it can be assumed that knocking out acetate production (ackA) in Figure 2 would divert the carbon 330 flux towards shikimate production. However, this is counterintuitive as seen in Figure 9A. With FBA models a 331 stoichiometric assumption can be made since 2 molecules of glucose ($C_6H_{12}O_6$) are needed to produce one 332 molecule of shikimate ($C_7H_{10}O_5$). But, the atomic excess of $C_5H_{14}O_7$ has to be converted or excreted in other 333 metabolic forms else, the reaction would reverse in order to maintain the cell viability. With the POSYBAL 334 platform it is feasible to find out if a knockout, knockdown or overexpression is required for optimal production 335 of shikimate (or any other metabolite). It is observed that despite intuitively choosing ackA as a knockout for 336 producing more shikimate, but the in vitro output shows otherwise. Through POSYBAL platform it is seen that 337 poxB, pykA/F and aroL can be knocked out to produce high amount of shikimate whereas an 'intermediate' flux 338 through ackA and ptsG (Figure 9A) produces higher shikimate than its knockout. Figure 9B shows that limited 339 amount of acetate (ackA expression) is required for the production of Shikimate. When it comes to glucose

340 media the acetate is upstream in production and subsequently lactate is also increased. But in case of LB media

the input is predominantly consists of nitrogen sources and hence the carbon 'system' in turn is downstream. The

enzymatic journey taken for production of acetate is lesser in case of glucose medium than LB media. Along

with this principle, it is important to consider the limiting effects of nitrogen for production of non-nitrogenous

344 metabolites. This, along with an understanding of the anaerobic *E. coli* biochemistry, helps us in devising a 345 strategy of N-swap coupled with partial aerobic flow through the fermentation process to maximize

345 strategy of N-swap coupled with partial aerobic flow through the fermentation process to maximize 346 isobutanol/shikimate production. It is also important to note that the POSYBAL simulations were don

isobutanol/shikimate production. It is also important to note that the POSYBAL simulations were done with
 BW2511 strain where as with the exception of initial experimentation, the BL21 strain was used for producing

BW2511 strain where as with the exception of initial experimentation, the BL21 strain was used for producing
 shikimate and isobutanol. However, these strains have near identical genes and pathways and hence, the

349 simulations stick to the in vitro repertoire.

350

351 The results from the insilico simulations were tested in the lab by creating a triple knockout

 $\Delta ackA: \Delta adhE: \Delta ldhA using p1 transduction method. Additionally, ketoisovalerate decarboxylase (kivD) was$

introduced to convert ortho-isovalerate to isobutanal. Minimal media was utilized to convert glucose (carbon

source) to isobutanol in shake flasks. Initial experiments showed higher titre of Isobutanol yield in media devoid

of Nitrogen (Figure 4). Whereas, in Shikimate, ΔaroA:ΔaroK:ΔaroL knockouts fail to survive as an external

source of aromatic amino acids is required for continued survival of the cells.

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The validation of computer predictions showed an increased flux of isobutanol away from ethanol production as
seen in Figure 7 and 8. All the bioconversions and knockouts were done in BL21 strain. The overall yield of
shikimate increased from ~50 to 100 ppm in BL21 strain compared to the BW25113 (See supplementary figure
S2) also, aroK and aroL knockouts of BW and BL21 showed the same 'signature' of higher yield and BL21 was
used to make ΔaroA:ΔaroK:ΔaroL triple Knockout. The optimal biomass to isobutanol reduction ratio is

363 observed in 3% nitrogen in minimal media. Similarly, higher yield of shikimate was seen in the predicted

 $\Delta aroA:\Delta aroK:\Delta aroL$ triple knockout compared to the double knockout ($\Delta aroK:\Delta aroL$) and wild type (Figure 9).

365

366 Based on the threshold parameters as mentioned and selected earlier by POSYBAL simulations, it was found 367 that the triple knockout of $\Delta adh E: \Delta ack A: \Delta ldh A$ triple knockout corresponds to 2770 solutions of the 100000 368 (Figure 5a) solutions obtained, which is basically 2.7% of the solution set. Similarly, for shikimate 300 solutions 369 were obtained of which five solutions (Figure 5b) were seen as optimal (16%). We see that there is no central 370 governing systemic intelligence to a collection of the reaction set that has a small section (probability) wherein 371 the system produces the metabolite of interest (in this case isobutanol) and this probability increases when the 372 stem is reengineered (in case of shikimate) wherein certain gene expressions are blocked (knocked out). A filter 373 to locate solutions that have reduced flux through specific reactions below a threshold finally points to an 374 optimal knockout.

376 Conclusion

377 The in-silico platform for various species of bacteria like E. coli, M. tuberculosis, P. aeruginosa, C. 378 acetobutylicum have been described previously. They are either based on the non-linearity of interconnected 379 ordinary differential equations (ODE) that represent various enzymes which describe the cellular interactions 380 with its intrinsic kinetic parameters or in a Flux based mode where stoichiometry of each metabolite is linearly 381 connected to another of the ensemble. Both these in silico modes essentially describes the functionality of a 382 single cell and assumes homogeneity of behaviour in a population of an isogenic bacteria. In reality, a 383 population of bacteria is not only asynchronous in its physiological state, no two cells in a population are in 384 metabolic congruence. In conventional FBA the optimal solutions derived out of maximization/minimization of 385 a particular reaction gives an understanding of the system required to achieve a theoretical maximum/minimum 386 in a utopian environment. A living cell can be visualized as an ensemble of underdetermined equations that 387 connects metabolites which produces an infinite array of possible solutions. Each cell in the population may use 388 one set of these solutions. This tacitly entails that no two cells have identical expression levels in a given 389 environment. These varying behavioural signatures enables the system to be robust enough to handle stress 390 factors such as nutritional deficiency, osmotic imbalance, temperature shock or presence of antibiotics. These 391 kind of competitive growth "advantage-disadvantage" simulation can be generated in our POSYBAL population 392 model. It is seen that even in media with optimum nutrient availability cells diverge in their growth rate and 393 rapidly move away from being synchronous. This divergence that is seen experimentally and is a natural 394 outcome of our POSYBAL platform. It is also seen that this divergence of synchronicity is dependent on the 395 flux through some key non-essential pathways wherein specific knockouts produce altered divergence.

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Figures

Figure 1: Depiction of ODE, FBA and POSYBAL models. A. The ODE model takes in the kinetic parameters and gives solutions based on those parameters. The solution is singular (green) however, gene lethality can also be found (red) for a given knockout. B. The FBA model uses stoichiometric constraints and gives a single optimal solution for the entire population. The essentiality/lethality is often found through literature or experimental evidence. C. The POSYBAL model may not predict essentiality but it considers multiple solutions for each stochiometric constraint and gives multiple combination of flux the best knockouts are chosen through the population. D. Concept behind constraint-based modelling where a solution space is limited given the 'dimensions' such as stochiometric flux, metabolite produced and biomass



Figure 2: Pathway for Isobutanol and Shikimate production. The genes coloured blue is native to E. coli whereas the genes labelled in red are heterologously expressed and metabolites in green are products of interest.







■Without Nitrogen (0%) ■With Nitrogen (100%)

Figure 4: The figure represents the scatter plot of the population distribution for normalized isobutanol/shikimate production vs biomass A. Scatter plot of the population distribution for isobutanol production vs biomass comprising the triple knockouts adhE, ackA and ldhE with the threshold considered (encircled). B. Scatter plot for shikimate production comprising of the triple knockout aroK, aroA and aroL which are one of the solutions (encircled) in the plot.



Figure 5: POSYBAL simulations for isobutanol production with the predicted knockouts. A. The population of bacterial cells producing isobutanol increases as flux decreases across adhE, ackA and ldhA genes. B. Graph depicting the normalized HPLC peaks for understanding the flux of Ethanol, acetate, lactate and Isobutanol in Wild type BL21 and Δ ackA: Δ adhE: Δ ldhA triple knockout expressing KivD. There is considerable reduction of flux towards acetate, lactate and acetate production compared to Isobutanol.



Figure 6: POSYBAL simulation for minimal requirements necessary in shikimate production. A: Scatter plot representing fluxes through each gene and the corresponding shikimate produced. It is observed that despite intuitively choosing ackA as a knockout for producing more shikimate, but the in vitro output proved otherwise. Through POSYBAL platform it is observed that poxB, pykA/F and aroL can be knocked out to produce high amount of shikimate whereas an 'intermediate' flux through ackA and ptsG produces higher shikimate than its knockout. B: Normalized invitro graphs show that acetate production is required for the flux to move towards shikimate production. Also, glucose cannot be consumed when a nitrogen source (LB) isn't available.



Figure 7: A. Production of Shikimate (mg/L or PPM) in BL21, ΔaroK:ΔaroL, ΔaroA:ΔaroK:ΔaroL, ΔaroA:ΔaroK:ΔaroL with pntAB and tktA overexpression in LB/5 or 20% LB (nitrogen source) in bioreactor. B. Production of Isobutanol in 0% and 3% nitrogen after 24 hours in pilot scale fermenter.

