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# Genome-scale metabolic modelling identifies reactions mediated by SNP-SNP interactions associated with yeast sporulation

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Srijith Sasikumar 🕲 <sup>1,2,3</sup>, S Pavan Kumar<sup>2,3,4</sup>, Nirav Pravinbhai Bhatt<sup>2,3,4,5</sup> & Himanshu Sinha 🕲 <sup>1,2,3</sup> 🖂

Genome-scale metabolic models (GEMs) are powerful tools used to understand the functional effects of genetic variants. However, the impact of single nucleotide polymorphisms (SNPs) in transcription factors and their interactions on metabolic fluxes remains largely unexplored. Using gene expression data from a yeast allele replacement panel grown during sporulation, we constructed co-expression networks and SNP-specific GEMs. Analysis of co-expression networks revealed that during sporulation, SNP-SNP interactions impact the connectivity of metabolic regulators involved in glycolysis, steroid and histidine biosynthesis, and amino acid metabolism. Further, genome-scale differential flux analysis identified reactions within six major metabolic pathways associated with sporulation efficiency variation. Notably, autophagy was predicted to act as a pentose pathway-dependent compensatory mechanism supplying critical precursors like nucleotides and amino acids, enhancing sporulation. Our study highlights how transcription factor polymorphisms interact to shape metabolic pathways in yeast, offering insights into genetic variants associated with metabolic traits in genome-wide association studies.

The phenotypic characteristics of an organism are influenced by metabolic reprogramming, which modifies metabolic pathways to support various cellular functions and adapt to environmental changes. Genome-scale metabolic models (GEMs) that connect metabolic genes, proteins, and metabolites based on gene-protein-reaction associations are practical systems biology tools for describing cellular metabolism<sup>1</sup>. GEMs are an in silico tool for studying metabolic reaction networks, metabolic engineering, enzyme function prediction, drug discovery, and understanding human diseases<sup>2-4</sup>. Recent algorithm advancements have led to more precise GEMs that accurately reflect the metabolism of specific cell types or tissues. While GEMs encompass all reactions in an organism, not all enzymes are active in every cell line or tissue. Context-specific models, a subset of the GEMs, capture the metabolism of individual cell types or tissues (also known as tissue-specific models) by removing inactive reactions based on factors such as gene expression levels, the presence of proteins or metabolites, experimental data, literature knowledge, or predefined metabolic functions specific to the cell type<sup>5-9</sup>. Therefore, context-specific models provide a more accurate representation of metabolism for particular cell types or tissues.

Recently, GEMs have gained traction in exploring genotypephenotype relationships. For example, Scott et al.<sup>10</sup> demonstrated how intracellular flux variation in specific reactions influences aroma production among different vineyard yeast strains by constraining yeast GEM with exometabolomics and leveraging the flux sampling approach. Similarly, Jenior et al.<sup>11</sup> leveraged transcriptomics-integrated GEMs to compare metabolic adaptations in laboratory and clinical isolates of *Klebsiella pneumoniae*, revealing increased valine catabolism as a key feature of the pathogenic strains. Moreover, constraint-based modelling approaches are increasingly used to predict the functional effects of genetic variants on metabolism, thereby providing deeper insights into variant-to-function relationships<sup>12</sup>. Øyås et al.<sup>13</sup> developed a methodology that uses structural sensitivity analysis to model the impact of nonsynonymous SNPs on metabolic networks. Focusing on deleterious mutations and their downstream effects on reaction

<sup>1</sup>Systems Genetics Lab, Department of Biotechnology, Bhupat and Jyoti Mehta School of Biosciences, Indian Institute of Technology Madras, Chennai, Tamil Nadu, India. <sup>2</sup>Centre for Integrative Biology and Systems Medicine (IBSE), Indian Institute of Technology Madras, Chennai, Tamil Nadu, India. <sup>3</sup>Wadhwani School of Data Science and Artificial Intelligence (WSAI), Indian Institute of Technology Madras, Chennai, Tamil Nadu, India. <sup>4</sup>BioSystems Engineering and Control (BiSECt) Lab, Department of Biotechnology, Bhupat and Jyoti Mehta School of Biosciences, Indian Institute of Technology Madras, Chennai, Tamil Nadu, India. <sup>5</sup>Department of Data Science and Artificial Intelligence, Indian Institute of Technology Madras, Chennai, Tamil Nadu, India. fluxes, they applied this method to 18 *Mycobacterium tuberculosis* strains, identifying functional SNPs that impact metabolic behaviour. Similarly, Sarkar and Maranas<sup>14</sup> introduced the SNP-effect method. This technique models the impact of SNPs in enzyme-coding genes by constraining reaction fluxes based on steady-state assumptions and relative growth rates across genotypes. This approach provided a framework for predicting the phenotypic consequences of SNPs and understanding how they alter metabolic fluxes across different genetic backgrounds.

While these studies primarily focused on the cis effects of enzymecoding regions, the potential trans effects of SNPs in both coding and noncoding regions of transcription factors on reshaping the metabolic landscape remain largely unexplored. In this study, we test how SNPs in transcription factors have an impact on the metabolic landscape of a cell. We studied four naturally occurring SNPs causal for high sporulation efficiency in yeast. Sporulation efficiency, the percentage of cells sporulating in a culture in yeast, is a quantitative trait and is highly heritable across yeast strains. Sporulation efficiency is extensively studied and serves as a model for understanding complex traits<sup>15-19</sup>. During sporulation, cells undergo meiosis, where the parent diploid cell divides into four haploid spores. Several metabolic changes happen during the entire course of sporulation orchestrated by five major metabolic pathways-glutamate synthesis, tricarboxylic acid cycle, glyoxylate cycle, gluconeogenesis, and glycogenolysis<sup>20</sup>. Studies have identified several SNPs and their genetic interactions as causal for sporulation efficiency variation<sup>16,17,19,21,22</sup>. Since sporulation efficiency is a metabolically driven process, it is crucial to investigate how SNPs, individually and in combination, modulate coexpression patterns and alter the connectivity of metabolic regulators, as well as how these interactions affect metabolic flux distribution.

Our study aimed to investigate how, individually and in combination, SNPs affect differential expression and co-expression patterns of metabolic regulators, as well as the distribution of intracellular metabolic flux. We explored whether interactions between SNPs influence the distribution of intracellular metabolic flux. We used gene expression data from a previously published allele replacement panel to create context-specific genome-scale metabolic models. The goal was to investigate the interactions between SNPs and their impact on sporulation efficiency variation. While metabolic flux analysis and network-based approaches have previously been used to study the molecular mechanisms underlying specific traits or diseases, our study represents one of the few attempts to integrate gene expression data from an allele replacement panel with context-specific GEMs to explore the molecular basis of SNP interactions.

#### Results

#### Molecular model of sporulation efficiency under study

The previous studies dissected genetic variants causal for differences in sporulation efficiency in a segregant population of a cross between two yeast strains, namely, a natural oak isolate (100% sporulation efficiency) and a vineyard strain (3.5% sporulation efficiency)<sup>17,21,22</sup>. The studies identified four causal quantitative trait nucleotides (SNPs) responsible for high sporulation efficiency by performing linkage mapping<sup>17,21,22</sup>. The four causal SNPs (oak alleles) were in three sporulation genes: IME1 (Initiator of Meiosis 1), RME1 (Repressor of Meiosis 1) and RSF1 (Respiratory Factor 1). There were two alleles of IME1, namely coding IME1<sup>L325M</sup> (denoted as IME1) and a non-coding *IME1nc*<sup>A-548G</sup> (denoted as *IME1nc*), and a non-coding polymorphism *RME1*<sup>indel-308A</sup> (denoted as *RME1nc*) and a coding polymorphism in RSF1<sup>D181G</sup> (denoted as RSF1, Fig. 1A). All three genes are known transcription factors involved in the sporulation process. The IME1<sup>L325M</sup> impacts the functional equilibrium of the Ime1 protein, influencing its capacity to regulate early meiotic genes as a trans-eQTL<sup>22</sup>. The IME1nc<sup>A-548G</sup>, located in a potential Rme1 binding site within the promoter region, affects the affinity of Rme1 for IME1, modulating the strength of IME1 repression. This interaction adjusts the initiation of sporulation by altering the transcriptional activity of IME1. The RME1<sup>indel-308A</sup> acts as a cis-eQTL, impacting the RME1 expression and protein concentration, which influences its repressive activity on IME1<sup>22</sup>. RSF1 is involved in respiration and regulation of mitochondrial

genes, which is required for sporulation<sup>23,24</sup>. Since these three genes are involved at similar points in the sporulation regulatory network, extensive genetic interactions have been observed among them<sup>21,22</sup>.

By swapping these four corresponding alleles in the low sporulation vineyard strain (++++) with oak alleles (OOOO), the authors generated 16 isogenic strains with all possible allelic combinations (Fig. 1B). Based on the sporulation efficiency data of these 16 allele replacement strains<sup>25</sup>, we classified them as very high, high, medium, low, and very low sporulating strains (Fig. 1C).

In the same study by Sudarsan and Cohen<sup>25</sup>, gene expression data for all these 16 strains were obtained after 2 h of incubation in the sporulation medium, which is considered an early time point in the sporulation process. The study showed that the genetic variants explained more of the phenotypic variance proportion than the gene expression variation<sup>25</sup>. However, the study highlighted that these SNP interactions contributed significantly to gene expression variation, underscoring the complex interplay between SNPs and gene expression variation.

The phenotype and gene expression data from the Sudarsan and Cohen study were used to investigate how these SNPs and their combinations contribute to phenotypic variation by modulating the connectivity of metabolic regulators and intracellular metabolic fluxes. A three-step approach was employed to investigate the impact of SNP interactions on cellular function. First, specific differential gene expression patterns associated with these interactions were identified. Second, gene expression networks were constructed for each allele replacement strain to analyse changes in co-expression patterns. Finally, a context-specific GEM modelling approach was used to explore how these SNPs and their interactions influence intracellular metabolic fluxes, ultimately affecting the quantitative trait. The workflow, from preprocessing RNA-Seq data to applying GEM modelling techniques, is detailed in Fig. 1D.

### SNP interactions elicit distinct sporulation and metabolic gene expression patterns

First, we aimed to ascertain whether particular combinations of SNPs uniquely influence specific gene expression patterns. Second, we sought to determine if different SNP combinations elicit differential expression in distinct sets of genes or if a consistent core of genes was affected. The strains were termed based on the oak alleles as 'O' and vineyard as '+' in the gene order '*IME1 IME1nc RME1nc RSF1*'. To address this, we analysed differential gene expression between the wildtype vineyard strain and the allele replacement strains using DESeq2 after adjusting for the false discovery rate (<10% using the Benjamini-Hochberg method). We identified a total of 380 significantly up and down-regulated differentially expressed genes (DEGs) across 15 SNP combinations with wildtype vineyard strain (S<sup>++++</sup>) as the baseline (Fig. 2A and Supplementary Data 1).

Since we were interested in determining the number of differentially expressed sporulation-related genes between wildtype vineyard and allele replacement strains, we compared the above 380 DEGs list with 362 genes known to influence sporulation efficiency phenotype<sup>15</sup>. We identified 48 sporulation-specific DEGs across all SNP combinations, each with a different set of these genes. The strain with all four causal oak alleles (S<sup>0000</sup>), which had the highest sporulation efficiency, had the maximum number of these sporulation-specific DEGs (Figs. 2B and S1). In all strains except for the strain S<sup>+OO+</sup>, the presence of the RME1nc allele resulted in the downregulation of RME1 gene expression (Fig. S1). The RME1 gene is a repressor of sporulation and its downregulation in the presence of the RME1nc allele highlights the cis effect of this allele in enhancing sporulation efficiency in oak strains. Upregulation of the IME1 gene, whenever the RSF1 allele was in combination with either IME1 or IME1nc (Fig. S1), could be due to a transeffect of the RSF1 allele on the expression of the RIM4 gene, a major regulator of early sporulation genes and IME1 gene expression.

Interestingly, no common genes were found to be upregulated across all five high sporulating strains ( $S^{0000}$ ,  $S^{+000}$ ,  $S^{0+00}$ ,  $S^{00+0}$  and  $S^{000+}$ ) even though several genes were uniquely activated in each SNP combination strain (Fig. 2C). We found that *IME2*, a meiosis initiator gene, was activated



**Fig. 1** | **Schematic representation outlines the key steps and methodologies employed in our study. A** Overview of yeast sporulation efficiency model where four causal Oak SNPs in three genes *IME1*, *RME1* and *RSF1* contribute to sporulation efficiency variation. **B** The table represents the notation of strains and models used in the study and their respective genotypes. **C** The sporulation efficiency of each allele replacement strain (the data were from Sudarsan and Cohen<sup>25</sup>). **D** The raw transcriptomic data of 16 allele replacement panels were preprocessed and corrected for batch effects

using CombatSeq. Differentially expressed genes were obtained using DESeq2, and the critical regulators were obtained from RIF analysis and further studied for connectivity using PCIT analysis. An evaluation of model extraction and thresholding methods was performed to identify the best combination for integrating transcriptomic data into the yeast genome-scale metabolic model to reconstruct SNP-specific metabolic models. The change in intracellular fluxes between the wild type and the SNP models was studied using GS-DFA. The figure was created with BioRender.com.

and uniquely upregulated only when all four SNP were combined ( $S^{OOOO}$ ). We identified five upregulated genes, *HSP26*, *IME1*, *REC8*, *RIM4*, and *YIR016W*, common across the subset of strains ( $S^{OOOO}$ ,  $S^{OO+O}$ ,  $S^{O+OO}$ ,  $S^{O++O}$ ) with *IME1* and *RSF1* variants in combination (Fig. 2D). In particular, *REC8*, a gene that mediates sister chromatid cohesion, homologous recombination, and chromosome synapsis<sup>26</sup>, was uniquely upregulated only in this subset of strains (Fig. 2E).

To investigate which metabolic genes were differentially expressed across the 16 SNP combinations, we took 1150 from the Yeast8 metabolic model<sup>27</sup>, which had all metabolic reactions in the yeast genome. Using the Yeast8 model, we found that 164 metabolic genes were differentially expressed across all SNP combinations. We focused on identifying the metabolic genes upregulated in the strain ( $S^{0000}$ ), as these genes could be crucial for high sporulation efficiency. We found 14 upregulated genes,

including *FKS3* (1-3-beta-D-glucan synthase), involved in ascospore wall assembly formation<sup>28</sup> and were uniquely upregulated in this strain. We also identified that the expression of *SRT1* (involved in dolichol biosynthesis and linked to reduced sporulation in null mutants)<sup>29</sup>, *SPF1* (a P-type ATPase with null mutant showing sporulation defects)<sup>29</sup>, and *ATP15* (an ATPase) was upregulated whenever the *RSF1* allele was in combination with *IME1* and *IME1nc*, i.e., in S<sup>OO+O</sup> and S<sup>OOOO</sup> strains (Fig. S1B).

# The connectivity of critical metabolic regulators changes with SNP interactions

To elucidate the underlying regulatory mechanisms driving the differential expression patterns, we tested how different allelic combinations can impact the co-expression patterns of genes and regulators that control the sporulation phenotype.



Fig. 2 | Transcriptional differences between the allele replacement strains. A Number of upregulated and downregulated genes (LFC > 0.5 and LFC < -0.5) identified using DESeq2 between wildtype vs. each SNP combination. **B** The number of differentially expressed (LFC > 0.5 and LFC < 0.5) sporulation-specific and metabolic genes identified using DESeq2 between wildtype vs. each SNP. **C** Venn diagram analysis of upregulated genes in all strains with *RSF1*<sup>o</sup> and *IME1*<sup>o</sup> SNP combinations (S<sup>0000</sup>, S<sup>00+0</sup>, S<sup>0+00</sup>, S<sup>0++0</sup>) compared to wildtype vineyard

strain. **D** Venn diagram plot for upregulated genes identified using DESeq2 with LFC > 0.5 and *p* adj > 0.1 for three SNP combination strains (S<sup>+000</sup>, S<sup>0+00</sup>, S<sup>00+00</sup> and S<sup>000+)</sup> and four oak SNP combination strain (S<sup>0000</sup>) strains in comparison with the wildtype vineyard strain. **E** Heatmap representing the LFC values of *IME1*, *RIM4*, and *REC8* genes across all allele replacement strains compared to the wildtype vineyard strain. The Venn diagrams were generated using a web tool 'Venn'.

Given that the S<sup>0000</sup> strain with four causal oak SNPs had the highest sporulation efficiency, we examined the connectivity of metabolic regulators that control the expression of differentially expressed genes in this strain varied across other combinations of SNPs. For this, we applied Regulatory Impact Factor (RIF)<sup>30</sup> analysis to determine the critical transcription factors (TF) that control the expression of differentially expressed genes in the S<sup>0000</sup> strain. The RIF analysis has been used in various studies to identify the critical transcription factors that control gene expression<sup>31,32</sup>. The algorithm assigns a score to each transcription factor and classifies them as RIF-I and RIF-II metrics. In brief, the RIF-I metric classified TFs that were differentially co-expressed with highly abundant differentially expressed genes. RIF-II was used to identify the TFs with the altered ability to predict the abundance of differentially expressed genes<sup>30</sup>. Using *z*-score cutoffs of the RIF-I and RIF-II, we obtained 15 TFs as regulators of DEG expression (Supplementary Data 2). The RIF-I analysis identified *GCR1* (*z*-score = 2.40), which has the highest *z*-score value and is a transcriptional activator of glycolysis<sup>33</sup>. The top candidate transcription factor from RIF-II analysis was *BAS1* (*z*-score = 2.18), a Myb-related TF that regulates the basal and induced expression of genes of the purine and histidine biosynthesis pathways<sup>34</sup>. Of these 15 regulators, we were able to identify seven metabolic regulators: *GCR1*<sup>33</sup>, *GCR2*<sup>35</sup> in glycolysis, *ARG80*<sup>36</sup>, *LYS14*<sup>37</sup>, and *MET32*<sup>38</sup> in amino acid metabolism, *UPC2* involved in sterol biosynthesis<sup>39</sup>, and *BAS1* in histidine biosynthesis<sup>34</sup>.

Next, we explored how SNP interactions alter co-expression patterns. We constructed co-expression networks for the wildtype vineyard and allele replacement strains using the Partial Correlation and Information Theory (PCIT) algorithm, as detailed in the 'Methods' section. We chose PCIT for its ability to discern significant gene-to-gene associations by evaluating all possible gene triplets, which helps filter out indirect correlations and reduce false positives. Further, due to limited replicates (4 replicates per condition), PCIT is advantageous over other conventional correlation analyses like WGCNA as they may yield a high number of spurious associations. Our analysis covered 6307 genes, applying a stringent correlation threshold greater than |0.95|, which revealed distinct numbers of correlated pairs across the wildtype and allele-specific networks, highlighting differences in gene co-expression influenced by SNP interactions.

To investigate the co-expression patterns among key regulators, we focused on the interactions between the critical regulators identified through Regulatory Impact Factor (RIF) analysis and the differentially expressed genes between S<sup>++++</sup> and S<sup>0000</sup> strains. Specifically, we filtered co-expressed gene pairs to include only those involving the 143 differentially expressed genes and the 15 identified TFs from the RIF analysis. This approach enabled us to construct allele-specific sub-networks with varying numbers of nodes and edges (Supplementary Data 3). These allele-specific sub-networks were further analysed to understand how these TFs regulated gene expression by varying their connectivity across the SNPs and their combinations (Fig. 3A). The term 'connectivity' refers to the number of connections each node has in a network relative to the node with the highest number of connections in that network, which is a proxy for the importance of that node in the network<sup>31,32</sup>. We found that each metabolic regulator has different connectivity across SNP combinations. In particular, we could only identify that the connectivity of the UPC2 gene was higher for high and medium sporulating strains than for low sporulating strains, while other regulators exhibited varying connectivity without a pattern (Fig. 3B). This finding highlighted that the regulatory influence of transcription factors could differ based on SNP combinations, which might lead to the same phenotypic outcome but through distinct regulatory pathways or coexpression patterns. Therefore, it was essential to thoroughly understand the key metabolic shifts by studying the critical intracellular metabolic flux alterations driven by the genetic interactions between the SNPs associated with sporulation efficiency variation.

# Context-specific metabolic models for each SNP and their combinations show metabolic heterogeneity

The gene expression levels do not directly correlate with the enzyme levels. Hence, we leveraged GEMs to identify the metabolic flux alteration. As previously established, several genes and pathways undergo deregulation during yeast sporulation<sup>29,40</sup>. Furthermore, our analysis showed that the combinations of the SNPs studied here activate different genes (Fig. 2A).

Hence, extracting only the active genes and pathways from the generalised yeast metabolic model was crucial to reconstructing SNP-specific metabolic models. We generated context-specific metabolic models by integrating the gene expression data of SNPs and their combination into yeast GEM (Yeast 8.4.2)<sup>27</sup> using three model extraction methods: iMAT<sup>5</sup>, INIT<sup>9</sup>, and FASTCORE<sup>8</sup>. Briefly, iMAT (Integrative Metabolic Analysis Tool) creates context-specific models by balancing the inclusion of core reactions and the exclusion of non-core reactions. It requires the gene or protein expression data to be classified into high, moderate, and low expression categories, and it identifies a subnetwork that is abundant in high-expression reactions and has few low-expression reactions. iMAT maximises the number of reactions whose activity corresponds to their expression level, assigning non-zero flux to the active reactions and zero flux to inactive ones. FASTCORE uses an iterative linear programming (LP) approach to solving two LP problems. One maximises the number of core reactions with non-zero flux, and the other minimises the number of reactions required for functional support of core reactions. In this way, FASTCORE constructs a consistent context-specific model with all the predefined core reaction sets (highly expressed) in the given context. Other non-core reactions are added only to provide support to the core reactions. The extracted metabolic subnetwork will have no blocked reactions, ensuring all reactions are flux-consistent and have a non-zero flux distribution under the given conditions. INIT (Integrative Network Inference for Tissues) frames the model extraction problem as a mixed-integer LP optimisation problem. It requires weights to be derived from the gene expression data for all the reactions in the GEM. A positive weight corresponds to evidence for the presence of the reaction, and a negative weight corresponds to evidence for the absence of the reaction. INIT finds the optimal trade-off between the inclusion of reactions with positive weights and the exclusion of reactions with negative weights to identify the metabolic network that defines the specific context.

Since these algorithms require the identification of core reactions (or reaction weights) based on threshold and gene expression data, we employed two thresholding methods, Standep<sup>41</sup> and LocalGini<sup>42</sup>, which were more effective at capturing housekeeping reactions than other thresholding methods. Integrating the model extraction and thresholding methods resulted in six combinations, namely, LocalGini-iMAT, LocalGini-INIT, LocalGini-FASTCORE, Standep-iMAT, Standep-INIT and Standep-FASTCORE, to extract SNP-specific metabolic models. The number of reactions extracted by each of the six combinations is shown in Fig. 4A. The three combinations, namely, LocalGini-iMAT, LocalGini-FASTCORE, and Standep-FASTCORE, effectively captured most of the 56 meiosis-specific reactions mentioned in Ray et al.<sup>20</sup> (Fig. 4B). Further, through a comparative enrichment analysis using a hypergeometric test, we found that models derived with the LocalGini-iMAT combination exhibited higher enrichment for meiosis-specific reactions (Fig. 4C), making LocalGini-iMAT more specific compared to LocalGini-FASTCORE. Therefore, we proceeded with the SNP-specific models extracted using the LocalGini-iMAT combination to gain further biological insights into how genetic interactions modulate intracellular metabolic fluxes to drive sporulation efficiency variation.

SNP-specific models extracted from LocalGini-iMAT with the number of genes, reactions, and metabolites present in each model are shown in Table 1. We termed the M<sup>++++</sup> model generated from the wildtype vineyard strain the 'null model'. The other SNP-specific models were termed based on the oak alleles 'O' and vineyard as '+' in the gene order 'IME1 IME1nc RME1nc RSF1' (Table 1). To comprehensively analyse the similarities and differences among the SNP-specific models generated by LocalGini-iMAT, we quantified their metabolic heterogeneity and identified a core set of shared reactions. The metabolic heterogeneity among the SNP-specific models was captured by calculating the Jaccard distance between each model (Fig. S2). We identified 401 common reactions across all the extracted models. We hypothesised that these reactions form the core meiosis network, which was crucial for the survival and sporulation of yeast strains in an acetate medium. As expected, these reactions were significantly enriched with pathways that play essential roles in sporulation, which include carbon, glyoxylate and dicarboxylate, glycerolipid and glycerophospholipid metabolism, pentose phosphate pathway, and amino acids biosynthesis (Fig. S3).

# SNPs activate unique subsystems associated with sporulation efficiency variation

Since each SNP-specific model had different metabolic reactions, we hypothesised that the observed sporulation efficiency variation could be either due to the activation of new reactions in each model or variations in intracellular flux within the meiosis core network. Using the M<sup>++++</sup> null model as a baseline, a subnetwork topology analysis of each SNP model was used to investigate the contribution of each SNP combination to the sporulation efficiency variation. This analysis revealed that each SNP model had uniquely enriched subsystems, a set of reactions involved sharing a common function obtained using the flux enrichment analysis function (FEA.m) provided in the COBRA toolbox (Supplementary Data 4). For example, the model with RSF1nc SNP (M<sup>+++0</sup>) showed enrichment of valine, leucine, and isoleucine degradation pathways, glycolysis, gluconeogenesis, pyruvate metabolism pathways, and the TCA cycle (Fig. 5A). Using the subnetwork topology analysis, we found metabolic pathways uniquely enriched for respiration in the  $M^{+++0}$  model (Fig. 5Å), consistent with the previous reports highlighting the role of the RSF1 gene in respiration during

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the  $M^{++O+}$  model with *RME1nc* SNP (Fig. 5A).

0.36

0.91

0.36

0.80

0.57

0.38

0.76

0.71

0.72

0.78

0.62

0.53

0.45

0.68

0.79

LYS14

1.0

Α

В





**Fig. 3 | Analysis of connectivity of critical metabolic regulators. A** Heatmap showing the Connectivity of critical metabolic regulators identified using RIF analysis across combinations of SNPs. **B** Boxplot showing the connectivity of various metabolic regulators across three sporulation levels: High, Medium, and Low. Each box represents the distribution of connectivity values for individual regulators

sporulation. Furthermore, purine metabolism is uniquely enriched only in

the interactions between Rim11 and Ume6 and is crucial for initiating the

sporulation<sup>22,43</sup>. The study by Sudarsanam and Cohen<sup>25</sup> revealed that IME1

coding and non-coding variants contribute to sporulation efficiency varia-

tion, with the coding variant explaining a more significant proportion of the

The *IME1* coding variant affects the equilibrium between the functional and non-functional forms of the Ime1 protein. *IME1nc* variant affects within each sporulation category, with jittered points indicating individual data points. A Kruskal-Wallis test was used to assess significant differences in connectivity across sporulation levels 'ns' denotes non-significant differences, while '\*' indicates significant differences.

phenotype. Hence, we wanted to test how these variants in coding and noncoding regions of the same protein influence the metabolic fluxes. A comparison of SNP-specific metabolic models, *IME1* ( $M^{O+++}$ ) and *IME1nc* ( $M^{+O++}$ ), revealed significant differences in the metabolic activity of these two variants. Enrichment analysis of active reactions indicated that the *IME1* coding variant activated autophagy, one carbon pool by folate, phagosome, glycerolipid metabolism, and propionate metabolism pathways (Fig. S4A). In contrast, the *IME1nc* did not activate any of these metabolic



Fig. 4 | LocalGini-iMAT shows higher enrichment for meiosis-specific reactions.
A Fraction of reactions extracted from each MEM and thresholding method.
B Fraction of meiosis-specific reactions extracted from each MEM and thresholding

method. C Box plot representing the p values obtained using a hypergeometric test for models enriched for meiosis-specific reactions.

pathways, which were activated in the  $M^{O+++}$  model. As the model,  $M^{+O++}$  did not have an *IME1* coding variant necessary to form a functional protein; only the non-functional form of Ime1 protein would have been expressed. This explains the observed differences between the models  $M^{O+++}$  and  $M^{+O++}$ , suggesting a link between the metabolic plasticity of Ime1 protein functional and non-functional forms.

Further, we investigated whether, in comparison to their individual effects, interactions between SNPs had the potential to selectively activate or deactivate particular cellular pathways. In other words, we investigated whether these interactions could result in synergistic or antagonistic effects on specific metabolic reactions. To test this idea, we examined the combination of two coding SNPs, IME1 RSF1 (M<sup>O++O</sup>) and two non-coding SNPs, IME1nc RME1nc (M<sup>+00+</sup>), as well as their single SNP models, namely RSF1 (M<sup>+++O</sup>), *IME1* (M<sup>O+++</sup>), *RME1nc* (M<sup>++O+</sup>), and *IME1nc* (M<sup>+O++</sup>). By comparing reactions that were activated exclusively in the IME1 RSF1  $(M^{O^++O})$  model but not in the RSF1  $(M^{+++O})$ , IME1  $(M^{O^+++})$ , and  $M^{++++}$ null model, we identified 113 reactions. These reactions were notably enriched in purine and pyrimidine metabolism pathways (Fig. 5B and Supplementary Data 5). Conversely, when comparing reactions that were inactive in the  $M^{O++O}$  model but active in the  $M^{+++O}$ ,  $M^{O+++}$ , and  $M^{++++}$ models, we determined 271 reactions. These reactions were enriched in various subsystems, including fatty acid biosynthesis, lipid metabolism, the

phosphatidyl inositol signalling pathway, steroid biosynthesis, etc. (Supplementary Data 5 and Fig. 5C). Similarly, when comparing the *RME1nc*  $(M^{++0+})$ , *IME1nc*  $(M^{+0++})$ , and *IME1nc RME1nc*  $(M^{+00+})$  models, we uncovered the combined role of *RME1nc* and *IME1nc* in synergistically modulating the one-carbon pool by folate subsystems, encompassing the glycine cleavage complex and tetrahydrofolate aminomethyl transferase (Fig. S5 and Supplementary Data 6).

#### The impact of SNP-SNP interactions on intracellular flux variation

The analysis of the unique sub-network topology showed that combinations of SNPs could activate shared metabolic pathways. However, it was unclear whether SNPs could modify the flux distribution patterns of these pathways to modulate phenotypic variation, i.e., whether a pathway was deregulated in response to SNP-SNP interactions. Therefore, flux variations in common reactions among SNP and null models were analysed to investigate flux distribution patterns.

While the flux balance analysis (FBA)<sup>44</sup> for a given constraint can find multiple potential flux values for reactions in a metabolic model, it cannot ensure the uniqueness of a solution. To overcome this limitation in FBA, we used a flux sampling approach to assess variations in metabolic flux distribution using the genome-scale differential flux analysis (GS-DFA)<sup>45</sup>. Briefly, we first sampled the whole solution space of each reaction in the

Table 1 | Number of reactions, metabolites, and genes in each SNP-specific model generated using LocalGini-iMAT thresholding

Models	Genes	Reactions	Metabolites	Genotypes
$M^{++++}$	596	1728	1231	IME1 <sup>v</sup> IME1nc <sup>v</sup> RME1 <sup>v</sup> RSF1 <sup>v</sup>
$M^{O+++}$	571	1720	1219	IME1 <sup>o</sup> IME1nc <sup>v</sup> RME1 <sup>v</sup> RSF1 <sup>v</sup>
$M^{+O++}$	504	1247	1012	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>v</sup> RSF1 <sup>v</sup>
$M^{++O+}$	565	1644	1155	IME1 <sup>v</sup> IME1nc <sup>v</sup> RME1 <sup>o</sup> RSF1 <sup>v</sup>
$M^{+++O}$	591	1731	1237	IME1 <sup>v</sup> IME1nc <sup>v</sup> RME1 <sup>v</sup> RSF1 <sup>o</sup>
$M^{OO++}$	579	1717	1222	IME1°IME1nc°RME1 <sup>v</sup> RSF1 <sup>v</sup>
$M^{O+O+}$	515	1560	1088	IME1°IME1nc <sup>v</sup> RME1°RSF1 <sup>v</sup>
$M^{O++O}$	570	1604	1139	IME1 <sup>o</sup> IME1nc <sup>v</sup> RME1 <sup>v</sup> RSF1 <sup>o</sup>
$M^{+OO+}$	602	1750	1247	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>o</sup> RSF1 <sup>v</sup>
$M^{+O+O}$	595	1742	1238	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>v</sup> RSF1 <sup>o</sup>
$M^{++00}$	594	1733	1233	IME1 <sup>v</sup> IME1nc <sup>v</sup> RME1 <sup>o</sup> RSF1 <sup>o</sup>
$M^{+000}$	557	1737	1242	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>o</sup> RSF1 <sup>o</sup>
$M^{O+OO}$	569	1696	1209	IME1°IME1nc <sup>v</sup> RME1°RSF1°
$M^{OO+O}$	595	1745	1246	IME1°IME1nc°RME1 <sup>v</sup> RSF1°
M <sup>000+</sup>	598	1737	1240	IME1°IME1nc°RME1°RSF1 <sup>v</sup>
M <sup>0000</sup>	603	1726	1227	IME1°IME1nc°RME1°RSF1°

'O' denotes the oak allele, and 'V' indicates the vineyard allele.

model using optGpSampler, a random flux sampling method. We chose optGpSampler because of its advantages of faster convergence and reduced run time compared to traditional samplers like the GP sampler and ACHR, facilitating efficient sampling of large solution spaces. This allowed us to generate 10,000 flux solutions for each model, which spans the whole flux space (see 'Methods' for details). While nucleotide synthesis could be considered an objective during the early stages of sporulation, the progression of meiosis can vary between strains due to SNP effects. For example, a strain with a single SNP may not be undergoing active meiosis at the same time as another strain with four SNPs that is progressing rapidly. Therefore, selecting a single objective function to predict flux distributions would not adequately capture the dynamic and SNP-dependent variations in sporulation. Consequently, we performed flux sampling without an objective function to bypass these limitations and more accurately represent the metabolic fluxes influenced by SNPs across different strains.

Next, we employed GS-DFA for its capability to detect significant flux dysregulation by comparing flux distributions between SNPspecific models and the null model ( $M^{++++}$ ) using a stringent statistical test and flux change cutoff (see methods). This approach provided a robust framework for identifying reactions significantly modulated by specific SNPs and their combinations. We identified upregulated (Supplementary Data 7) and downregulated (Supplementary Data 8) reactions in each SNP model compared to the null model. GS-DFA analysis showed that the upregulated reactions were primarily enriched in the six subsystems involved in the biosynthesis of amino acids, antibiotics, secondary metabolites, glycerophospholipid metabolism, glycerolipid metabolism, and pentose phosphate pathway in the strain with the four oak allele combination  $S^{0000}$  ( $M^{00000}$ , Fig. 6A and Supplementary Data 9).

As  $M^{0000}$  model created from strain  $S^{0000}$  had the highest sporulation efficiency, we checked if these six subsystems were present in the other high sporulation efficiency strain models:  $M^{+000}$ ,  $M^{0+00}$ ,  $M^{00+0}$  and  $M^{0000}$ . We found that these models contained different subsets of the six subsystems observed in the  $M^{0000}$  model, emphasising the role of intracellular flux variations in these pathways as significant contributors to the observed differences in sporulation efficiency across the strains (Fig. 6A). The two subsystems, glycerophospholipid and glycerolipid metabolism, present in all models suggested that they were essential for sporulation. Further, we explored how the functional (with the *IME1* coding variant) and non-functional forms (without the *IME1* coding variant) of the Ime1 protein could change the flux distribution patterns to modulate sporulation efficiency variation. From the GS-DFA analysis, we found that reactions involved in glycerolipid, glycerophospholipid metabolism, and biosynthesis of secondary metabolites were upregulated only in *IME1 IME1nc* ( $M^{OO+++}$ ) and *IME1* ( $M^{O+++}$ ) models but not in *IME1nc* ( $M^{+O++}$ ) model (Fig. S4B). Interestingly, we could also predict the metabolic consequence of the non-functional form of *IME1* (*IME1nc*,  $M^{+O++}$ ), which upregulated the reactions involved in the peroxisome, fatty acid metabolism, fatty acid degradation, and fatty acid biosynthesis (Fig. S4B).

Upon hierarchically clustering the flux change (FC) values of upregulated reactions in the M<sup>0000</sup> model across other models, we found that models were clustered into two groups (Fig. S6). The presence or absence of the IME1nc variant primarily determined this clustering. Further, we found that SNP combinations without the IME1nc variant, i.e., M<sup>O+OO</sup>, M<sup>O++O</sup>, M<sup>0+0+</sup>, M<sup>++00</sup>, M<sup>++0+</sup>, M<sup>0+++</sup>, and M<sup>0000</sup> models led to an increased flux through reactions involved in the pentose phosphate pathway (PPP, Table 2 and Supplementary Data 9). This result was consistent with the flux balance analysis performed using nucleotide biosynthesis as the objective function and flux variability analysis<sup>46</sup>, where we found that the PPP reactions were upregulated in these models (Fig. S7 and Supplementary Data 10). The increased flux through PPP reactions was inversely coupled with the glucose-6-phosphate isomerase reaction. This result indicated that the flux was redirected through the PPP for biosynthesis rather than glycolysis for energy generation (Fig. 6B). In highly sporulating SNP-specific models M<sup>00+0</sup>, M<sup>000+</sup>, and M<sup>+000</sup> that had *IME1nc* variant and where the PPP was not upregulated (Fig. 6A, B), showed enrichment of reactions involved in autophagy and phagosomes (Fig. 6C). These findings were interesting since previous studies have established a correlation between autophagy and yeast sporulation, and the deletion of several autophagy-related genes has shown reduced sporulation<sup>47,48</sup>. These findings demonstrated the significant impact of SNPs and their interactions on the metabolic flux distribution, explaining how these interactions affected sporulation efficiency.

#### Discussion

By analysing the gene expression data from an allele replacement panel of four SNPs, we showed how SNPs interacted and modulated the metabolic pathways associated with sporulation efficiency variation. We showed a few cases where specific SNP combinations could synergistically or antagonistically modulate certain metabolic reactions. We identified that nucleotide metabolism (purine and pyrimidine metabolism) was synergistically modulated, while steroid biosynthesis was antagonistically modulated when coding variants of *RSF1* and *IME1* were combined. These insights gave us a better understanding of how SNP interactions could redirect metabolic flux, prioritising nucleotide synthesis over steroid synthesis in the early stages of meiotic progression during sporulation. Previous studies have underscored the pivotal role of pyrimidine metabolism in facilitating efficient meiotic progression<sup>49</sup>. Further, we showed the SNP-specific modulation of the glycine cleavage system, where the SNPs of *RME1nc* and *IME1nc* were synergistic.

We characterised the intracellular metabolic flux states of each SNPspecific model<sup>10,45</sup>. Using GS-DFA analysis, we identified flux variation in six major pathways as significant contributors to sporulation efficiency variation. Interestingly, our GS-DFA analysis did not identify the fundamental reactions that were known to constitute the meiosis-specific metabolic model<sup>20</sup>. These included the TCA cycle, glyoxylate cycle, and acetate uptake, which have been shown to influence sporulation at an early stage through gene-knockout approaches. However, we captured these pathways in our core meiosis-specific network by comparing the common reactions shared among SNP-specific models. These findings suggested that SNPs and their interactions modulate sporulation efficiency variation by influencing metabolic reactions beyond the essential ones required for sporulation. This observation highlighted the specific effects of SNP-level variations, revealing regulatory influences often ignored in gene knockout studies in metabolic



### B Synergistic effect of *IME1*c and *RSF1c* variants on nucleotide metabolism



#### Models

### C Antagonistic effect of *IME1*c and *RSF1c* variants on Steroid metabolism



**Fig. 5** | **SNPs activate unique subsystems associated with sporulation efficiency variation. A** Metabolic subsystems enrichment analysis for reactions that are overrepresented in each comparison that are present exclusively in single SNP models compared with the wildtype model. The size of the dots represents the reaction ratio, while the colour intensity indicates the adjusted *p* value obtained using the hypergeometric test. **B** Synergistic effect of *RSF1*<sup>o</sup> and *IME1*<sup>o</sup> in modulating nucleotide metabolism. Here, the violin plots represent the flux distribution obtained by

optGpSampler for representative key reactions involved in purine and pyrimidine metabolism compared across null (M<sup>++++</sup>), *RSF1<sup>0</sup>IME1*<sup>o</sup> (M<sup>O++O</sup>), *RSF1*<sup>o</sup> (M<sup>+++O</sup>) and *IME1*<sup>o</sup> (M<sup>O++++</sup>) models. C Antagonistic effect of *RSF1*<sup>o</sup> and *IME1*<sup>o</sup> in modulating Steroid metabolism. Here, the violin plots represent the flux distribution obtained by optGpSampler for representative key reactions involved in Steroid biosynthesis compared across null (M<sup>++++</sup>), *RSF1*<sup>o</sup>*IME1*<sup>o</sup> (M<sup>O++O</sup>), *RSF1*<sup>o</sup> (M<sup>+++O</sup>) and *IME1*<sup>o</sup> (M<sup>O+++</sup>) models.



Proposed pentose phosphate pathway dependent compensatory mechanism

Fig. 6 | Pentose phosphate pathway is differentially regulated in SNP models. A Flux enrichment plots for upregulated reactions in three and four SNP combination models ( $M^{+000}$ ,  $M^{0+00}$ ,  $M^{000+0}$ ,  $M^{000+}$  and  $M^{0000}$ ) compared to the null ( $M^{++++}$ ) model obtained after GS-DFA analysis. The size of the dots represents the reaction ratio, while the colour intensity indicates the adjusted *p* value obtained using the hypergeometric test. **B** Box plots representing the flux distribution of key reactions involved in the pentose phosphate pathway obtained from optGpSampler

for three and four SNP combination models ( $M^{+000}$ ,  $M^{0+00}$ ,  $M^{000+0}$ ,  $M^{000+}$  and  $M^{0000}$ ) compared to the null ( $M^{++++}$ ) model. **C** Flux enrichment plots for unique reactions in highly sporulating strains compared to the null model ( $M^{++++}$ ). The size of the dots represents the reaction ratio, while the colour intensity indicates the adjusted *p* value obtained using the hypergeometric test. **D** Proposed mechanism of autophagy as a pentose-phosphate pathway-dependent compensatory mechanism during the early phase of yeast sporulation (created with Biorender.com).

Table 2 | Differential regulation of pentose phosphate pathway subsystem across SNP models ('O' denotes the oak allele, and 'V' indicates the vineyard allele)

PPP status	Genotype	Models
PPP similar to the null model	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>o</sup> RSF1 <sup>v</sup>	$M^{+OO+}$
(M <sup>++++</sup> ) (IMF1 <sup>v</sup> IMF1nc <sup>v</sup> RMF1 <sup>v</sup> RSF1 <sup>v</sup> )	IME1°IME1nc°RME1 <sup>v</sup> RSF1 <sup>v</sup>	$M^{OO++}$
(	IME1°IME1nc°RME1°RSF1 <sup>v</sup>	$M^{OOO+}$
	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>o</sup> RSF1 <sup>o</sup>	$M^{+000}$
	IME1°IME1nc°RME1 <sup>v</sup> RSF1°	M <sub>00+0</sub>
	IME1 <sup>v</sup> IME1nc <sup>v</sup> RME1 <sup>v</sup> RSF1 <sup>o</sup>	M+++0
	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>v</sup> RSF1 <sup>v</sup>	$M^{+O++}$
	IME1 <sup>v</sup> IME1nc <sup>o</sup> RME1 <sup>v</sup> RSF1 <sup>o</sup>	$M^{+O+O}$
PPP upregulation	IME1 <sup>v</sup> IME1nc <sup>v</sup> RME1 <sup>o</sup> RSF1 <sup>v</sup>	$M^{++O+}$
	IME1°IME1nc <sup>v</sup> RME1 <sup>v</sup> RSF1 <sup>v</sup>	$M^{O+++}$
	IME1°IME1nc <sup>v</sup> RME1°RSF1 <sup>v</sup>	$M^{O+O+}$
	IME1 <sup>v</sup> IME1nc <sup>v</sup> RME1 <sup>o</sup> RSF1 <sup>o</sup>	$M^{++00}$
	IME1°IME1nc <sup>v</sup> RME1 <sup>v</sup> RSF1°	$M^{O++O}$
	IME1°IME1nc <sup>v</sup> RME1°RSF1°	M <sup>0+00</sup>
	IME1°IME1nc°RME1°RSF1°	M <sup>0000</sup>

models, which focus on complete pathway loss. Our analysis also predicted the differential regulation of the pentose phosphate pathway in an IME1ncspecific manner. The pentose phosphate pathway is critical in supplying the cell with the necessary building blocks and reducing equivalents needed for the biosynthesis of nucleotides and amino acids<sup>50</sup>. Our hypothesis suggested that autophagy could produce the essential precursors for nucleotide and amino acid biosynthesis by degrading and recycling cellular components compensating for PPP, particularly in models where PPP upregulation was absent (Fig. 6D). Previous studies have shown autophagy as a compensatory mechanism for the PPP in tumour cells<sup>51</sup>. It would be interesting to experimentally validate whether autophagosomes are induced when the PPP reaction is downregulated. The reactions identified as upregulated or downregulated based on flux change do not necessarily correspond to differential gene expression. These changes can also be attributed to the posttranscriptional regulation of genes. Thus, integrating SNP and their combination-specific gene expression data in GEM provided several novel mechanisms by which phenotypic variation in a well-studied trait was regulated at the molecular level.

Genome-wide association studies (GWAS) revealed that multiple SNPs, each exerting a modest impact on a trait, contributed to the complexity of genetic effects through additive and epistatic interactions<sup>52,53</sup>. Several studies have reported complex genetic interactions between identified SNPs in metabolic diseases<sup>54,55</sup>. It is crucial to examine intermediate phenotypes such as intracellular metabolic flux to address the challenge of understanding how SNPs and their interactions influence complex traits, particularly the variant-to-function (V2F) relationship<sup>56</sup>. A recent study has used GWAS variant data to develop personalised organ-specific metabolic models for 524,615 individuals of the INTERVAL and UK Biobank cohorts to clarify the impact of genetic variations on the metabolic processes implicated in coronary artery disease<sup>57</sup>. Applying SNP-specific genomescale metabolic models that integrate transcriptomic and metabolomic data will help identify known and novel pathways perturbed by SNPs and their combinations, leading to clearer variant-to-function pathways.

#### Methods

### RNA-Seq data, preprocessing, and differential expression analysis

Raw RNA sequence read data for single SNP and their combinations was downloaded from NCBI GEO (accession number GSE55409)<sup>25</sup> for 16 isogenic allele replacement strains of the oak alleles of four SNPs *IME1*,

IME1nc, RME1nc and RSF1. In the mentioned study, cultures of the 16 allele replacement strains containing oak causal variants (described in the manuscript) in all possible combinations in the vineyard background were grown in a YPD medium and incubated for 2 h in 1% potassium acetate to initiate sporulation. Total RNA was extracted from the cell pellets on different days until four biological replicates were obtained for each strain. There were 63 samples, with each strain having four replicates, except for the strain harbouring the IME1c (S<sup>++O+</sup>) variant, which had only three replicates. The reads were converted to Fastq format using the command 'Fastqdump' in the SRA toolkit. The FastQC and MutiQC were performed for quality control<sup>58</sup>. The reads were mapped to reference Saccharomyces cerevisiae R64-1-1.105 downloaded from the Ensembl database (https://ftp. ensembl.org/pub/release110/fasta/saccharomyces\_cerevisiae/cdna/Saccha romyces\_cerevisiae.R64-1-1.cdna.all.fa.gz) using Salmon (an alignmentfree tool), accounting for GC and sequence bias<sup>59</sup>. The transcript TPM values obtained from Salmon were converted to gene-level counts using Tximport v1.4.0<sup>60</sup>. The effect of day on the gene expression values was removed using ComBatseq<sup>61</sup>.

To identify the genes that are differentially expressed between the wildtype vineyard strain and each allele replacement strain, we performed differential gene expression analysis using DESeq262. In brief, the batch-corrected untransformed raw counts were given as input to the DESeq2. After filtering for low counts with less than 100 counts in all 63 samples, 6305 genes out of 6571 were selected for further processing. The effect of the day was included in the design model of DESeq2 to account for the contribution of covariates to the total gene expression. log<sub>2</sub> fold changes (LFC) shrinkage using the alpegm method<sup>63</sup>, with a false discovery rate of 10% (p value adjusted using the Benjamini-Hochberg method), was used to identify differentially expressed genes (DEGs). Genes with LFC values greater than 0.5 and less than -0.5 are considered upregulated and down-regulated, respectively. The upregulated DEGs obtained for each allele replacement strain are given as input to ClueGO<sup>64</sup> from Cytoscape<sup>65</sup> for gene ontology and functional annotations analysis.

#### Regulatory impact factor (RIF) analysis

The CeTF package in R was utilised to perform the Regulatory Impact Factors analysis<sup>66</sup>. The Variance stabilising transformation (VST) was performed on the gene expression data of wildtype vineyard strain and strain with four oak SNPs. It was given as input for the RIF analysis as it required two contrasting conditions. The TFs, downloaded from the Yeastract database (http://www.yeastract.com)<sup>67</sup> were contrasted to the list of DEGs identified by comparing the wildtype vineyard strain (i.e., a strain with no SNPs) to the strain with all four oak SNP combinations. RIF analysis classifies TFs into two categories: RIF-I and RIF-II. RIF-I focuses on TFs consistently exhibiting the strongest differential co-expression with highly abundant and differentially expressed genes. In contrast, RIF-II identifies TFs that provide predictive evidence for changes in the abundance of DE genes. The TFs identified using the RIF analysis with scores deviating  $\pm$  1.96 SD from the mean were considered significant and termed 'critical regulators'.

#### PCIT and connectivity analysis

The Partial Correlation and Information Theory (PCIT) approach<sup>68</sup> was used to arrive at co-expression networks for wildtype vineyard strain and 14 SNP combinations except for strain with  $IME1^{O}$ , which had only three replicates. The correlation of |0.95| and based on the *p* value < 0.05 was used as a threshold for arriving at a significant association (this threshold was chosen to arrive at a maximum number of significant edges)<sup>31</sup>. Further, the DEGs (specific for wild type vs. four SNP combination strains) and critical regulators identified using the RIF analysis are filtered to arrive at sub-networks for each SNP-specific network. The individual co-expression sub-networks for wildtype vineyard strain and 14 allele replacement strains were visualised using Cytoscape<sup>65</sup> version (3.9.1). Each subnetwork was analysed for network properties using the Cytoscape plug-in NetworkAnalyser 4.4.8.

The connectivity for each critical regulator was identified by dividing the degree of that regulator in any specific network with the highest degree in that network<sup>31,32</sup>. The connectivity of each critical regulator was then compared across each SNP combination and was visualised using the Pheatmap package in R (v1.0.12, https://CRAN.R-project.org/package=pheatmap).

#### An evaluation of model extraction and thresholding methods for reconstructing SNP-specific metabolic models

Yeast 8.4.2<sup>27</sup>, a commonly used yeast GEM model, was used in this study (https://github.com/sysbiochalmers/yeast-gem). This GEM had 1150 genes, 4058 reactions, and 2742 metabolites. The bounds of exchange reactions of the GEM were changed to reflect the early stages of the sporulation state appropriately. For this, we restricted the intake of glucose and nitrogen (Lower bound = Upper bound = 0 [mmol/(g DW h)]) while allowing an unrestricted supply of oxygen and acetate (Lower bound = -1000 [mmol/(g DW h)])<sup>20</sup>.

The gene expression data (TPM values) averaged across replicates of wild type, and 15 allele replacement strains were preprocessed using our novel preprocessing algorithm LocalGini thresholding<sup>42</sup> and Standep<sup>41</sup> and integrated into Yeast 8.4.2 using the model extraction methods (MEMs), iMAT<sup>5</sup>, INIT<sup>9</sup> and FASTCORE<sup>8</sup> algorithms that did not require any objective function were used to build the models.

#### LocalGini thresholding

The LocalGini algorithm employs an inequality metric, the Gini coefficient, to identify the high and low expressed genes as defined below. Given the gene expression matrix (*G*) of size  $m \times n$ , the rows represented m genes, and the columns represented n strains. Hence, the element,  $G_{i,j} \in [0, \infty)$  was the expression value of the *i*th gene for the *j*th strain. The Gini coefficient of the gene *i* (*GC<sub>i</sub>*) was computed as:

$$GC_{i} = \frac{\sum_{k=1}^{n} \sum_{j=1}^{n} \left| G_{i,k} - G_{i,j} \right|}{2n^{2} \bar{G}_{i}}$$

where, *n* was the total number of strains, and  $\overline{G}_i$  was the mean expression of the *i*th gene across the strains. An *m*-dimensional vector of Gini coefficients, GC, was constructed. GC was further scaled to percentile values. The percentile value of the corresponding rows in the gene-expression matrix, *G*, was taken as the LocalGini threshold.

For iMAT, if a gene's expression was greater than the obtained threshold value, it was included in the high-expressed gene set (HG-set), and if a gene's expression value was lesser than the obtained value, it was included in the low-expressed gene set (LG-set). Further on, genes with expression values above the 90th percentile were included in the HG-set, and genes with expression values below the 10th percentile were included in the LG-set, regardless of their Gini coefficient. For FASTCORE, if a gene's expression was greater than the obtained threshold value, it was included in the core gene set. Further on, genes with expression values above the 90th percentile were included in the core gene set. Genes with expression values below the 10th percentile were obtained by normalising the gene expression values with respect to the thresholds. The minimum value of the reaction activity scores was set to -10.

The Gene-Protein-Reaction (GPR) rules in Yeast 8.4.2 were utilised to map the genes to the reactions. Reactions without gene evidence, such as spontaneous, diffusion, and transport reactions, as well as reactions without GPR relation, were assigned to the moderately expressed reaction set (MRset) in the case of iMAT, non-core reactions in the case of FASTCORE and given a score of 0 in case of INIT.

#### Preprocessing using Standep

StanDep hierarchically clusters the genes with similar expression patterns across the tissues and provides a cluster-specific threshold. Euclidean

distance and complete linkage metrics were used for hierarchical clustering. The cluster-specific threshold,  $\emptyset_c$ , for the cluster *c*, was calculated as follows:

$$\phi_{c} = (\theta_{c} - \min(\theta_{c})) * \frac{100}{\max(\theta_{c} - \min(\theta_{c}))}; \phi_{c} \in [0, 100]$$
$$\theta_{c} = f(\sigma_{c}) + g(\mu_{c})$$
$$f(\sigma_{c}) = \frac{(\sigma_{c} - \Delta)}{\max(\sigma_{c} - \Delta)}$$
$$g(\mu_{c}) = \mu_{c} - M$$

here,  $\theta_c$  was the raw threshold value for cluster c,  $\mu_c$  and,  $\sigma_c$ , which was the mean and standard deviation of the cluster, respectively. M and  $\Delta$ , were the mean and standard deviation of the entire dataset, respectively. These thresholds were applied to the enzyme expression values, i.e. only the 'AND' rule in the GPR relation was resolved.

For FASTCORE and iMAT, the core and high-expressed reaction sets included enzyme expression values greater than the threshold. For INIT scores, the distance between the threshold and the expression value was provided as scores.

The yeast cells in the sporulation medium showed an increased rate of nucleotide synthesis during the early phase<sup>20</sup>, the reaction catalysed by glucose-6-phosphate dehydrogenase ('r\_0466') being the first step of the pentose phosphate pathway. Hence, we ensured the glucose-6-phosphate dehydrogenase ('r\_0466') and non-growth associated maintenance reaction ('r\_4046') were present in all the extracted context-specific models. This resulted in the reconstruction of 15 SNP-specific models and a null model for each thresholding and MEM combination. Further, we evaluated the performance of each method for its ability to capture most of the meiosis-specific reactions<sup>20</sup> in each SNP-specific model. Then, we employed a hypergeometric test to compare the enrichment of the meiosis-specific reactions in each method. The models generated from the optimal thresholding and MEM identified in our evaluation were used for further analysis.

To understand the metabolic heterogeneity of each extracted model, we identified the Jaccard distance between each SNP-specific model. Further, we identified reactions in each SNP-specific model that were not in the null model. These reactions, active in SNP models while not in the null model, were searched for significantly enriched subsystems using the flux enrichment analysis function (*FEA.m*) provided in the COBRA toolbox<sup>69</sup>. In brief, *FEA.m* used a hypergeometric 1-sided test ( $\alpha = 0.05$ ) and FDR correction (adjusted *p* value < 0.05) to obtain significantly enriched subsystems.

# Genome-scale differential flux analysis (GS-DFA) using a flux sampling approach

To discern how each SNP and their combinations can modulate the intracellular fluxes to show sporulation efficiency variation, we leveraged the random flux sampling approach, which acquires a significant number of evenly distributed solutions over the whole solution space and does not require an objective function. We adopted a previously developed protocol<sup>45</sup> to identify the most significantly dysregulated reactions in the presence of a particular SNP and their combination by comparing the null and each SNP models. However, we modified the pipeline to use an optGpSampler<sup>70</sup> instead of an ACHR sampler. This modification was based on previous studies demonstrating the optGpSampler's advantage in faster convergence and reduced run time compared to the ACHR sampler<sup>71</sup>. The optGpSampler in COBRApy version 0.10<sup>72</sup> was used in this study to sample 10,000 flux solutions, and the thinning factor was set to 100 for all models. This was to ensure the spanning of the whole flux space and to obtain sparse and uncorrelated fluxes<sup>45</sup>. Let,  $X_{SNP}$  and  $X_{null}$ , be the flux distributions of a particular reaction in SNP-specific and null models. We identified the

significantly deregulated reactions in each SNP model compared to the null model using a two-sided Kolmogorov-Smirnov test with a significance level 0.05 (adjusted *p* value using the Benjamini-Hochberg method). Upregulated and downregulated reactions were identified from these dysregulated reactions using the flux change value obtained. The reactions with flux change greater than 0.82 were considered upregulated, and flux change less than -0.82 were considered downregulated, indicating ten times higher or lower fold change in flux in the SNP model compared to the null model.

The flux change was calculated for each reaction as follows:

$$Flux \ change = \frac{\bar{X}_{SNP} - \bar{X}_{null}}{|\bar{X}_{SNP} + \bar{X}_{null}|}$$

where,  $\bar{X}_{SNP}$  and  $\bar{X}_{null}$ , the arithmetic means of the flux distributions for a given reaction in SNP and null models, respectively.

#### Flux balance and flux variability analysis

Using the LocalGini-iMAT-derived models, we performed flux balance analysis<sup>44</sup> using nucleotide biosynthesis ('r\_0466') as objective function<sup>20</sup> and flux variability analysis<sup>46</sup> for the reactions in the null and each SNPspecific model. The function *fluxVariability.m* in the COBRA toolbox was used to get the minimum and maximum possible flux values for each reaction. We used NGAM (Non-Growth Associated Maintenance Reaction) as the objective function, with the upper bound set at 1000 and the optimal percentage value set at 2%. This ensured that the metabolic reactions required for the cell's survival but not essential for their growth were active in the models. As previously described, we calculated the flux span ratio (FSR) for each reaction that was common between the null and SNPspecific models<sup>2</sup>. FSR for a reaction was the ratio of flux ranges (max flux-min flux) of the SNP-specific model to the null model. Reactions with FSR less than 0.5 and greater than two were considered downregulated and upregulated reactions, respectively. Further, the upregulated and downregulated reactions were checked for subsystem enrichment using flux enrichment analysis as described in previous sections.

#### COBRA toolbox and COBRApy setup

COBRA Toolbox version 3.0.0 and COBRApy version 0.10.1 on MATLAB R2021b and Python version 3.5.2, respectively. The optimiser used for linear programming and mixed-integer linear programming optimisation was Gurobi, version 8.0.0. optGpSampler was run using parallel processes. We used customised Python scripts for statistical analysis and visualisation of flux sampling results.

#### Data availability

All the RNAseq data used in this paper are available from NCBI GEO accession number GSE55409. All other datasets used in the paper are mentioned in the 'Methods' section.

#### Code availability

The codes used in this paper and all generated models are available in Zenodo (https://doi.org/10.5281/zenodo.13917841)<sup>73</sup>.

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### Author contributions

Conceptualisation: S.S. and H.S.; Formal analysis: S.S., S.P.K. and N.P.B.; Investigation: S.S., S.P.K., N.P.B. and H.S.; Writing original draft: S.S. and H.S.; Supervision: H.S.; All authors reviewed and approved the final paper.

### **Competing interests**

The authors declare no competing interests.

### **Additional information**

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**Correspondence** and requests for materials should be addressed to Himanshu Sinha.

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