
UNDERSTANDING HOW GENETIC VARIATION AMONG INDIVIDUALS LEADS TO THE PHENOTYPIC DIVERSITY OBSERVED WITHIN A SPECIES IS A LONG-STANDING AND FUNDAMENTAL GOAL IN BIOLOGY. ADVANCES IN WHOLE GENOME SEQUENCING HAVE ALLOWED GENOME-WIDE ASSOCIATION STUDIES (GWAS) TO CORRELATE GENETIC VARIANTS WITH VARIOUS TRAITS, INCLUDING MANY HUMAN DISEASES. HOWEVER, WHILE THOUSANDS OF CAUSAL VARIANTS HAVE BEEN IDENTIFIED, THEY OFTEN ONLY EXPLAIN A FRACTION OF THE OBSERVED PHENOTYPIC VARIANCE. WHILE THERE ARE A NUMBER OF POSSIBLE EXPLANATIONS FOR THIS “MISSING HERITABILITY”, RARE VARIANTS, GENETIC INTERACTIONS AND STRUCTURAL VARIANTS ARE LIKELY TO BE A SIGNIFICANT CULPRIT. DURING THIS REPORTING PERIOD, WE TOOK ADVANTAGE OF LESSONS LEARNED FROM SYSTEMATIC FUNCTIONAL GENOMIC ANALYSES IN S. CEREVISIAE AND WE USE IT TO EXPLORE THE ORIGIN OF THE MISSING HERITABILITY FROM POPULATION DATA. IN PARALLEL WE BUILT NEW TOOLS THAT ALLOW US TO PERFORM LARGE SCALE FUNCTIONAL GENOMICS IN DIVERSE STRAIN BACKGROUNDS.

SPECIES-WIDE EXPLORATION OF THE GENETIC AND PHENOTYPIC LANDSCAPES

TO LAY THE FOUNDATION OF THE EXPLORATION OF THE GENETIC COMPLEXITY OF TRAITS, WE FIRST COMPLETELY SEQUENCED AND PHENOTYPED A LARGE COLLECTION OF 1,011 YEAST ISOLATES, COMING FROM DIVERSE ECOCLOGICAL NICHEs ACROSS THE WORLD ON PLATES CONTAINING DIFFERENT STRESSORS. THE GENERATED DATASET REVEALED AN UNDESCRIPTED EVOLUTIONARY HISTORY AS WELL AS THE DRIVING FORCES OF GENOME EVOLUTION, AND HAS PROVIDED INSIGHTS INTO THE GENOTYPE–PHENOTYPE RELATIONSHIP.
To dissect the genetic basis of complex traits, we developed an efficient, standardized and quantitative high-throughput screening strategy based on an automated robotic platform (Singer RoToR HDA robot) to replicate the set of strains onto various media (more than 50 growth conditions). In total, we determined and analyzed 34,956 phenotypic measurements covering a large number of traits providing a comprehensive analysis of their inheritance patterns. The combination of these two datasets allowed to perform genome-wide association studies. We performed mixed-model association and detected 35 variants significantly associated with 14 conditions, with an enrichment and high association scores for Copy Number Variants (22 CNVs vs. 13 SNPs). In addition, some of the detected variants are linked to variable ORFs, which are not present in the reference genome. Phenotypic variance explained was estimated by running a new association with a similarity matrix containing the significantly associated markers. For five of the tested traits, the phenotypic variation explained is surprisingly greater than 25%. In fact, CNVs explained larger proportions of trait variance compared to SNPs, with a median of 36.8% and 4.49% of the variance explained, respectively. Our genome-wide association analyses, including an exhaustive catalog of genome content and CNVs present in the 1,011 genomes, highlighted the overall importance of these genetic variants on the phenotypic diversity.

Interestingly, the difference between the estimated genome-wide heritability and explained phenotypic variance gives an overview of the extent of missing heritability. Many SNPs are present at low frequencies, which echoes observations previously made in human GWAS and raised the question of whether rare SNPs have an important role in modulating the phenotypic landscape.

**Rare variants contribute significantly to quantitative trait variation**

Based on the genomic and phenotypic data from the collection of 1,011 *S. cerevisiae* isolates, a total of 55 isolates that are diploid, homozygous, genetically diverse and present unbiased population structure have been selected. In total, we created 3,025 hybrids, representing 2,970 heterozygous hybrids with a unique parental combination and 55 homozygous hybrids. We screened the entire set of isolates, and hybrids for high-resolution quantification of mitotic growth ability across 53 conditions, using the automated, standardized robotic platform. The conditions included different carbon sources and chemical compounds impacting various physiological and cellular responses such as membrane and protein stability, signal transduction, sterol biosynthesis, transcription, translation as well as osmotic and oxidative stress. This phenotyping step led to the characterization of more than 160,00 hybrid/trait combinations.

Our dataset gave us the opportunity to have a species-wide view of the relative contributions of additive and non-additive components to overall phenotypic variation in our large sample of 3,025 hybrids. Separation of variance components of phenotypic variation is a key point in the understanding of genetic architecture of traits and a potential source of the missing heritability. In addition, because we tested many traits across a large population, we highlighted interesting variations between conditions as well as isolates, suggesting the presence of rare functional variants and potential expressivity cases.

Because all the 55 isolates used in the diallel cross were completely sequenced, we examined the genetic basis of traits across a large panel of traits using GWAS. The main point about using a diallel cross design was that we can use the redundancy of the haplotype which is intrinsic of the pairwise crosses to our advantage. Indeed, with only 34 base genomes, we could generate *in silico* the 595 genomes corresponding to a half pairwise matrix of 561 hybrids with 34 homozygous. Each parental genome is present 34 times hence creating haplotype mixing across the matrix. This high level of haplotype shuffling and repetition gives the advantage of offering allele overrepresentation compared to the use of a population with the same number of independent individuals. Minor allele frequency (MAF) will be fundamentally changed in a diallel compared to the species level because of the smaller number of parents involved. In our diallel panel, out of the total 31,632 SNPs retained, 3.5% (1,128) which had a MAF < 5% in the 1,011 *S. cerevisiae* genomes happen to
surpass this threshold in the diallel panel and thus are now detectable, going up to a MAF of 32%. Surprisingly, 12.1% of the significantly associated SNPs also surpassed this threshold meaning that they could not have been detected by a classical GWAS approach. Altogether, our results have major implications for our understanding of the genetic architecture of traits in the context of unexplained heritability. They clearly highlight the extensive role of low-frequency and rare variants on the phenotypic variation at the population level.

Global picture of inheritance patterns and the genetic complexity of traits

We then took advantage of the diallel panel to assess the overall genetic complexity of traits and the prevalence of phenotypic expressivity at a population-scale. In this context, we first selected a subset of the large diallel hybrid panel in order to have 190 unique hybrids coming from 20 natural isolates representative of the S. cerevisiae genetic diversity. For each of these hybrids, a large progeny of 160 individuals (corresponding to 40 full tetrads) was obtained, leading to a total of 30,400 offspring individuals. Their mitotic growth has been assessed on 40 growth conditions inducing various cellular stress.

The main objective of this work has been to infer complexity level of traits at a population scale and assess its dynamic across multiple genetic backgrounds. To do so, we conducted a large-scale phenotyping of the whole panel of 30,400 haploid progeny coming from 190 hybrids. Overall, more than three million phenotypic measurements were performed and grouped for each cross and condition (trait) to obtain 7,600 phenotypic distributions of haploid progenies i.e. one distribution for each cross/trait combination. Manually inferring the complexity level for each of the cross/trait combination would be tedious and error prone. To help us in this task, we based our analysis on a constructed decision tree to classify distributions into different inheritance categories based on their underlying genetic complexity. Yet, the first step of this process was the determination of unimodality vs. bimodality of the distribution. This distinction is far from trivial and required to be assessed in a very specific manner. To do so, we used a machine learning algorithm, more precisely, we build a random forest classifier.

We classified the distributions in one of three complexity level: monogenic, oligogenic and complex. Overall, 80.3% of the considered distributions displayed inheritance patterns corresponding to a complex inheritance pattern. In the meantime, 11.2% appear as monogenic and only 4% as oligogenic. The remaining 4.5% failed to be sorted into one of the previous categories for various reasons, either the parents could not be confidently attributed to one cluster or the tetrad segregation phenotype could not result in a confident classification. These results confirm the fact that inheritance patterns are mainly complex but also that in a non-negligible number of cases, one gene is actually responsible for most of the observed genetic variance. However, this overview can be completed by the fact that this repartition of the complexity is highly dependent on the condition considered. Indeed, extensive variation in the complexity repartition can be observed in the 40 conditions explored here. When further dissecting the 153 distributions corresponding to an oligogenic inheritance, we highlighted several types of digenic interaction. We detected 87 cases of recessive epistasis. In 66 cross/trait combinations, modifier gene suggesting the presence of a suppressor have been identified.

Overall, we were able to assess the complexity level of traits at a species-wide level. We also highlighted the prevalence of expressivity with most of the followed variants displaying departure from monogenic inheritance patterns. Finally, this works lays the ground for a more complete and in detail exploration of variants displaying different levels of expressivity by dissecting the genetic basis of the observed cases. This dissection is the next step and will allow to have a better insight into the phenotypic expressivity landscape.

PRODUCTION SCIENTIFIQUE DE LA PÉRIODE 2016-2021 :


Since 2016, I had the chance to be invited to more than 30 international conferences and research institutes to talk about the research enabled by IUF.

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AUTRES AVANCÉES SIGNIFICATIVES AU COURS DE LA PÉRIODE :

PRIX ET DISTINCTIONS SCIENTIFIQUES OBTENUS AU COURS DE LA PÉRIODE

This project and the preliminary results we obtained allowed to access to 3 new fundings:

2019-2023 **ANR - Projet BrettAdapt**, Partner, 270 k€
Populational and multi-dimensional survey of the evolution, impacts and consequences of polyploidization in a yeast model

2019-2023 **ANR - Projet RecombFun**, Partner, 240 k€
Partner Evolution of the recombinational landscape and functional patterns across yeast species.

2018-2023 **ERC (European Research Council) Consolidator**, Coordinator, 1999 k€
Inheritance, expressivity and epistasis hidden behind the phenotypic landscape of natural populations
AUTRES OBSERVATIONS :

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