INTRODUCTION

Sporulation is a remarkable instance of phenotypic plasticity. It enables organisms to adapt to adverse nutrient-depleted conditions and survive in a metabolically quiescent and stress resistant state until vegetative growth is once again permitted (Driks, 2002; Knight & Goddard, 2016). Whilst diploid cells of the ascomycete yeast *Saccharomyces cerevisiae* primarily replicate asexually via mitosis, upon inhospitable conditions, they can undergo meiosis to form a tetrad of four haploid spores of two distinct mating types (a and α) (Honigberg & Purnapatre, 2003; Neiman, 2005; Piccirillo & Honigberg, 2010). Sporulation is a quantitative trait exhibiting complex mapping from genotype to phenotype (Ben-Ari et al., 2006; Deutschbauer & Davis, 2005; Gerke et al., 2009); a number of candidate genes have been detected with nonsynonymous mutations associated with variation in sporulation (Tomar et al., 2013).
It is widely recognized that sporulation enables dispersal by insect vectors. Drosophilids are closely associated with yeast throughout their life history, and play a central role in their dispersal (Dobzhansky et al., 1956; Gilbert, 1980; Reuter et al., 2007; Starmer & Fogelman, 1986). Whilst the acidic conditions of insect digestive systems destroy vegetative cells, spores generally survive passage (Coluccio et al., 2008; Reuter et al., 2007). Sporulation is primarily a starvation response, as the absence of nitrogen and fermentable carbon initiate the developmental pathway of meiosis and sporulation (Freese et al., 1982; Mata et al., 2002; Neiman, 2011; Primig et al., 2000). However, spores encounter digestive enzymes and altered pH in the digestive tract, and they generally remain unscathed. This would suggest that predation by insects has influenced their evolution (Thomasson et al., 2021).

Whilst spores survive due to the protective chitosan and dityrosine layers of the spore wall, the ascus holding sister spores together is broken down in the digestive tract (Coluccio et al., 2008). This provides potential for outcrossing between spores from different ascpiring upon the return of resource-rich conditions (Reuter et al., 2007). For *S. cerevisiae*, this is typically achieved by dispersal to a new fruit resource. Sporulation, and subsequent dispersal by insect vectors, thus provides a method by which *S. cerevisiae* can increase genetic diversity (Stefanini et al., 2016); this is particularly important for an asexual homothallic species with a tendency for automixis (Herskowitz & Jensen, 1991; Magwene et al., 2011; Zeyl & Bell, 1997). Increasing genetic diversity is useful prior to dispersal as it increases the likelihood that one or more daughter cells will possess a fitness sufficient to persist in the novel environment to be encountered (Otto & Lenormand, 2002).

Sexual reproduction is risky, however, because it enables the spread of parasitic genetic elements. It is believed that RNA interference (RNAi) and programmed cell death have evolved to attenuate infective elements (Drinnenberg et al., 2009; Gao, Chau, & Meneghini, 2019). For example, dsRNA M satellite and L-A helper viruses are absent from yeast reconstituted with an RNAi pathway, which is naturally absent (Drinnenberg et al., 2011). Moreover, Gao, Chau, Chowdhury, et al. (2019) found the same viruses were attenuated during sporulation by the NUC1 nuclease.

Biotic influences on sporulation efficiency, defined as the proportion of cells in a culture to sporulate, have been overlooked. Whether providing a net benefit or cost to the host, the L-A virus (and indirectly the M virus) utilizes host resources. As starvation induces sporulation, the presence of resource-utilizing intracellular endosymbionts should enhance the onset of starvation and sporulation under resource-limiting environments. This should be a natural by-product of their infectivity and virulence. We refer to this as the “infection by-product hypothesis,” whereby enhanced sporulation is an unavoidable by-product of infection, with its onset determined by parasite virulence and environmental resource availability. As starvation is the trigger of the sporulation pathway, we expect endosymbiotic infections to enhance sporulation no matter the strain explored. On the contrary, endosymbiont-free yeasts should have lower sporulation efficiencies, which may be detrimental and non-adaptive in competitive environments.

To investigate whether endosymbiosis is linked to enhanced sporulation we made use of the *S. cerevisiae* killer yeast system, which is composed of the yeast host, and two dsRNA viruses (M satellite and L-A helper viruses). The combination of M and L-A viruses encodes an antimycotic toxin for use by their host in interference competition, known collectively as the killer phenotype (Boyon, 2019; Wickner, 1996; Zhu et al., 1993). The dsRNA L-A genome encodes replication and transcription of the M virus and itself. The M genome is hosted and propagated by the L-A virion on which it is completely dependent. The M virus encodes toxin production and immunity to said toxin. These viruses provide context-dependent benefits to their hosts, depending on the level of competition present. We hypothesize yeast exhibiting the killer phenotype will have significantly enhanced sporulation efficiency relative to non-killers. We also predict that removing the M virus from killers will not significantly lower sporulation efficiency because the L-A virus will continue to utilize the hosts resources in the absence of the M virus.

## 2 | MATERIALS AND METHODS

### 2.1 | Study system

In total, 11 *S. cerevisiae* strains – five non-killers and six killers – were used (see Table 1). Of the killer strains, three belonged to the K1 killer-type, two to K2 and one to Klus. For each killer strain sample, a corresponding M virus-free strain was created from treatment with elevated temperature (Wickner, 1974), anisomycin (Fredericks et al., 2021) or cycloheximide (Fink & Styles, 1972) to destroy genetic material of the M virus. This procedure is commonly referred to as “curing” the strains. Killer strains and their cured equivalent were donated and cured with the aforementioned methods by the groups of Paul Rowley and Nieves Rodríguez-Cousiño (Fredericks et al., 2021; Rodríguez-Cousino et al., 2013). For a control, all non-killer strains were additionally independently exposed to elevated temperature (a widely utilized curing method) prior to the experiment (Wickner, 1974), as this would allow us to establish whether curing of the killer factor per se or simply a curing process can influence sporulation efficiency. This also leads to a full factorial design of cured non-killers, non-killers, cured-killers and killers. Background information on each strain can be found in the respective collections of each strain (see Table 1).

### 2.2 | Sporulation

Three independent replicates per *S. cerevisiae* strain were cultivated on standard YPD media (1% yeast extract, 2% peptone, and 10% glucose) at 25°C overnight, and subsequently the optical densities (OD) of the growing yeast cells were measured (SpectraMax 19, software: SoftMax Pro 6.2.2). When the OD was between 0.6 and 0.8, 50 ml of yeast were washed according to the protocol of Knight and Goddard (2016), with the following modifications: washed twice...
with 3 ml of sterile water. 0.5 ml of each yeast strain was plated on sporulation plates (40 mm plates, sporulation media: 1% potassium acetate, 0.1% yeast extract, 0.05% glucose, 2% agar, 25 mg/L zinc sulfate). Yeast plates were inoculated for 7 days at 25° C. Finally, we evaluated 100 cells for each strain replicate under the microscope and classified their sporulation status to calculate the sporulation efficiency, defined as the proportion of cells in a culture that have sporulated (Knight & Goddard, 2016).

### 2.3 Statistical analysis

All analyses were performed in R version 4.0.1 (R Development Core Team, 2020). To establish whether the presence of viruses or heat stress affects the proportion of sporulating cells, a linear mixed effect model was constructed in the package “lme4” (Bates et al., 2015), with strain as a random effect. We used the proportion of sporulating cells as the response variable. An interaction term of previous killer phenotype and previous killer type, and exposure to a curing treatment was created, leading to a full factorial design. Using simultaneous inference in general parametric methods (contrast analysis) in the package multcomp (Horthorn et al., 2008), any significant differences in sporulation rate between factors was established. Bonferroni corrections were applied. We treated strain as a random variable throughout to establish any effects on sporulation efficiency of the killer phenotype after accounting for variation in strain. The percentage of total variance explained by the strain was also calculated.

### 3 RESULTS

The proportion of cells to sporulate was consistently higher for killers than for non-killers (see Table 2, Killers vs. L-A only and
initial non-killers, and killers vs. non-cured non-killers) (Figure 1).

In general, curing did not have a significant effect on sporulation
(see Table 2, non-cured vs. cured, killer vs. L-A only, non-cured non-
killer vs. cured non-killer), though non-cured killers tended to have
higher sporulation rates than their cured equivalents ($F = 3.500,
p = .0669$; linear mixed effect model; Figure 1). This indicates that
killer yeasts have enhanced sporulation rates, but that the curing
methods used (which remove the M, but not the L-A) are not ca-
pable of lowering this rate significantly. Strain was found to only
explain 10.39% of variance in sporulation efficiency after the fixed
effects.

Killer type did not affect sporulation rate, but in general the dif-
f erent killer types had higher sporulation efficiencies compared to
non-killers (see Figure 2, non-cured non-killers vs. non-cured K1,
K2, and Klus: Estimate: $-0.352078$, SE: $0.101233$, Z-value: $-3.478$,
$p = .0091$). Furthermore, non-cured K1 was the only killer found to
be significantly different from non-killers (non-cured non-killers vs.
non-cured K1: Estimate: $-0.195765$, SE: $0.057697$, Z-value: $-3.393$,
$p = .0124$).

Finally, we find significant variation in the propensity to sporu-
late across all the strains used (see Figure 3). For non-killer strains,
there appears to be no general pattern concerning whether curing
increases or decreases sporulation efficiency. All killer strains ex-
cept YJM 1307 generally had higher sporulation efficiencies prior
to curing.

**4 | DISCUSSION**

The dsRNA M satellite and L-A helper viruses in *S. cerevisiae* are
known for encoding toxin production as an interference competi-
tion mechanism. Recently, Buser et al. (2021) found that yeast host-
ing dsRNA viruses were more attractive to *Drosophila simulans*,
a Drosophilid responsible for yeast dispersal (Buser et al., 2014).
Surviving dispersal via gut passage requires sporulation. Here, we
find evidence that the killer phenotype (the presence of the M sat-
eellite and L-A helper viruses) is significantly associated with higher
rates of sporulation. This finding hints at an acutely intertwined re-
lationship between the sexual life-stage of *S. cerevisiae* and infection
by viruses. It suggests that the dsRNA viruses play roles in *S. cerevi-
siae* life history and evolutionary ecology beyond interference com-
petition, despite the potential survival ramifications for the viruses
(Gao, Chau, Chowdhury, et al., 2019). In particular, it could mean that
viruses play a role in their host’s expansion into highly competitive
fermentation sites by enhancing sporulation rates, as is theorized of
endosymbionts (Joy, 2013; Sudakaran et al., 2017). Nonetheless, we
merely present a positive statistical association between the pres-
ence of dsRNA viruses and the sporulation efficiency, with the cause
as of yet unknown. Here, we outline three possible explanations for
this observation.

Our finding that non-cured killers have significantly higher sporu-
lation rates than the non-killers supports our infection by-product
hypothesis, that endosymbiont infectivity and virulence enhance sporulation efficiency (see Table 2). Starvation halts cell growth by mitotic division, inducing in its place the meiotic regulatory pathway (Honigberg & Purnapatre, 2003). Nitrogen and fermentable carbon limitations are primarily responsible for inciting sporulation, though limitations of phosphate, sulfate, guanine, and methionine are also demonstrated causes (Freese et al., 1982). The presence of endosymbiotic viruses may cause resource limitation, as resource usage is part of any infection. LA helper and M satellite viruses make use of free-floating nucleotides (Marquina et al., 2002; Schmitt & Breinig, 2006). If dsRNA viruses are diverting resources away from their host, meiotic gene expression may be triggered by a limitation of guanine or another nucleotide.

We found no significant difference in sporulation rate between current and former killer yeast, though there does appear to be a slight general decline in sporulation after curing (see Figure 1). The curing techniques used (elevated temperature, anisomycin, or cycloheximide application) remove the M satellite virus, but the LA virus typically persists. The M virus (often termed a virophage) is housed and propagated in the host cell by the LA virus (Schmitt & Breinig, 2006). The LA virus should harness a set quantity of resources from the host independent of the presence of the M virus; however, if the requirement to share resources with the M virus is absent, the LA viral copy number should be higher. One killer (YJM1307) had higher sporulation efficiency after curing which could be due to curing-induced modifications to sporulation genes (Lorenz & Cohen, 2014), which may also apply to some non-killers (see Figure 3). The high variation in the sporulation efficiency of yeast carrying a virus could be determined by viral load, which plausibly influences host metabolic costs of endosymbiosis. Though not explored in this paper, the degree of coadaptation likely affects whether hosts have evolved stabilizing mechanisms, such as sanctions of viral load (Gao, Chau, Chowdhury, et al., 2019; Wickner & Edskes, 2015), in order to compensate for the metabolic costs of endosymbiosis.

Alternatively, enhanced sporulation in killer yeast may be an adaptation to predation. Thomasson et al. (2021) demonstrated that sporulation is an adaptive trait, and that insect passaging selects for yeasts with higher rates of spore production. Drosophila attraction varies across yeast strains (Buser et al., 2014). Strains that are more attractive would therefore have higher sporulation rates as insect vectors disperse them more frequently. It was recently shown that killer yeasts are generally more attractive than non-killers (Buser et al., 2021). An alternative explanation for the higher sporulation rate in killers could therefore be that it is an adaptive trait making up for their higher attraction to insect vectors. A proximate explanation for their higher attraction may be a change in their volatile
composition resulting from virus infection, which remains untested. An ultimate explanation could be that killer yeasts are found more frequently in highly competitive fermentation environments meaning fruit flies consume them more regularly.

A third explanation is that viruses do not induce higher rates of sporulation. There is standing variation in the propensity of hosts to sporulate, demonstrated here (see Figure 3). This is likely due to variation in their abilities to acquire resources from the external environment and subsequent variation in the timing of starvation. Candidate genes have been linked to variation in sporulation efficiency (Tomar et al., 2013), whereby the significance of the non-synonymous mutation is determined by the stage of sporulation at which the gene is expressed (Lorenz & Cohen, 2014). As these strains sporulate more frequently, they are also more likely to undergo sexual reproduction, and outcross with non-clonal spores. If sexual reproduction is the only means of transmitting dsRNA viruses beyond the host lineage, viruses are only transmitted to strains that are sporulating. Therefore, strains with higher sporulation rates are more likely to be infected by viruses and other genetic elements. However, the cytosolic nucleases produced during meiosis degrade the M and L-A viruses of S. cerevisiae (Gao, Chau, Chowdhury, et al., 2019); it seems counterintuitive for frequently sporulating strains to host viruses when immune responses in sporulation actively degrade any foreign genetic elements. It appears that NUC1 does act as a host stabilizing mechanism to prevent uncontrolled proliferation of their cytosolic mycoviruses. However, it remains to be seen whether these dsRNA viruses can persist in their host if their pre-sporulation viral load is above a certain threshold. If so, this explanation is quite probable; however, if not possible, it gives credence to our infection by-product hypothesis, that sporulation is an unavoidable consequence of infection.

5 | CONCLUSION

Among the detrimental consequences of sexual reproduction is the transmission of parasitic genetic elements, of which resource exploitation is a consequence. Here, we provide evidence linking dsRNA viruses to enhanced sporulation in S. cerevisiae. Our findings have three possible explanations: (1) high sporulation rates are a consequence of the virulence of infectious viornes; (2) strains with higher sporulation rates outcross more frequently, thus providing an opportunity for infection by parasitic genetic elements; (3) sporulation is an adaptive trait of killer yeast to account for their higher attractiveness to dispersal vectors. Our findings hint at a deeply interwoven relationship between sexual reproduction and parasitism in S. cerevisiae. Future work is required to establish causation.

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CONFLICT OF INTEREST

The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

AUTHOR CONTRIBUTIONS

Thomas J. Travers Cook: Conceptualization (equal); Data curation (equal); Formal analysis (equal); Investigation (equal); Methodology (equal); Validation (equal); Writing – original draft (lead); Writing – review & editing (lead). Christina Skirgaila: Conceptualization (equal); Data curation (equal); Formal analysis (supporting); Methodology (equal); Writing – original draft (supporting). Oliver Y. Martin: Conceptualization (equal); Supervision (equal); Validation (equal); Writing – review & editing (equal). Claudia C. Buser: Conceptualization (lead); Data curation (supporting); Formal analysis (equal); Funding acquisition (lead); Investigation (equal); Methodology (equal); Project administration (lead); Resources (lead); Supervision (lead); Validation (equal); Writing – original draft (supporting); Writing – review & editing (equal).

DATA AVAILABILITY STATEMENT

The Data used in the analysis of this paper can be found in the Dryad Data Repository (https://doi.org/10.5061/dryad.x0k6djhm8).

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