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Protocol

Development of a ribosome profiling protocol to study translation in Kluyveromyces marxianus

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One sentence summary: Development of a ribosome profiling protocol to study gene expression in the thermotolerant yeast Kluyveromyces marxianus.

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Abstract

Kluyveromyces marxianus is an interesting and important yeast because of particular traits such as thermotolerance and rapid growth, and for applications in food and industrial biotechnology. For both understanding its biology and developing bioprocesses, it is important to understand how K. marxianus responds and adapts to changing environments. For this, a full suite of omics tools to measure and compare global patterns of gene expression and protein synthesis is needed. We report here the development of a ribosome profiling method for K. marxianus, which allows codon resolution of translation on a genome-wide scale by deep sequencing of ribosome locations on mRNAs. To aid in the analysis and sharing of ribosome profiling data, we added the K. marxianus genome as well as transcriptome and ribosome profiling data to the publicly accessible GWIPS-viz and Trips-Viz browsers. Users are able to upload custom ribosome profiling and RNA-Seq data to both browsers, therefore allowing easy analysis and sharing of data. We also provide a set of step-by-step protocols for the experimental and bioinformatic methods that we developed.

Keywords: nonconventional yeast (NCY), Ribo-Seq, RNA-Seq, transcriptome visualization, global analysis, industrial biotechnology

Introduction

As with other microbes, yeasts have evolved elaborate mechanisms to sense and respond to changing extracellular and intracellular environments. External influences include phenomena such as altered nutrient availability, toxic molecules, temperature fluctuations, low pH and high osmotic pressure, while internally, cells can experience changes such as reduced intracellular pH, ion fluxes, energy depletion or nutrient starvation (Martínez-Montañés et al. 2010, Broach 2012, Ljungdahl and Daignan-Fornier 2012, Morano et al. 2012, de la Torre-Ruiz et al. 2015, Sui et al. 2015, Taymaz-Nikerel et al. 2016). The best-studied response mechanisms in yeast involve sensor systems, signal transduction pathways and changes in gene expression (de Nadal and Posas 2010). Ultimately, this gives rise to a new set of proteins that enable the cell to adapt, if necessary, and to survive as well as proliferate in this new environment. Dissecting these response mechanisms is central to understanding the fundamental biology of a species, but it is also important for the development of yeast for biotechnological applications (Liu and Nielsen 2019). Yeasts are used for diverse applications in the food, biopharma and industrial biotechnology sectors (Arevalo-Villena et al. 2017, Nandy and Srivastava 2018, Parapouli et al. 2020) and, very often, they need to perform under suboptimal conditions or deal with a fluctuating environment. This is a particular problem when scaling engineered yeast cell factories in industrial biotechnology (Takors 2012, Delvigne et al. 2014, Wehrs et al. 2019). Developing a comprehensive understanding of adaptive responses is a key requirement for the construction of yeast cell factories that are both robust and resilient, and capable of optimal performance in an industrial bioprocess.

Most adaptive responses involve increased or reduced activity of specific proteins, which can be achieved at the level of synthesis, stability or activity. While some specific responses can be at the protein level, for example mediated by allosteric regulation, adaptation usually requires changes in the expression of many genes, and is considered to be a 'global' response. Changes in global gene expression can occur at various levels, most notably via transcription, translation or mRNA stability. Transcriptional changes are due to chromatic restructuring or changes in the activity of particular transcriptional regulators leading to increased or decreased levels of mRNA (Hahn and Young 2011). This is by far the best-understood adaptive process, deployed in response to heat shock (Masser et al. 2020), osmotic stress (de Nadal and Posas 2010), oxidative stress (Morano et al. 2012) and cell wall challenges (Sanz et al. 2017, Jiménez-Gutiérrez et al. 2020). Transcriptional responses can be studied at the level of individual genes using northern blots and reverse transcription quantitative realtime PCR (RT-qPCR), or globally by DNA microarrays or massively parallel sequencing (RNA-Seq), the latter of which has become the method of choice to study changes in gene expression (Schena et al. 1995, Gibson et al. 1996, Wang et al. 2009). Translation results in protein synthesis and, as such, is a better indicator of protein levels than transcription, though in many cases, higher abundance of mRNA due to increased transcription leads to a corresponding increase in the amount of translation. This is not always the

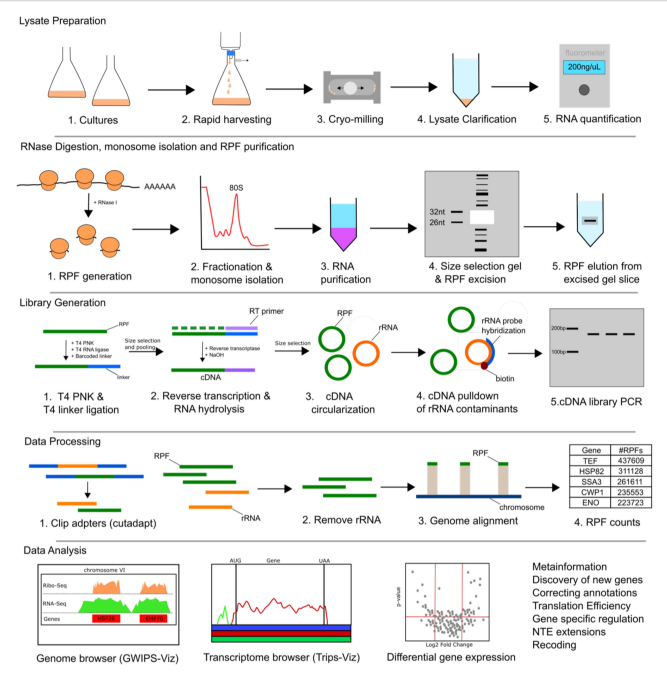


Figure 1. Summary of the ribosome profiling workflow. This summary is broken into five parts: lysate preparation, RPF generation and purification, library generation, data processing and data analysis. Lysate preparation includes culturing, lysis and the quantification of total RNA in a lysate. RNase digestion, monosome isolation and RPF purification represent the generation of RPFs. Library generation involves the conversion of small RNAs (RPFs) to a cDNA library, ready to be sequenced on an Illumina sequencing platform. Data processing involves removing of the sequencing adapters to leave only RPF sequences that are aligned to the genome. Data analysis typically involves visualization of data via genome and/or transcriptome browser, differential gene expression and a range of others as listed in the figure.

case, however, and there are also instances where translation is regulated without any changes in the mRNA abundance. A welldocumented example of this in yeast is regulation of the translation of the transcriptional activator encoded by GCN4. In this case, short open reading frames (ORFs) upstream of the main GCN4 coding sequence regulate the rate of GCN4 translation in response to intracellular amino acid levels (Hinnebusch 2005). The target of rapamycin (TOR) growth control system also mediates some of its effects by regulating translation via controlling access of the small ribosomal subunit to the cap structure at the 5' end of the mRNA (Merrick 2015). Indeed, as will be mentioned later, there is

an increased awareness that translational regulation is a central part of the yeast system for controlling gene expression.

Ribosome profiling, sometimes termed Ribo-Seq, is a method that allows for the visualization and quantification of translation at a global level. First developed in Saccharomyces cerevisiae (Ingolia et al. 2009), it has since been widely used in bacteria, yeast and mammalian systems for genome-wide studies of translation (Andreev et al. 2017, Ingolia et al. 2019, Mohammad et al. 2019). During mRNA translation, a ribosome translocates an mRNA one codon at a time and protects a fragment of mRNA within its mRNA tunnel (Steitz 1969). Ribosome profiling is a method to

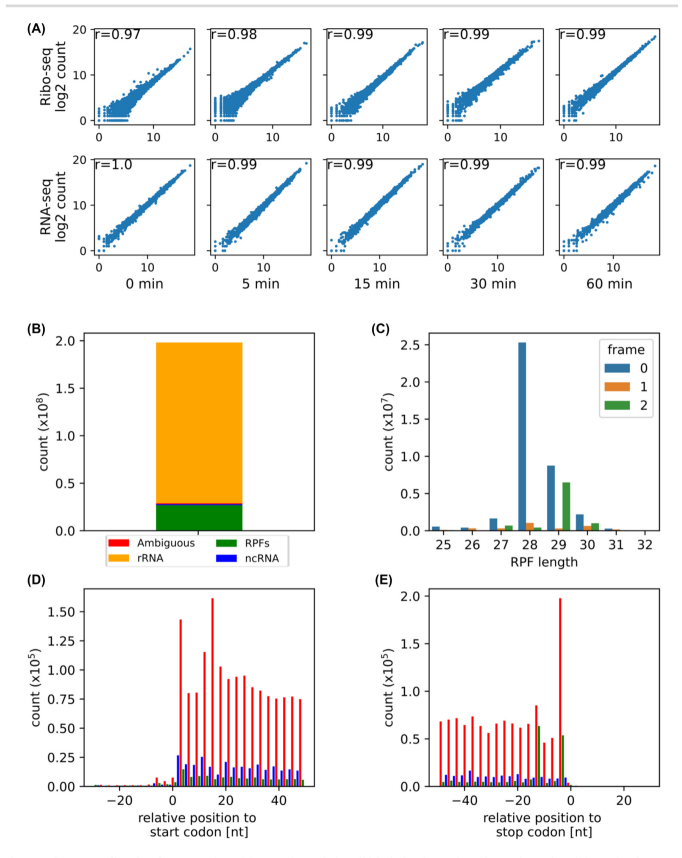


Figure 2. Ribosome profiling data from K. marxianus. (A) Pearson's correlation of biological replicates for each experimental condition. Axis values represent log₂ read counts. (B) Composition of ribosome profiling library with rRNA depletion. (C) Triplet periodicity of aligned RPFs for each read length. Panels (D) and (E) display a metagene profile of aligned RPFs near the start codon and stop codon, respectively.

Table 1. Biotinylated oligos for rRNA depletion. These oligos contain a 5' biotin modification to allow pulldown of specific rRNA contaminants using magnetic streptavidin beads.

Name	rRNA target	Sequence
rRNA#1	26S	5'AAGGGTGCATCATCGACCGATCCTG
rRNA#2	26S	5'GTTTCTTTACTTATTCAATTAAGCGGA
rRNA#3	Mitochondrial	5'TAAAGAATGGTACAGCTATAAATATT
rRNA#4	18S	
		5'GCTCGAATATATTAGCATGGAATAATGGA
rRNA#5	26S	5'TATAGAAGGATACGAATAAGGCGTC
rRNA#6	26S	5'TTTCCACGTTCTAGCATTCAAAGTCCT

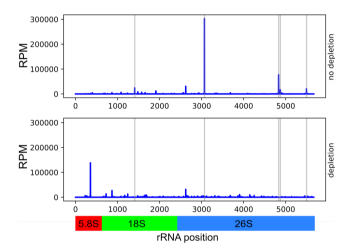


Figure 3. Targeted removal of nuclear-encoded rRNA contaminants. Abundance is represented in reads per million (RPM) and position is relative to the generated rRNA index presented in the bottom track. Top panel represents rRNA composition and abundance with no targeted rRNA depletion employed. Bottom panel represents rRNA composition and abundance with targeted rRNA depletion protocol. Targets for rRNA depletion are highlighted as dark grey areas. Abundance is represented in RPM and position is relative to the generated rRNA index presented in the bottom track. The grey vertical lines highlight the rRNA contaminants that are targeted in the oligo depletion step.

identify these ribosome-protected fragments (RPFs), thereby reporting which mRNAs are being translated at a given point in time. When applying the method, translation is arrested, usually by the addition of translation inhibitors, ribosomes are isolated and the RPFs are identified by deep sequencing. RNA-Seq is usually carried out in parallel to ribosome profiling, allowing estimation of changes in mRNA translation efficiency (Ingolia et al. 2009). In S. cerevisiae, ribosome profiling has uncovered widespread translation of upstream ORFs (Ingolia et al. 2009), non-AUG initiation at canonical genes (Monteuuis et al. 2019, Eisenberg et al. 2020) and small translated ORFs throughout the genome (Smith et al. 2014). Ribosome profiling combined with RNA-Seq has been useful in deciphering both transcriptional and translation regulation in the yeast meiotic programme (Brar and Weissman 2015) and in the response to oxidative stress (Blevins et al. 2019). Ribosome profiling has also been carried out on a range of other yeast species, including Saccharomyces paradoxus (McManus et al. 2014), Schizosaccharomyces pombe (Duncan and Mata 2014), Saccharomyces uvarum (Spealman et al. 2018), Komagataella phaffii (Alva et al. 2021) and Candida albicans (Sharma et al. 2021).

We are especially interested in another budding yeast, Kluyveromyces marxianus, which originally attracted interest be-

cause of its role in food fermentation (Coloretti et al. 2017) but is now increasingly being considered as a platform of industrial biotechnology (Fonseca et al. 2008, Lane and Morrissey 2010, Karim et al. 2020). Kluyveromyces marxianus has some intrinsic traits such as thermotolerance, a broad substrate range and rapid growth that are useful for biotechnology (Groeneveld et al. 2009), and molecular and genomic tools to aid its development as an industrial platform (Cernak et al. 2018, Rajkumar et al. 2019, Rajkumar and Morrissey 2020). To date, all studies that addressed gene expression in this yeast focused on transcriptional effects via RNA-Seq experiments. Aspects that have been studied include growth and ethanol production on alternative sugar substrates such as xylose (Schabort et al. 2016, Kwon et al. 2019) and inulin (Gao et al. 2015), ethanol tolerance during adaptive laboratory evolution (Mo et al. 2019), response to growth inhibitors derived from lignocellulosic substrates (Wang et al. 2018) and the ability to grow at high temperatures (Fu et al. 2019). Recently, mainly using transcriptome analysis, we determined that young genes specific to K. marxianus are enriched in the response to stresses such as high temperature, low pH and high osmolarity (Doughty et al. 2020).

To complement the molecular toolbox, and as a resource to study the biology of this yeast, here we report the development of a protocol to carry out ribosome profiling in *K. marxianus*. For this, we adapted and applied the methods previously used for *S. cerevisiae* (Ingolia et al. 2009). We also developed a suite of bioinformatics tools to visualize and analyse *K. marxianus* RNA-Seq and ribosome profiling results. This involved addition of the *K. marxianus* data to publicly available genome (GWIPS-viz) and transcriptome (Trips-Viz) browsers, which, in turn, can be uploaded with usergenerated expression data and used in private or public configurations. To facilitate the use of ribosome profiling as a very valuable tool to explore gene expression, we also include a detailed step-by-step protocol for users.

Materials and methods Strains and growth conditions

Kluyveromyces marxianus strain CBS 6556 (CBS-KNAW culture collection, Westerdijk Institute) was used in these studies following standard growth and handling procedures. This particular strain is also available from other collections under the strain names ATCC 26548, NRRL Y-7571, KCTC 17555 and NCYC 2597 and has been quite widely used as a representative K. marxianus strain. For ribosome profiling experiments, standard growth conditions used synthetic minimal medium (Verduyn et al. 1992) and an incubation temperature of 30°C with shaking. The mineral medium consisted of the following per litre amounts: (NH₄)₂SO₄, 5.0 g; KH₂PO₄, 3.0 g; MgSO₄·7H₂O, 0.5 g; trace elements (EDTA, 15 mg; ZnSO₄·7H₂O, 4.5 mg; MnCl₂·2H₂O, 0.84 mg; CoCl₂·6H₂O, 0.3 mg; CuSO₄·5H₂O, 0.3 mg; Na₂MoO₄·2H₂O, 0.4 mg; CaCl₂·2H₂O, 4.5 mg; FeSO₄·7H₂O, 3.0 mg; H₃BO₃, 1.0 mg; KI, 0.1 mg); and silicone antifoam, 0.05 mL. It was adjusted to pH 6.0 with KOH before autoclaving (121°C, 20 min). The medium was cooled to room temperature and a filter-sterilized solution of vitamins prepared in demineralized water was added to a final concentration, per litre, of D-biotin, 0.05 mg; calcium pantothenate, 1.0 mg; nicotinic acid, 1.0 mg; myo-inositol, 25 mg; thiamine HCl, 1.0 mg; pyridoxine HCl, 1.0 mg; and para-aminobenzoic acid, 0.20 mg. Glucose was sterilized separately and added to a final concentration of 10 g/L. One hundred fifty microlitres of cultures in 500-mL conical flasks were grown to early log phase at $A_{600} \sim 0.8$ and either harvested or transferred to a shaking water bath at 40°C, with cells harvested

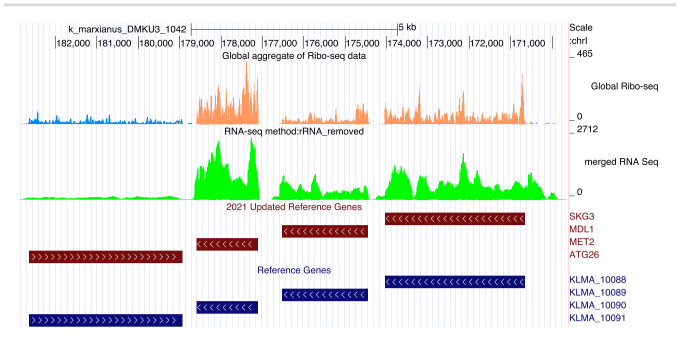


Figure 4. GWIPS-viz browser screenshot surrounding the SKG3, MDL1, MET2 and ATG26 locus of chromosome 1. Arrows on reference gene bars represent strand orientation. For ribosome profiling, orange reads represent positive strand RPFs, while blue reads represent negative strand RPFs.

at 5, 15, 30 and 60 min. All experiments were carried out with two biological replicates.

Ribosome profiling

For cell harvesting, cultures were quickly poured into a Durapore glass filter assembly (Merck, New Jersey, USA) using $0.45-\mu m$ pore nitrocellulose filter membrane (#7184-009, General Electric, New York, USA). A vacuum pump was immediately turned on and once liquid medium was removed, cells were quickly scraped into a 50mL Falcon tube filled with liquid nitrogen. Once a 150 mL culture is added to the filtration assembly, it takes \sim 9–12 seconds until the medium is removed and the scraped cell pellet is collected and submerged in liquid nitrogen. After harvesting, 1.5 mL of polysome lysis buffer (5 mM MgCl₂, 150 mM KCl, 20 mM Tris-HCl, 100 µg/mL cycloheximide (Sigma-Aldrich, St. Louis, USA), 1 mM DTT and 1% Triton X-100) was slowly added dropwise to the liquid nitrogen and cells to create a frozen mixture of buffer and cells. The 50-mL Falcon tube (with pierced cap from screwdriver) was placed in -80°C to allow boiling off of the liquid nitrogen. Frozen cells/buffer was disrupted using cryogenic grinding using a Retsch Mixer Mill 400 (Haan, Germany) and 10-mL steel grinding jars and balls. Samples were ground for six cycles of 3 min each at 20 Hz; the steel jars were submerged in liquid nitrogen to cool samples between each cycle. After lysis, lysates were gently thawed on ice and quantified with Qubit 4.0 fluorometer and BR assay kit (#Q10211, Waltham, Massachusetts, USA). Thirty micrograms of lysate was diluted to 200 μL in polysome buffer (lysis buffer without Triton X-100) and 1.5 μ L RNase I was added (#N6901K, Epicentre/Lucigen, Middleton, Wisconsin, USA). RNase digestion was carried out at 200 rpm at room temperature for 45 min. To halt digestion, SUPERase•In (Invitrogen) was added, and samples were placed on ice before loading onto cold 10-50% sucrose gradients, which were prepared using a Biocomp gradient master. Gradients were spun for 3 h at 4°C and $36\,000~\text{rpm}$ $(221632 \times q)$ on SW 41 Ti rotor (Beckman Coulter). Monosome fractions were isolated from each sucrose gradient with Brandel density gradient fractionator using 1.5 mL/min flow speed and 60%

CsCl, aliquoting fractions every 12 s on a UV-visible 96-well plate. Reading the 96-well plate at 260 nm determined which well(s) contained the monosome fractions. RNA from monosome fractions were isolated using TRIzol (Invitrogen) (Chomczynski and Sacchi 2006). Ribosome footprints were size selected with a 15% PAGE-urea (polyacrylamide gel electrophoresis) gel (70 min in 1× TBE and 300 V constant) using a 26 and 34 nt RNA marker (IDT) as a guide for excision. Using a scalpel, a slice representing the RPFs was cut from gel and placed into a 1.5-mL RNase-free Eppendorf tube and 500 μ L of RNA elution buffer (300 mM NaOAc, pH 5.5, 1 mM EDTA and 0.25% v/v sodium dodecyl sulfate (SDS)) was added. Following overnight shaking at room temperature to elute the RPFs, RPFs were precipitated using standard alcohol precipitation using ice-cold isopropanol, 80% ethanol and 1.5 μ L GlycoBlue coprecipitant (#AM9515, Ambion, Austin, Texas, USA).

Library construction

cDNA library construction with ribosome footprints is based on McGlincy and Ingolia (2017) protocol with minor modifications. In brief, size selected ribosome footprints were treated with T4 Polynucleotide Kinase (#M0201L, New England Biolabs (NEB), Ipswich, Massachusetts, USA)) followed by ligation to a DNA linker using T4 RNA ligase 2, truncated K227Q (#M0351L, NEB). The footprints were reverse transcribed using ProtoScript II (#M0368L, NEB). cDNA products were circularized using CircLigase II (#CL9025K, Epicentre/Lucigen). The major rRNA contaminants were removed using subtractive hybridization with custom biotinylated oligos (Sigma-Aldrich) and streptavidin beads (#65001, Invitrogen) as described in Ingolia et al. (2012). The remaining circularized products were amplified by PCR using Phusion polymerase (#M0530L, NEB). In a pilot experiments, libraries were sequenced on MiSeq platform at the Teagasc Next Generation DNA Sequencing Facility, Moorepark, Moorepark West, Fermoy, Co. Cork, Ireland. Prepared libraries using the protocol described within were sequenced on Illumina HiSeq 4000 using SE-75 sequencing at the Genomics & Cell Characterization Core Facility (GC3F), University of Oregon, Eugene, Oregon, USA.

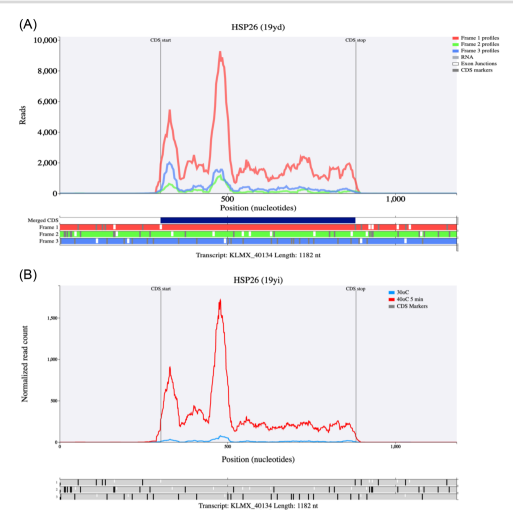


Figure 5. Trips-Viz transcript plots of HSP26. (A) Ribosome profiling coverage for aggregated data is displayed. Note that the dominant red line corresponds to the HSP26 CDS region. (B) Normalized transcript comparison plot of HSP26 showing increased mRNA translation at 30 and 40°C temperatures. Unique Trips-Viz plot identifiers are presented as '19yd' and '19yi' for panels (A) and (B), respectively. These identifiers can be added at the end of the following link to regenerate these specific plots (https://trips.ucc.ie/short/) on a web browser (example for panel (A): https://trips.ucc.ie/short/19yd).

Ribosome profiling data analysis preprocessing and genome annotation update

Adapter sequences were removed from reads using Cutadapt (Martin 2011). For genomic alignments, rRNA contaminants were removed and remaining reads were aligned to the *K. marxianus* DMKU3-1042 reference genome (Lertwattanasakul et al. 2015) with Bowtie (Langmead et al. 2009) using parameters -n 2 -m 1. Transcriptome alignments were made with Trips-Viz (Kiniry et al. 2019). For all genomic and transcriptome analysis, Ribosome profiling and RNA-Seq reads were aligned to the *K. marxianus* DMKU3-1042 reference strain (Lertwattanasakul et al. 2015).

RNA-Seq

RNA was isolated from clarified lysates using TRIzol (Chomczynski and Sacchi 2006) (Invitrogen #15596026) and quantified with a Qubit 4.0 fluorometer (Invitrogen). 1 μ g of total RNA from each sample was analysed on an agarose bleach gel to determine RNA quality. Samples were sent to BGI Hong Kong for yeast rRNA removal (Ribo-Zero Gold rRNA Removal kit by Illumina, San Diego, California, USA (now discontinued)), library generation and sequencing with paired-end chemistry. Alternatively, RNA-Seq was carried out using polyA selection using Poly(A)Purist Mag Kit

(#AM1922, Ambion) as per the manufacturer's instructions. PolyA selected cDNA library was generated and sequenced in the same method as ribosome profiling.

Results and discussion

Development of a ribosome profiling protocol to study translation in K. marxianus

The previously described *S. cerevisiae* protocol (McGlincy and Ingolia 2017) was used as the basis for development of a ribosome profiling procedure for *K. marxianus*. An overview of the pipeline from culturing to downstream bioinformatic analyses is shown in Fig. 1. Summarizing the first part of the protocol, cultures are rapidly harvested and flash frozen to preserve the translational state of the cell. Frozen cells are then lysed cryogenically in the presence of cycloheximide, which ensures that ribosomes remain stalled even if the cells thaw, and the clarified lysate containing polysomes is treated with RNase I to digest unprotected mRNA surrounding the ribosomes, retaining the RPF. Monosomes are isolated from a sucrose gradient and loaded on a polyacrylamide gel to allow size selection of RPFs of ~28nt. A cDNA library of these fragments is created and sequenced to identify the RPFs.

A limited-scale pilot experiment was first carried out to validate the methods and to identify the most abundant rRNA contaminants in the library. These arise because of RNase I digestion of rRNA and subsequent copurification of fragments of the same size as the RPFs. Due to natural polymorphisms in the rRNA encoding genes between species, the sequence of the major contaminated rRNA fragments needs to be determined empirically for each yeast. Knowing these sequences allows the design of synthetic biotinylated oligos that can be used to reduce rRNA contamination (Ingolia et al. 2012). In our pilot library, we identified six highly abundant rRNA fragments, four from the 25S rRNA, and one each from the 18S rRNA and from the mitochondrial 21S rRNA (Table 1). One of these sequences from 25S rRNA (GGGT-GCATCATCGACCGATCCT) comprised ~33% of all rRNA contaminants. By reducing rRNA contamination, the proportion of RPFs in a library is increased and thus more usable data are generated per experiment. The ribosome profiling protocol with rRNA depletion was then tested on a larger scale using 150 mL cultures of K. marxianus growing at different temperatures to increase the total number of genes that would be expressed. The focus at this time was on assessing the quality of the data generated and the robustness of the protocol rather than on analysis of changes in gene expression. Ribosome profiling was performed in duplicate on flask cultures at 30°C and at 5, 15, 30 and 60 min after a transfer from 30 to 40°C and, as is standard, RNA-Seq was also performed to measure transcript levels. Several analyses were performed to assess the robustness of the data that were obtained. First, the degree of correlation of the number of mapped reads per gene between biological replicates for each condition was assessed and found to be high with a Pearson's correlation of >0.96 (Fig. 2A). Second, we checked whether the RPFs actually represented known protein coding genes (Fig. 2B). We found that 14% of total reads aligning to the genome represented uniquely mapping RPFs; only $\sim 0.5\%$ of reads represented ambiguous RPFs, aligning to more than one location on the genome/transcriptome; and ~85% of the reads mapped to rRNA encoding genes. Third, we examined whether our data showed the distinctive triplet periodicity (or subcodon phasing) of the aligned reads reflecting the 'codon-wise' movement of elongating ribosomes that is seen in ribosome profiling data. In the dataset, footprints of length 28 nt (approximately half of total footprints) displayed a remarkable strong periodicity signal with \sim 95% of RPFs in phase with one of the three subcodon positions (Fig. 2C). Finally, RPFs are expected to be massively enriched in coding sequence (CDS) regions of genes. Using metagene profiles, we found that RPFs are largely present with CDS regions (Fig. 2D and E). In combination, these data demonstrate that the protocol generates robust ribosome profiling

Despite the oligo rRNA depletion, in our dataset from the largescale experiment, ~85% of the total reads were rRNA fragments. To determine the efficiency of the targeted rRNA contamination depletion, specific rRNA contaminant sequences were analysed before and after depletion. After depletion, we see almost 100% efficiency in removal of targeted rRNA contaminants, but there is not complete removal of all reads mapping to rRNA genes (Fig. 3). In part, this is because the rRNA abundance and composition may vary between samples and experiments due to slicing of RPFs from size selection gels by free hand. If desired, more oligonucleotides could be designed to further reduce rRNA contamination, thus increasing the proportion of RPFs in the sequencing pool. It was interesting to note that while ambiguously mapped reads can represent >10% of all reads in many studies from S. cerevisiae (seen looking at data in the Trips-Viz browser; https://trips.ucc.ie/) these comprised <1% of all reads in K. marxianus. Ambiguous mapping, whereby an RPF maps to two or more loci in the genome or transcriptome, arises because of the very short reads generated by ribosome profiling. As a result, it is not possible to determine the origin of the reads and these are generally discarded/ignored. This difference is most likely due to the large number of paralogous genes in S. cerevisiae, which arose through the proposed whole-genome duplication/hybridization (WGD) event in the evolutionary history of this species (Wolfe and Shields 1997, Marcet-Houben and Gabaldón 2015). As K. marxianus is a pre-WGD yeast, the same issue does not

Visualization of K. marxianus ribosome profiling data on public browsers

Visualization of ribosome profiling data is important to examine translation/transcription of particular loci of interest. We previously developed two tools to allow visualization of these data at a genome level (GWIPS-viz) and at the level of individual RNAs (Trips-Viz). These tools are freely accessible via RiboSeqOrg portal at https://riboseq.org. GWIPS-viz is a genome browser that displays RPFs mapped to each chromosome of a reference genome (Michel et al. 2014). The GWIPS-viz database already contained reference genomes for ~24 animal, plant, protozoal, fungal and viral genomes, and we added K. marxianus using the genome sequence and annotation from K. marxianus DMKU3-1042 strain as this was the most complete genome sequence available (Lertwattanasakul et al. 2015). It is possible to search GWIPS-viz by gene name or gene ID, and to zoom in/out of loci and as an extra feature that is new to GWIPS-viz, we included strand orientation of our ribosome profiling data to allow users to determine the strand to which an RPF is mapped (orange for +/forward strand, blue for -/negative strand) (Fig. 4). The browser is free to use and any user who generates their own ribosome profiling data or RNA-Seq tracks (bigWigs) can upload those data as custom tracks that can be viewed privately or made public. Once uploaded, a user is able to visualize and analyse their data using all the functionality of GWIPS-viz.

GWIPS-viz is mainly designed for analysis at a global level, allowing users to visualize any part of a genome, regardless of whether or not it is included in the annotations. In contrast, the second tool Trips-Viz is a transcriptome-level browser that focuses on individual mRNAs and allows a deep analysis of translation of each mRNA (Kiniry et al. 2019, 2021). This transcriptome browser allows users to generate single transcript plots displaying the ORF that is being translated. It also allows users to visualize the distribution of RPFs along an individual mRNA while also utilizing the triplet periodicity signal and differential colouring to identify potential translation in each ORF. As Trips-Viz did not include a reference transcriptome for K. marxianus, we created this reference transcriptome using our data. The application of Trips-Viz to study an individual mRNA is illustrated with an analysis of HSP26, using the (ribosome profiling) data for translation at 30 and 40°C (Fig. 5). The top panel uses aggregate data and shows the distribution of RPFs between each reading frame and across the transcript. It is clear that reads from the first ORF (red) dominate, which match the position and frame of the annotated CDS. The increase in the number of reads (RPFs) at certain positions indicates ribosome stalling during translation, for example at difficult-to-translate codons. The bottom panel compares the normalized read count of the correct ORF between the samples coming from cells grown at 30 and 40°C. The huge increase in translation at 40°C is evident. This was to be expected as HSP26

encodes a heat shock protein and is strongly transcriptionally induced by temperature shift. Thus, the increase in translation in this case is due to an increase in mRNA abundance. Although not shown in this simple example, in addition to single transcript plots, the Trips-Viz browser contains a large amount of metadata analyses such as triplet periodicity, read breakdowns, metaplot, protein count tables and differential expression analysis that is useful for detailed studies of translation and its regulation (Kiniry et al. 2019, 2021).

Integrated omics studies with K. marxianus

Ultimately, a full suite of omics technologies, ranging from genomics to proteomics, is a requisite for comprehensive studies of any microbe. Analysis of genome sequences and transcriptomes is now relatively straightforward for diverse yeasts, but the development of other tools still lags. Now, with the laboratory and in silico methods that we developed, it is possible to perform ribosome profiling with a nonmodel yeast, K. marxianus. By including both transcriptome (RNA-Seq) and translatome (ribosome profiling) analyses in future studies, it will be possible to generate a comprehensive view of gene expression at a point of time and in response to a perturbation. This can be very useful to understand biological processes and for the development of strains for biotechnology. The strategy taken, and the pipeline used, can also serve as a prototype for the development of ribosome profiling methods for other yeasts of biological and biotechnological interest. To facilitate the application of the tools by as many users as possible, a comprehensive step-by-step protocol is provided in the Supporting Information.

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Supplementary data

Supplementary data are available at FEMSYR online.

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Data availability

The data underlying this article are available in European Nucleotide Archive at www.ebi.ac.uk, and can be accessed with the project assession number PRJEB45612. The data have also been deposited to GWIPS-viz (https://gwips.ucc.ie/) and Trips-Viz (https: //trips.ucc.ie/).

Conflict of interest statement. PVB is a cofounder of RiboMaps Ltd, a company that provides ribosome profiling as a service.

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