Saccharopolyspora erythraea’s genome is organised in high-order transcriptional regions mediated by targeted degradation at the metabolic switch

Esteban Marcellin1†, Tim R Mercer1†, Cuauhtemoc Licona-Cassani1, Robin W Palfreyman1, Marcel E Dinger2, Jennifer A Steen1, John S Mattick2 and Lars K Nielsen1*

Abstract

Background: Actinobacteria form a major bacterial phylum that includes numerous human pathogens. Actinobacteria are primary contributors to carbon cycling and also represent a primary source of industrial high value products such as antibiotics and biopesticides. Consistent with other members of the actinobacterial phylum, Saccharopolyspora erythraea undergo a transitional switch. This switch is characterized by numerous metabolic and morphological changes.

Results: We performed RNA sequencing to analyze the transcriptional changes that occur during growth of Saccharopolyspora erythraea in batch culture. By sequencing RNA across the fermentation time course, at a mean coverage of 4000X, we found the vast majority of genes to be prominently expressed, showing that we attained close to saturating sequencing coverage of the transcriptome. During the metabolic switch, global changes in gene expression influence the metabolic machinery of Saccharopolyspora erythraea, resetting an entirely novel gene expression program. After the switch, global changes include the broad repression of half the genes regulated by complex transcriptional mechanisms. Paralogous transposon clusters, delineate these transcriptional programs. The new transcriptional program is orchestrated by a bottleneck event during which mRNA levels are severely restricted by targeted mRNA degradation.

Conclusions: Our results, which attained close to saturating sequencing coverage of the transcriptome, revealed unanticipated transcriptional complexity with almost one third of transcriptional content originating from un-annotated sequences. We showed that the metabolic switch is a sophisticated mechanism of transcriptional regulation capable of resetting and re-synchronizing gene expression programs at extraordinary speed and scale.

Keywords: RNA-sequencing, actinobacteria, Saccharopolyspora erythraea, Erythromycin, Metabolic switch

Background

Actinobacteria represent one of the most dominant and distinct bacterial phylum. The evolutionary divergence of actinobacteria is an ancient event, reflected in their unique biology. The majority of actinobacteria are found in soil, where they play a vital role in carbon cycling. In human medicine, they comprise a large number of pathogens and provide a major source of antibiotics [1]. They have a large, high GC content genome that equips them for growth in hostile, nutrient poor environments. They are also able to synthesise secondary metabolites that have important commercial and medical significance [2].

Saccharopolyspora erythraea (S.erythraea) is a model gram-positive filamentous actinomycete that has served for the study of antibiotic production for several decades. The bacteria produces erythromycin A, the first macrolide antibiotic to be discovered, and the backbone...
for numerous modern antibiotics [3]. *S. erythraea* exhibits a distinct and complex lifecycle, in which an initial growth phase is followed by a transition period, known as the metabolic switch, followed by a secondary growth phase. Like in *Streptomyces coelicolor* (*S. coelicolor*), the switch in *S. erythraea* is followed by morphological changes that coincide with potential cell death and the transcription of secondary metabolite gene clusters [4]. A detailed study of transcriptional activity during this transitional period is of major importance to understand the complex, yet little understood, life cycle of this sophisticated bacterial phylum. Studies have profiled actinomycete transcriptomes using microarrays [5,6]. In fact all annotated coding sequences have been profiled using Affymetrix arrays in *S. coelicolor* [7]. However, using long probes which are less likely to produce false negatives has proved challenging due to the high GC content of actinobacterial genomes. The recent advent of RNA sequencing provides a new opportunity to profile the global transcriptome of high GC content microorganisms.

Here, we present a detailed transcriptional description of *S. erythraea* life cycle from in-depth RNA sequencing. Our study shows that the major morphological changes observed at the metabolic switch are accompanied by an extreme transcriptional phenomenon, with the repression of large regions of the genome. In addition, we characterise a restrictive bottleneck event mediated by targeted RNase activity that results in a wholesale and rapid global transcriptional rearrangement event.

**Results**

**RNA sequencing shows pervasive transcription of the genome**

Like most actinobacteria, *S. erythraea* has a 71.1% GC content genome which prevented us designing and performing microarray experiments that contained all ORFs (Additional file 1: Figure S1). After performing microarray experiments in a control sample, we employed RNA sequencing to profile the whole transcriptome. We first assessed the ability of two alternative RNA sequencing approaches, duplex-specific thermostable normalisation (DSN) [8] and MicrobExpress depletion [9], to deplete rRNA in GC-rich transcriptomes. Enriched mRNA samples were compared to untreated matched reference controls (Additional file 1: Figure S1B,G,H). We did not observe any systematic divergence in nucleotide composition of sequenced reads between treated and untreated libraries \((r^2 = 0.01,\) Additional file 1: Figure S1B-H). We observed a high correlation between gene abundance estimates by independent validation by qRT-PCR and the expression of low GC content genes represented in the microarray (Additional file 1: Figure S1I,J), thereby confirming the fidelity of both strategies to profile GC rich transcriptomes.

Prior to harvesting RNA, several fermentations were performed to carefully characterize the metabolic switch in bioreactors. We determined that RNA harvested at seven time-points, (three points immediately before, one during and three following the metabolic switch) would accurately describe the entire genetic program (Figure 1A ). We prepared unstranded RNA libraries [10] and sequenced 363 million reads for a total of 32.7 Gb and a mean ~4,000 fold coverage of the genome (Additional file 2: Table S1, Additional file 2: Table S2). A replicate fermentation was sampled across the metabolic switch (Additional file 1: Figure S2C). The replicate samples were used to prepare strand-specific libraries that were sequenced to produce 86.1 million alignable reads and an additional 7.5 Gb of sequencing (Additional file 2: Table S1).

Our comprehensive sequencing showed the *S. erythraea* genome to be prevalently transcribed, with at least 85.8% of genome sequence from either strand being represented (98.7% of the genome without considering strand). This is similar to recent studies in yeast and other eukaryotes that show pervasive transcription of complex genomes that express intergenic and noncoding RNA [11,12]. Furthermore, given less than 0.3% of expressed nucleotides were represented singly suggests that close to saturating sequencing coverage of the transcriptome was attained, thereby permitting us to detect rare and transient transcriptional events.

**The transcription of genomic macro-regions alternates at the switch**

We next applied RNA sequencing to resolve large-scale topological features of the *S. erythraea* transcriptome. Our sequencing clearly illustrates the strand-specific enrichment for transcription from the leading strand, consistent with co-directional transcription of essential and highly expressed genes with replication [13] (Figure 1B). Like other actinomycetes, the genome is divided into two large regions, with a core region thought to contain the majority of genes essential for survival, and a non-core region enriched for genes for conditionally adaptive functions [2,14]. Alignment of sequenced reads across these regions allowed the resolution of genome-scale, contiguous transcriptional regions associated with these core/non-core regions (Figure 1C). These macro-regions undergo alternative enrichment across the switch, with marked repression of the non-core transcriptional loci following the switch (Figure 1C).

The borders of the core/non-core macro regions are delineated by transposable sequences (Figure 2 and Additional file 1: Figure S2A) [2]. Given the capacity for RNA sequencing to resolve the expression of repetitive sequences, we were able to detect a notably strong and specific induction \((p = 6.5\times 10^{-7})\) of paralogous transposon
clusters at these region borders, coincident with the shifting of non-core to core transcriptional regions. These borders orientate the transcriptional regions perpendicular to the strand-specific enrichment for leading/lagging strands (Figure 1B). When superimposed, these regions dissect the genome into four distinct quarters; with each region containing distinctly co-expressed and functionally related gene families (Figure 1C,D). Collectively this demonstrates a broad regulatory context provided by contiguous transcriptional macro-regions for the functional organisation and alternate expression of genes across the metabolic switch.

The metabolic switch defines a distinct gene expression program

The high temporal resolution and sequencing depth with which we profiled the growth cycle and metabolic switch using RNA-seq (with >4000 fold genome coverage), revealed considerable complexity in gene expression profiles. Although we detected transcription from all genes, for clarity we restricted our analysis to prominently expressed genes (top 50% of the total most prominently expressed genes). Shifts to gene expression invoked at the switch were so broad as to affect 60.5% of the genes that were profiled (>2-fold change). To delineate cogent trends within this complexity, we identified major gene groups that exhibited a normalised enrichment at exponential, transitional or stationary phases across the *S.erythraea* life cycle (Additional file 1: Figure S3A-C, S4A-C). A comparative analysis delineated two distinct gene expression programs with diverging functional trends before and following the switch (Additional file 2: Table S2). For example, the greater number of genes expressed prior to the switch were enriched for programs associated with central carbon metabolism, such as those involved in carbohydrate synthesis.
(p = 1.75x10^{-16}), cell wall biogenesis (p = 0.043), energy production (p = 6.80x10^{-4}) and lipid metabolism (p = 0.011). In contrast, genes associated with secretion and transport (p = 0.037), posttranslational modification (p = 8.98x10^{-6}) and ribosomal structure and biogenesis (p = 1.12x10^{-16}) were enriched after the switch. Similarly, the erythromycin gene cluster was enriched after the metabolic switch (Additional file 1: Figure S3). During the metabolic switch the greatest enrichment was for genes associated with signal transduction (p = 0.0011) and transcription regulation (p = 1.94x10^{-9}). These alternate programs were reciprocally enriched for core/non-core and leading/lagging strand organisation (Figure 1D), reflecting the overarching context these macro-regions provide for gene expression. Interestingly we found similar patterns of up or down regulation to previously reported homologous genes in closely related microorganisms [7,15], suggesting that the transcriptional phenomenon described here comprise a characteristic event that punctuates and defines gene expression in related microorganisms (Additional file 1: Figure S8).

RNA sequencing facilitates operon annotation
By the contiguous transcription across shared and co-expressed genes, we looked at the predicted organization of the 1,134 previously predicted operons that encompass 4,919 genes [16]. Our de novo assembly using Oases[17], from the strand specific RNA sequencing after trimming all reads to 70 bases and assembling them using a 45 bases k-mer, resulted in 9,170 contigs. We found several of these contigs aligned with previously predicted operons (Additional file 2: Table S3). Moreover, using the strand specific sequencing, we found 2.9% of total transcription to be antisense to these annotations, with an enrichment of antisense transcription at the 3' end of the genes. Lastly, we employed a range of metrics, including presence, size and structure to annotate novel independent ncRNAs within intergenic regions. By Combining small RNA sequencing and long RNA-seq data, we identified 190 putative ncRNA from transcribed sections in intergenic genomic regions (Additional file 2: Table S7 and ncRNA track in the genome browser). The novel ncRNAs display distinct CPC scores [18], dynamic transcriptional pattern and 14 of them displayed a distinct ncRNA secondary structure (Additional file 1: Figure S7; Additional file 2: Table S4, S7) [18].

RNA degradation mediates a transcriptional bottleneck event
The two alternative genetic programs are demarcated at the switch by a bottleneck event that is readily apparent in transcriptomic data. Significant enrichment for functional gene categories at different stages of the growth cycle is indicated.
phenotypic changes that occur during this phase. Furthermore, the wide imbalance between the highest and lowest expressed genes (the top 1% genes contribute 27.8% of total mRNA) suggests the majority of mRNAs to be present at less than one copy per cell, consistent with recent reports of heterogeneity between individual cell transcriptomes [19]. This imbalance narrows during the bottleneck event, suggesting decreased diversity between individual cell transcriptomes. This observation is in accordance with the previously described programmed cell death that occurs at the switch and involves expression of enzymes for cellular dismantling [20]. Notably, we observed that BioAnalyzer traces from two independent fermentations showed degraded profiles at the switch (Figure 3C, S4H). Furthermore, we observe the pronounced expression of ribonucleases and proteases at the inception of the switch (21 out of 24 ribonucleases; p < 0.0001 paired t-test (Figure 3A,B)), including RNase E and RNase III. Given the critical roles ascribed to these ribonucleases in regulating the half-life of developmental transcripts associated with the translational process, including ribosomal genesis (4.6-fold, p = 2.12x10^-5), ribosomal proteins (1.4-fold, p = 6.3x10^-7) and other associated

Degradation targets specific mRNA classes

Our analysis of RNA integrity across the *S. erythraea* life cycle revealed extensive RNA degradation coincident with the metabolic switch (Figure 3C and Additional file 1: Figures S4H). Given that samples were treated with RNA-later and that we obtained high RIN numbers (except for the sample corresponding to the metabolic switch), it was unlikely that degradation could occurred post-extraction. Although the Qiagen columns have less binding affinity for RNAs smaller than 200 bp, it does not completely remove sRNAs [23]. Thus, to provide an insight into the degradation pattern, we isolated and sequenced these degraded RNA fractions. We firstly isolated fragments corresponding to 20-50nt in size, thereby depleting the preparation of small regulatory RNAs that were greater than 50nt in length [24]. We also reasoned that, unlike the 5’ monophosphate termini of a fragmented transcript, primary RNA transcripts with a 5’ triphosphate would be refractory to library preparation and sequencing (Additional file 1: Figure S5D) [25]. For comparison we also sequenced matching fractions from before (40 hours) and after (93 hours) the switch, returning approximately 16 million sequenced small RNAs (Additional file 1: Figure SSC). We next omitted any reads that did not align in sense to transcribed regions from matched RNA sequencing libraries, reasoning they may reflect bona-fide small RNAs. Alignment of the remaining small RNA fragments exhibit no apparent size or nucleotide enrichment to annotated coding genes previously resolved.

We next identified those mRNAs subject to targeted degradation and turnover across the switch by comparing the ratio of full-length and fragmented reads that align to each transcript. Although we observed broadly accelerated mRNA degradation at the switch (Additional file 1: Figure S6A), we noted this degradation to be relatively slower or faster for specific genes and functional classes and to often occur in concert with our previously described gene expression profiles. For example, consistent with their enrichment at the switch, we found that while transcripts encoding transcription factors generally exhibit higher stability (2.1-fold, p = 4.5x10^-12, Additional file 1: Figure S6), some specific classes, such as XRE transcription factors, known to regulate developmental life cycle and secondary metabolism [26], undergo a cohesive and accelerated degradation (Additional file 1: Figure S6F). This observation establishes that the degradation can be targeted to specific mRNA classes, thereby contributing to the stage-specific enrichments observed across the *S. erythraea* growth cycle. Gene ontology analysis of *S. erythraea* genes most intensively targeted for destruction at the metabolic switch returned a large enrichment for transcripts associated with the translational process, including ribosomal genesis (4.6-fold, p = 2.12x10^-5), ribosomal proteins (1.4-fold, p = 6.3x10^-7) and other associated
translational factors. We detected the targeted degradation of additional functional gene networks, such as energy production and regulation, which undergo suppression at the bottleneck. Collectively, these trends demonstrate the contribution of targeted mRNA degradation to the transcriptional bottleneck and the modulation of genetic programs subsequently deployed (Figure 4). They also anticipate a pivotal and global role for ribonucleases, such as RNase E and III, in regulating the metabolic switch [21,22,27,28].

Discussion

Our analysis of the *S. erythraea* transcriptome describes a major transcriptional phenomenon occurring during fermentation in bioreactors. A genome-wide transcriptional non-core region, delineated by genes of markedly lower GC content, sparse gene density and inverted repeats, is broadly repressed following the switch. This region organization is present in related microorganisms, including some with linear genomes, such as the terminal regions of the *S. coelicolor* linear chromosome where analogous insertion elements mark the inner boundary of terminal inverted repeats [29]. Similar low GC regions have been reported in *Streptomyces ambifaciens*, where these regions have been associated to genome plasticity and may be used to elucidate mechanisms of evolution [30]. The presence of analogous transposable elements delineating the core/non-core regions is apparent in other related microorganisms anticipating a role in the maintenance and possible regulation of these macro-regions. The alternating expression of these regions, previously observed in *S. erythraea* using microarrays [6], is able to mediate a broad coordinated suite of genetic changes, representing a broad architectural mechanism by which a cell can rapidly modulate large changes in the transcriptome in response to environmental cues [31].

This transcriptional reorganization is complemented by a targeted bottleneck event where mRNAs are degraded. The bottleneck event establishes that the degradation may be targeted to specific mRNA classes, thereby contributing to the stage-specific enrichments following the *S. erythraea* growth cycle. The importance of these degradation processes in global transcriptomic regulation has been recently dissected in yeast [32,33] and individual cases described in *S. coelicolor*, whereby endoribonuclease activity of RNase III has been shown to bind to specific transcript and to regulate antibiotic production [21,34]. Studies have also described the role of (p)ppGpp in regulating mRNA half-life in *S. coelicolor* by inhibiting PNPase activity which preserves mRNA for crucial cellular processes [35-37]. Such widespread degradation may also indicate a role for programmed cell death of a distinct cellular subpopulation, in mediating the changes at the metabolic switch. The new transcriptome observed after the switch may be reflective of the transcriptome of the surviving subpopulation. Collectively, these observations of cellular degradation and targeted mRNA degradation supports the emergence of key, yet little studied, mechanism by which gene expression are regulated.

Our results, which attained close to saturating sequencing coverage of the transcriptome, reveal unanticipated additional transcriptional complexity with almost one
third of transcriptional content originating from unannotated sequences. Combining RNA sequencing with sequencing of small RNAs showed the importance of non-coding RNAs in bacterial transcription.

Conclusions
Like other related microorganisms, *S. erythraea* undergoes a distinct transitional switch that is characterized by growth arrest, morphological changes and the production of secondary metabolites. We show this to be accompanied by a massive transcriptional modulation whose speed and scale, revealed here for the first time by RNA sequencing, represents a significant and novel transcriptional phenomenon. Given similarities shared between the life cycle and genome organization with other microorganisms, we suggest that this transcriptional phenomenon may be a widespread feature across the large critically important bacterial group of industrial microorganisms.

Methods

Bacterial strain, growth and fermentation conditions

*S. erythraea* (NRRL2338) was used in this study. All fermentations were conducted in 2L bioreactors (Applikon) in mineral salt medium MM-101 without casaminoacids, as described elsewhere [38,39]. Medium ISP 2 (yeast extract, 4 g/L; malt extract, 10 g/L; Dextrose, 4 g/L; Agar, 20 g/L) was used as solid media for spore germination and seed cultures. Approximately 0.5 mL of glycerol stock was used to inoculate starting cultures in a 500 mL baffled flask with 100 mL of ISP 2 media incubated at 30°C in a rotary shaker (INFORS HT, Bottmingen, Switzerland) at 220 rpm for 30 h. When the seed culture reached an OD₅₅₀ of 2.5 (early stationary phase), a second seed culture (1 L baffled flasks with 150 mL of ISP 2) was inoculated to an initial OD₅₅₀ of 0.3 and incubated under the same culture conditions for 72 h. Cells were then centrifuged at 10,000 rpm at room temperature (Allegra X-15R, Beckman Coulter, USA), washed and resuspended in MM-101 prior to inoculation. Temperature and pH remained constant at 30°C and pH = 7.0 respectively by addition of 20% NaOH or 10.9% HCl. Dissolved oxygen was maintained between 45 and 60% saturation. Oxygen uptake rate (OUR) and CO₂ production were measured using a DASGIP Off-Gas Analyzer GA4 (DASGIP AG, Jülich, Germany) or a mass spectrometer (Hiden HRPR20-QIC B, Warrington, England) attached to the bioreactor condensers. Cells were harvested from two biological replicates at seven time points for RNA extraction (see below). Erythromycin A was analysed by LCMS as described in [39].

Total RNA isolation and mRNA enrichment
To protect RNA from endogenous RNA degradation, RNA isolation was performed after 8 h incubation at 4°C in RNA-later (Ambion). Total RNA was extracted using two cycles of cellular lysis in RNase-free zirconia beads. RNA was purified using RNeasy midi kit (Qiagen) with on column DNase treatment and a second off column DNase treatment. Although the Qiagen columns have less binding affinity for RNAs smaller than 200 bp, it does not completely remove sRNAs smaller than 200 bp, and are therefore suitable for sRNA analysis. RNA quality was evaluated using BioAnalyzer (Agilent) and Nanodrop prior analysis. Ribosomal RNA was removed with MicroExpress Bacterial mRNA Enrichment kit (Ambion) or duplex-specific thermostable nuclease enzyme from Kamchatka crab (DSN) as described in the Illumina's protocol. RNA samples quality and integrity was analyzed using 2100 BioAnalyzer (Agilent) according to manufacturer’s instructions.

Microarrays and real-time quantitative PCR
Probes for microarrays, were designed using eArray and OligoRankPick [40] and evaluated using Agilent’s Base Composition (BC) algorithm. RNA was amplified using MessageAmp II-bacterial RNA Amplification kit (Life Technologies), reversed transcribed to cDNA, labeled using Agilent QuickAmp labeling kit (Life Technologies), hybridized, washed and scanned according to Agilent’s protocols. RT-PCR was performed from cDNA amplified from 600 ng of total RNA using Superscript III (Invitrogen) as per the manufacturer’s instructions for high GC% transcripts (50°C, 1 h). Diluted cDNA (1 in 50) was used for qRT PCRs performed in a Corbett Rotor-gene 3000 using SYBRgreen UDG master-mix (Invitrogen). Cₚ values were determined using the default values on the Rotor Gene- 6 software. A correction for DNA contamination was performed using Cₚ values from RT controls. The concentration of cDNA (ng/reaction) for each sample was determined using corrected values. These values were then log₂ transformed and normalized against eryBV (Additional file 1: Figure 1-i).

Library preparation and sequencing
Deep sequencing was performed from libraries prepared as follow: Two libraries were prepared for total RNA, seven after DSN treatment [8] and the second round of sequencing (from an independent fermentation), following mRNA enrichment using MicroExpress (Ambion), was sequenced strand specifically. Additionally, we ran a trial DSN-sequencing sample sequenced after preparation of three independent 90-nt, paired-end libraries. To achieve optimized clustering density, the trial run was sequenced with two 400mer libraries sequenced at two different concentrations following DSN normalization [41,42]. The third library was size selected from a gel after the final amplification to remove adapter multimer product and sequenced at 300mer. This optimized protocol for GC-rich
microorganisms was used for subsequent DSN-sequencing. Libraries prepared after MicrobExpress mRNA enrichment were directionally sequenced at 90bp as described in the direction mRNA-seq prep pre-realised protocol from Illumina [43]. RNAs corresponding from 15-50nt fractions relative to ladder were excised for from the PAGE gel and purified for sequencing using the Illumina small RNA sequencing protocol with minor modifications as previously described [44]. All sequencing was performed at Gene-works (Adelaide, Australia) on the Illumina GAII. All Data is available at GEO GSE39722.

Sequence read alignment and analysis

Reads were independently aligned to the S.erythraea genome using multiple software to assess for possible alignment bias caused by the high GC content of the genome. Reads were aligned using Burrows-Wheeler Aligner BWA [45] Bowtie [46] and Bowtie2 [47] requiring no more than 2 mismatches. Only reads that aligned to one genomic location were retained unless specified for the analysis. We employed CuffDiff [48] to determine gene expression using SAM files resulting from alignment, annotated gene file upper quantile normalization and masking of rRNA sequences. For expression of non-standard features, such as transcriptional macro-regions, read alignments were summed over genomic annotations. Hierarchal clustering of gene expression was performed by Pearson’s pair-wise complete linkage using Cluster3 [49]. Gene expression was normalized within libraries and between libraries as indicated in [50]. To determine significance of gene ontology enrichments, we performed Fisher’s exact test with multiple hypothesis correction. Additional statistical tests and figure generation were conducted using the Prism 5 (http://www.graphpad.com/prism/). Additional in-house perl scripts provided at http://matticklab.com/index.php?title=Marcelin_Dinger was also employed. We also employed for figure generation Circos to generate circular genomic figures [51]. Operon prediction was performed from de novo assembly of all reads resulting from the strand specific biological replicates using Oases [17]. Reads were trimmed to 70 bases and assembled using 45 bases kmer, from which 23% aligned with previously annotated operons (Additional file 2: Table S3, operon track in Gbrowser and de novo track in Gbrowser).

Additional files

Additional file 1: Supplementary Data are available online. All data is available on a dedicated S. erythraea genome browser (http://pathway. aibn.uq.edu.au/seythraea) and RNA-seq data has been submitted to GEO GSE39722. http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=pjcxjcsimegyze8&acc=GSE39722

Additional file 2: Table S1. Summary of sequenced libraries and alignments. Table S2. Gene expression profiles for annotated genes.

Gene identifier, description and normalised expression (RPKM) indicated. RPKM result from Cufflinks after removing multi-mappers (1), upper quantile normalization and masking of rRNA sequences. Bowtie2 alignments (2) are also presented. Table S3. Operon validated using Oases for de novo assembly of all reads. The majority of the DOOR operons present similarity with previous annotation (Mao et al., 2009).

Table S4. Novel ncRNAs annotated with secondary structure and dynamic profile across the fermentation. Unique identifier, chromosome location, and size indicated. Table S5. Gene-ontology analysis. Enrichment for GO terms at exponential, transitional and stationary phases within S.erythraea growth cycle. Table S6. Gene expression profiles for annotated genes. Gene identifier, description and expression before Quantile normalization (FPKM) indicated. Reads were aligned using Bowtie 2 and analysed using Cufflinks. Genes marked with ‘+’ in Column J were considered highly expressed.

Competing interests

The authors declare no competing financial interests.

Authors’ contributions

TRM, EM and LKN designed the experiments. EM and CLC performed all fermentation and extractions. EM and CLC perform protein purification and analysis. TRM, RP and MED performed alignment, transcript assembly and analysis. JS performed RT-PCR, EM, TRM, CLC, JSM and LKN wrote the manuscript. All authors read and approved the final manuscript.

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Author details

1Australian Institute for Bioengineering and Nanotechnology (AIBN), The University of Queensland, Brisbane, QLD 4072, Australia. 2Institute for Molecular Bioscience (IMB), The University of Queensland, Brisbane, QLD 4067, Australia.

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