

Structural and functional adaptation of *Haloferax volcanii* TFE α / β

Fabian Blombach^{1,*}, Darya Ausiannikava², Angelo Miguel Figueiredo¹, Zoja Soloviev¹, Tanya Prentice¹, Mark Zhang¹, Nanruoyi Zhou¹, Konstantinos Thalassinos¹, Thorsten Allers² and Finn Werner^{1,*}

¹Institute for Structural and Molecular Biology, Division of Biosciences, University College London, London, UK and

²School of Life Sciences, University of Nottingham, Queen's Medical Centre, Nottingham NG7 2UH, UK

Received July 11, 2017; Revised December 08, 2017; Editorial Decision December 15, 2017; Accepted December 19, 2017

ABSTRACT

The basal transcription factor TFE enhances transcription initiation by catalysing DNA strand-separation, a process that varies with temperature and ionic strength. Canonical TFE forms a heterodimeric complex whose integrity and function critically relies on a cubane iron-sulphur cluster residing in the TFE β subunit. Halophilic archaea such as *Haloferax volcanii* have highly divergent putative TFE β homologues with unknown properties. Here, we demonstrate that *Haloferax* TFE β lacks the prototypical iron-sulphur cluster yet still forms a stable complex with TFE α . A second metal cluster contained in the zinc ribbon domain in TFE α is highly degenerate but retains low binding affinity for zinc, which contributes to protein folding and stability. The deletion of the *tfeB* gene in *H. volcanii* results in the aberrant expression of approximately one third of all genes, consistent with its function as a basal transcription initiation factor. Interestingly, *tfeB* deletion particularly affects foreign genes including a prophage region. Our results reveal the loss of metal centres in Hvo transcription factors, and confirm the dual function of TFE as basal factor and regulator of transcription.

INTRODUCTION

The RNA polymerase (RNAP), auxiliary factors and molecular mechanisms of transcription are conserved between archaea and eukaryotes. Many Archaea inhabit extreme environments and their enzymes have adapted to cope with conditions of high temperature or salinity, while the underlying changes of protein chemistry enabling this tolerance remain poorly understood. The molecular engineering challenges of initiating transcription in a start site-specific

fashion are independent of the type of RNAP and the domain of life. RNAPs need to engage with the promoter region of the double-stranded DNA template, the two DNA strands need to be separated, the template strand loaded into the RNAP active site, and individual NTP substrates bound in a geometry conducive to catalysis. All cellular RNAPs depend on additional basal transcription factors to facilitate this process. In archaea three basal transcription factors (TBP, TFB and TFE) assist RNAP during initiation. TBP and TFB bind to the TATA and BRE motifs and are necessary and sufficient for RNA polymerase recruitment to the promoter and transcription initiation. TFE is not strictly required for transcription *in vitro*, but stimulates transcription initiation by enhancing open complex formation (OC), i.e. DNA melting and template loading (1).

Both monomeric and heterodimeric TFE variants have been characterized in archaea. In the crenarchaeon *Sulfolobus solfataricus* (Sso) TFE is a heterodimer consisting of TFE α and TFE β subunits (2). TFE α encompasses an N-terminal extended winged helix-turn-helix (eWH) domain (3) and a C-terminal Zn-ribbon domain (ZR) that is flanked by coiled-coil helices (4,5) (Figure 1). TFE β encompasses an N-terminal canonical winged helix-turn-helix (WH) and a C-terminal domain coordinating a structural cubane [4Fe-4S] cluster (2). The cluster is highly sensitive to oxidation and its integrity is critical for the stability of TFE α / β and its function (2). The broad phylogenetic distribution of genes encoding TFE α and TFE β homologues in archaeal genomes suggests that these genes were present in the last common ancestor of all archaea (1). While TFE α is conserved in the vast majority of archaeal species, several archaeal lineages appear to have lost the TFE β gene (*tfeB*). Canonical *tfeB* genes are conserved in all cren- and thaumarchaeal genomes (6). More divergent putative *tfeB* homologs have also been identified in different euryarchaeal lineages (Methanomicrobiales, Archaeoglobales and Halobacteriales), while the gene is absent altogether from other euryarchaea like *Methanocaldo-*

*To whom correspondence should be addressed. Tel: +44 20 7679 0147; Fax: +44 20 7679 7193; Email: f.werner@ucl.ac.uk
Correspondence may also be addressed to Fabian Blombach. Email: f.blombach@ucl.ac.uk

requirements for DNA strand separation during OC formation depend in large part on the physicochemical conditions of the environment. Repulsive forces within the phosphate backbones of the DNA strands destabilize the dsDNA helix, but can be counteracted by cations such as Mg^{2+} (17). The effect of human TFIIE on OC formation varies accordingly with KCl and Mg^{2+} concentration (18). Similarly, the effect of archaeal TFE α/β on OC formation increases at higher Mg^{2+} concentrations (F.B. and F.W., unpublished observations).

Haloarchaea thrive in a high salinity environment that is matched by high intracellular K^+ concentrations, which is thought to have caused global changes in the haloarchaeal proteomes such as low isoelectric points (19). Many studies have shed light on the function of TFB and TBP in the haloarchaeal transcription machinery, which is characterized by its broad expansion of TFB and TBP paralogues that contribute to global regulation of transcription (20–22). In contrast, the function of haloarchaeal TFE has not been investigated thus far. Interestingly, the putative haloarchaeal TFE β homologs are unusual in that they lack the cysteine residues that chelate the iron-sulphur cluster (Figure 1) (6). This raises the question whether the haloarchaeal homologues function as proper TFE β subunits in a dimeric complex or whether haloarchaea utilize instead monomeric TFE α variants as described for other members of the euryarchaea. Does the divergent primary sequence of haloarchaeal TFE β homologues reflect adaptations to the structure and function of haloarchaeal TFE?

Here, we show that the *tfeB* gene from *Haloferax volcanii* (Hvo) does encode a bona fide TFE β subunit, which does not encompass an FeS cluster and yet still forms a stable complex with TFE α . In addition, the second metal centre of TFE, the TFE α zinc ribbon that is crucial for RNAP binding, is highly degenerate with only a single cysteine residue contributing to zinc binding. This results in a very low affinity for zinc, yet collision induced unfolding experiments demonstrate that zinc binding stabilizes the protein structure. In contrast to *Sulfolobus*, the *Haloferax tfeB* gene is not essential for cell viability. The *tfeB* deletion causes a slow growth phenotype and leads to the transcriptional misregulation of one third of all genes. This substantiates the role of TFE as global regulator in the archaeal domain.

MATERIALS AND METHODS

Heterologous expression and protein purification of Hvo TFE α/β

The gene HVO_1174 coding for Hvo TFE α was PCR-amplified and cloned into a modified version of vector pRSF-1b (2) via NcoI and XhoI restriction sites (for all primer and plasmid details see Supplemental Tables S1 and S2, respectively) for the expression of native protein or as a C-terminal His₆-tag fusion, respectively. HVO_1090 coding for Hvo TFE β and its variant TFE β Δ 1–72 were cloned likewise via NdeI and XhoI restriction sites into vector pET-21a(+) (Merck). Plasmids were transformed into *Escherichia coli* Rosetta2 (DE3) (Merck) and heterologous expression was carried out in enriched growth medium for 2 h after induction at 37°C. Cells were resuspended in N(500) buffer (25 mM Tris/HCl pH 8.0, 500 mM NaCl, 10 mM

MgCl₂, 100 μ M ZnSO₄, 5 mM 2-mercaptoethanol), disrupted by sonication, cleared by centrifugation and Hvo TFE was isolated by Ni-affinity chromatography.

A TEV-protease cleavage site was introduced between Hvo TFE α and the C-terminal His-tag by the Phusion site directed mutagenesis method (ThermoFisher Scientific). The resulting protein TFE α TEV was expressed and purified as described above. Ni-affinity elution fractions were dialyzed against 20 mM Tris/HCl pH 8.0, 300 mM NaCl, 5 mM 2-mercaptoethanol in the presence of His-tagged TEV protease overnight. TEV protease was removed by Ni-affinity chromatography and successful cleavage was verified by SDS-PAGE.

Size exclusion chromatography was carried out in N(500) buffer with a flow rate of 0.5 ml/min on a Superose 12 hr 10/300 column (GE Lifesciences) with 0.5 ml fractions collected.

Gene deletion and growth conditions

Growth and transformation of *H. volcanii*, isolation of genomic DNA and construction of deletion mutants was carried out as described previously (23). The *tfeA* gene HVO_1174 is in an operon with HVO_1175 and HVO_1176, which was PCR-amplified (see Supplemental Table S1) and cloned in pTA1744, a derivative of pTA131 (23), to generate pTA1769. A deletion construct pTA1814 was generated using outward facing primers to amplify upstream and downstream flanking regions of *tfeA*, respectively. A novel NcoI site was used for insertion of a promoter-less *trpA* selectable marker from pTA1844, generating pTA1858; care was taken to not affect transcription of the remaining operon. pTA1858 was used to transform Hvo strain H53 (Δ *pyrE2* Δ *trpA*, see Supplemental Table S3) and Δ *tfeA::trpA*⁺ colonies were screened as described previously (23). All 129 colonies examined by hybridization were merodiploid and had retained the wild-type *tfeA*. Therefore, *tfeA* is most likely essential.

The genomic region containing the *tfeB* gene HVO_1090 was PCR-amplified and cloned in pTA131 (23) to generate pTA1706. A deletion construct pTA1708 was generated using outward facing primers to amplify upstream and downstream flanking regions of *tfeB*. A novel BglIII site was used for insertion of the *trpA* selectable marker from pTA298 generating pTA1709, which was used to transform Hvo strain H53 and Δ *tfeB::trpA*⁺ colonies were screened as described previously (23). 120 of the 160 colonies examined by hybridization were Δ *tfeB*, which was verified for 8 candidates by Southern blot (Figure 5A).

To measure growth of the Δ *tfeB* mutant strains H2644 and H2645 in Hv-YPC broth (23), 250 μ l cultures were incubated in 48-well plates at 45°C, with continuous double-orbital shaking at 425 rpm, using a BioTek Epoch2 microplate spectrophotometer. Optical density at 600 nm was measured every 15 minutes and generation times were calculated by curve-fitting to the log phase of the growth curve, as indicated by the straight-line portion between perpendicular marks in Figure 5B. The standard errors of generations times across replicates = 0.07 h for wild-type and 0.13 h for Δ *tfeB* mutant. To complement the Δ *tfeB* mutation *in trans*, a KpnI–NotI fragment of pTA1706 containing the

tfeB gene and its promoter was cloned in the shuttle vector pTA354 (24) to generate pTA2050, which was used to transform the $\Delta tfeB$ mutant H2644 and its parent H53; the pTA354 empty vector was used as a control. Growth was measured in a microplate spectrophotometer as described above (Figure 5C). Raw growth data are provided in Supplemental Table S4.

RNA-seq RNA isolation

Total RNA was isolated from *H. volcanii* using the TRIzol method. 50 ml of cells grown in Hv-YPC broth to early log phase ($OD_{650} = 0.2$) were resuspended in 250 μ l of unbuffered spheroplast solution (1 M NaCl, 27 mM KCl, 15% sucrose, pH 7.5) (23). 750 μ l of TRIzol LS (Invitrogen) was added and mixed by pipetting. 200 μ l of chloroform (Sigma) was added and vortexed for 15 s, then centrifuged at 20 000 $\times g$ for 15 min. The upper phase was transferred to a fresh tube, 500 μ l of isopropanol was added and incubated for 10 min at room temperature. RNA was pelleted by centrifugation at 20 000 $\times g$ for 10 min, washed with 75% ethanol, air-dried and resuspended in 50 μ l of RNase-free dH₂O. Total RNA quality was assessed using the RNA 6000 Nano Kit (Agilent) on an Agilent 2100 Bioanalyzer. Total RNA concentration was measured using Qubit RNA BR assay kit (Life technologies, Q10210). 1 μ g of total RNA was used for rRNA depletion using RiboZero rRNA Removal Kit (Illumina). Libraries were prepared using the dUTP method (25) with NEBNext Ultra Directional RNA Library Prep kit for Illumina (New England Biolabs) and library quality was analyzed on a Bioanalyzer High-Sensitivity DNA-chip (Agilent Biotechnologies). Sequencing was performed on the Illumina NextSeq 500 sequencing platform to generate 2 \times 75 bp paired end reads using V2 chemistry.

RNA-seq data analysis

Sequencing reads were trimmed to remove the adaptor parts using Trimmomatic (26) and the quality of the sequencing data was assessed using FastQC (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc>). Reads were mapped to the Hvo genome using TopHat (27) using parameters `-no-novel-juncs -no-mixed -library-type fr-first strand` with filtering out reads mapping to tRNA and rRNA genes as well as reads that do not map uniquely. Principal component analysis (PCA) was used to assess biological replicate concordance. RPKM values (reads per kb per million) were calculated using HTseq-count. Data were analyzed using DESeq2 to identify significant differentially expressed genes ($P_{adj} < 0.01$) (28).

In order to calculate transcript abundance, we combined an available transcriptome map of Hvo (29) with recently published transcription start site (TSS) mapping data (30) to create an updated map of Hvo mRNA transcriptome. For 1331 transcription units (TUs) the 5'-end coordinates were adjusted based on mapped primary TSSs. For these cases, the median adjustment was by 5 bp, whereas in 117 cases an adjustment of >100 nt was required, in the vast majority leading to an extension of the TU. Six annotated polycistronic TUs harboured additional internal primary

TSSs. In these cases the TU annotation was revised and shortened to the primary TSS position. 62 protein encoding genes with assigned TSSs were not included in the existing TU map (29). This updated map (Supplemental Table S7) was used to estimate expression levels using RSEM in paired end mode and $-$ -strandedness reverse for libraries prepared with the dUTP method (31).

The same transcription start site mapping data were used for promoter analysis. DNA sequences were extracted using BedTools getfasta (32) for positions -50 to $+10$ relative to the TSS and analyzed using the MEME software version 4.11.4 for motif identification (0 or 1 occurrence per sequence, 4–16 bp width, searching given strand only) (33). A 0-order background model based on the composition of the combined *H. volcanii* DS2 genome was employed. Aligned promoter sequences were depicted using WebLogo3 with correction for the genome composition (34). Synonymous codon usage data for Hvo genes have been described previously (19). AT% of genes was calculated using BedTools nuc (32). Data were plotted using R (<http://www.R-project.org>) with the LSD package v3.0.

Native mass spectrometry

For nESI mass spectrometry (MS) proteins were buffer changed to 0.5 M NH₄ acetate by ultrafiltration. Zn(II)-acetate was added to the samples at 2:1 molar ratio followed by 1 hr incubation at room temperature. All samples were analyzed on a first generation Synapt mass spectrometer (Waters corp., Manchester, UK) (35). Samples were introduced into the mass spectrometer by means of nESI at a concentration of 10–20 μ M. For protein unfolding experiments ZipTips (C4, Millipore) were used to buffer exchange samples into 50–70% acetonitrile containing 0.1% TFA according to the manufacturer's protocol. All samples were introduced into the mass spectrometer using *in-house* prepared gold-coated capillaries (borosilicate glass capillaries with dimensions 1.0 mm \times 0.78 mm from Harvard apparatus) created using a needle-puller (P-97, Sutter Instrument) and coated with gold using a sputter-coater (SC7620, Emitech), as described previously (36). The mass spectrometer was externally calibrated using a 30 mg/ml solution of caesium iodide. Instrument setting parameters were as follows: capillary voltage 1.2 kV, cone voltage 40 V, source temperature 40°C, extraction cone voltage 1 V, trap collision energy (CE) 6 V, transfer CE 4 V and bias voltage 4 V. For complexes with a mass >25 000 Da, the backing pressure was increased to 4 mbar. For collision induced unfolding (CIU) experiments bias voltage was increased to 17 V, and a CE ramp (6–36 V) in the trap collision cell was used. Mass spectra were analyzed in MassLynx software v4.1 (Waters) and CIU maps were analyzed using Amphitrite (37).

NMR spectroscopy

Isotopically labeled samples for NMR studies were prepared by growing BL21 (DE3) cells for heterologous expression of protein TFE α TEV in minimal (M9) medium using as nitrogen source ¹⁵NH₄Cl (CK Isotopes). The protein was purified as described above by Ni-affinity chromatography, TEV protease cleavage and size exclusion chromatography,

but the last step was carried out in an NMR compatible buffer (10 mM HEPES/KOH pH 7.5, 100 mM NaCl, 1 mM DTT).

NMR experiments were performed on a Bruker Avance III 600 MHz spectrometer, equipped with a 5 mm TXI room temperature probe at 25°C. The protein concentration for samples used for NMR studies was typically kept at 0.3 mM in the same buffer as above supplemented with 7% (v/v) $^2\text{H}_2\text{O}$. Zn^{2+} was added using a ZnSO_4 stock solution diluted to a final concentration of 165 and 190 mM corresponding to ~1:1 molar ratio for $\text{TFE}\alpha_{\text{TEV}}$ and $\text{TFE}\alpha_{\text{TEV}}$ C138S, respectively. 2D ^{15}N - ^1H HSQC spectra were acquired for each Zn^{2+} concentration. Data were processed using the NMRPipe/NMRDraw (38) package and analyzed with CCPNMR (39).

RESULTS

H. volcanii $\text{TFE}\alpha/\beta$ forms a stable heterodimer

The euryarchaeon *H. volcanii* (Hvo) harbours ORFs annotated in the arCOG database as $\text{TFE}\alpha$ and β subunit-encoding genes (arCOG04270 and arCOG04153, respectively) (40). In order to determine whether their gene products interact, we cloned the two genes *tfeA* (HVO_1174) and *tfeB* (HVO_1090) into compatible plasmids for coexpression in *E. coli*. Either of the two proteins was fused to a C-terminal His-tag in order to facilitate purification using Ni-affinity chromatography. SDS PAGE analysis of the elution fractions revealed that both $\text{TFE}\alpha$ and β subunits co-eluted irrespective of which subunit was tagged, suggesting that they form a stable complex. The fractions containing the co-eluting proteins were furthermore subjected to gel filtration (Figure 2). Both $\text{TFE}\alpha$ - and β -subunits eluted symmetrically with their predicted mobility, demonstrating that they form a stable heterodimeric $\text{TFE}\alpha/\beta$ complex (Figure 2A). The Sso $\text{TFE}\beta$ FeS cluster domain is essential for dimer formation, while the WH domain is dispensable. We constructed the corresponding WH deletion variant of Hvo $\text{TFE}\beta$ and verified its complex formation with $\text{TFE}\alpha$, which confirms that the WH domain is dispensable for dimer formation in Hvo and demonstrates that the domain requirements for dimerization are conserved between Sso and Hvo $\text{TFE}\alpha/\beta$ (Figure 2B).

Unusual metal centre binding properties of Hvo $\text{TFE}\alpha/\beta$

FeS clusters are commonly coordinated by cysteine residues, but alternatively arginine, histidine or aspartate residues can substitute for single cysteine residues (41,42). In the case of haloarchaeal $\text{TFE}\beta$ candidates, the absence of all four cysteine residues suggests that they do not contain FeS clusters (Figure 1). The $\text{TFE}\alpha$ ZR domain contributes the second metal centre of archaeal $\text{TFE}\alpha/\beta$ complexes. In the human TFIIE α ZR domain, four cysteine residues chelate a zinc ion (43). Likewise, the majority of archaeal $\text{TFE}\alpha$ homologues use four cysteine residues for zinc binding. Histidine, glutamate and aspartate residues have been shown to participate in zinc coordination in place of cysteine, and in some archaeal $\text{TFE}\alpha$ variants including Sso one of the four cysteines in the ZR domain has been substituted by an aspartate residue (44). Surprisingly, Hvo $\text{TFE}\alpha$

features only a single cysteine residue while the first, second and fourth cysteine positions are substituted by serine, alanine and threonine residues, respectively (Figure 1). This degenerate Hvo $\text{TFE}\alpha$ ZR domain has likely lost its ability to coordinate zinc altogether, or involves glutamate or other residues present in the loop of the two β -hairpins that form the canonical ZR motif. Notably the conservation of $\text{TFE}\alpha$ cysteine residues differs between the two major groups within the haloarchaea. While some halophilic archaea such as *Halobacterium salinarum* have $\text{TFE}\alpha$ variants with all four cysteines conserved, others such as *Haloquadratum walsbyi* $\text{TFE}\alpha$ have the same substitutions as in *H. volcanii* with a single conserved cysteine.

In order to characterize the nature of the metal binding properties of Hvo $\text{TFE}\alpha/\text{TFE}\beta$, we carried out native nano electrospray-ionization (nESI) mass spectrometry. The native mass spectrum of $\text{TFE}\alpha/\text{TFE}\beta$ -His confirms the formation of a dimeric complex (Figure 2C). The $\text{TFE}\alpha/\text{TFE}\beta$ -His complex was detected as a set of four peaks of calculated masses of 34 613.28 (\pm 1.12) Da, 34 743.97 (\pm 0.03) Da, 34 676.93 (\pm 0.03) Da and 34 807.17 (\pm 0.57) Da due to two different additional masses splitting the signal independently. One split is due to the partial N-Met cleavage of $\text{TFE}\alpha$ by methionine aminopeptidase during heterologous expression in *E. coli* (Supplementary Figure S1); this cleavage accounts for a mass difference of 131 Da. The second split is compatible with the presence or absence of a zinc ion, which accounts for an additional mass of 65.38 Da. The spectra revealed a small amount of free $\text{TFE}\beta$ -His that resulted in a second charge state series with a mass of 14 092.45 (\pm 0.31) Da coinciding well with the theoretical mass of $\text{TFE}\beta$ -His (14 090.5 Da, Figure 2C). This suggests that the zinc present in the $\text{TFE}\alpha/\text{TFE}\beta$ complex must be bound by the α subunit – in spite of the degeneracy of the zinc binding motif. We also noted that extended washing lead to complete loss of the heavier species indicative of zinc loss (data not shown).

Zinc binding stabilizes the degenerate $\text{TFE}\alpha$ ZR domain

In order to characterize the binding of zinc to the $\text{TFE}\alpha$ ZR domain, we produced a recombinant monomeric $\text{TFE}\alpha$ variant ($\text{TFE}\alpha_{\text{TEV}}$) that allowed the removal of the affinity tag following purification. To prevent zinc loss due to the additional purification steps, we re-incubated the protein with 1 mM ZnSO_4 before the buffer exchange into NH_4 acetate compatible with nESI MS. The native mass spectrum of $\text{TFE}\alpha_{\text{TEV}}$ showed two peaks of similar intensity with a calculated mass of 21 425.5 (\pm 0.4) Da and 21 490.6 (\pm 0.0) Da corresponding very well with the theoretical mass of $\text{TFE}\alpha_{\text{TEV}}$ (21 425.65 Da) and an additional zinc ion (65.38 Da, Figure 3A). In order to characterize any putative zinc chelating residues, we mutated the single conserved cysteine residue in the $\text{TFE}\alpha$ ZR domain to serine and recorded its native mass spectrum. The C138S variant showed one major peak with a calculated mass of 21 410.8 \pm 0.8 Da corresponding to the theoretical mass of the ligand-free protein (21 409.59 Da) and a very small population with a mass of 21 475.4 (\pm 1.4 Da) matching the theoretical mass of the zinc-bound form (21 474.97 Da, Figure 3B).

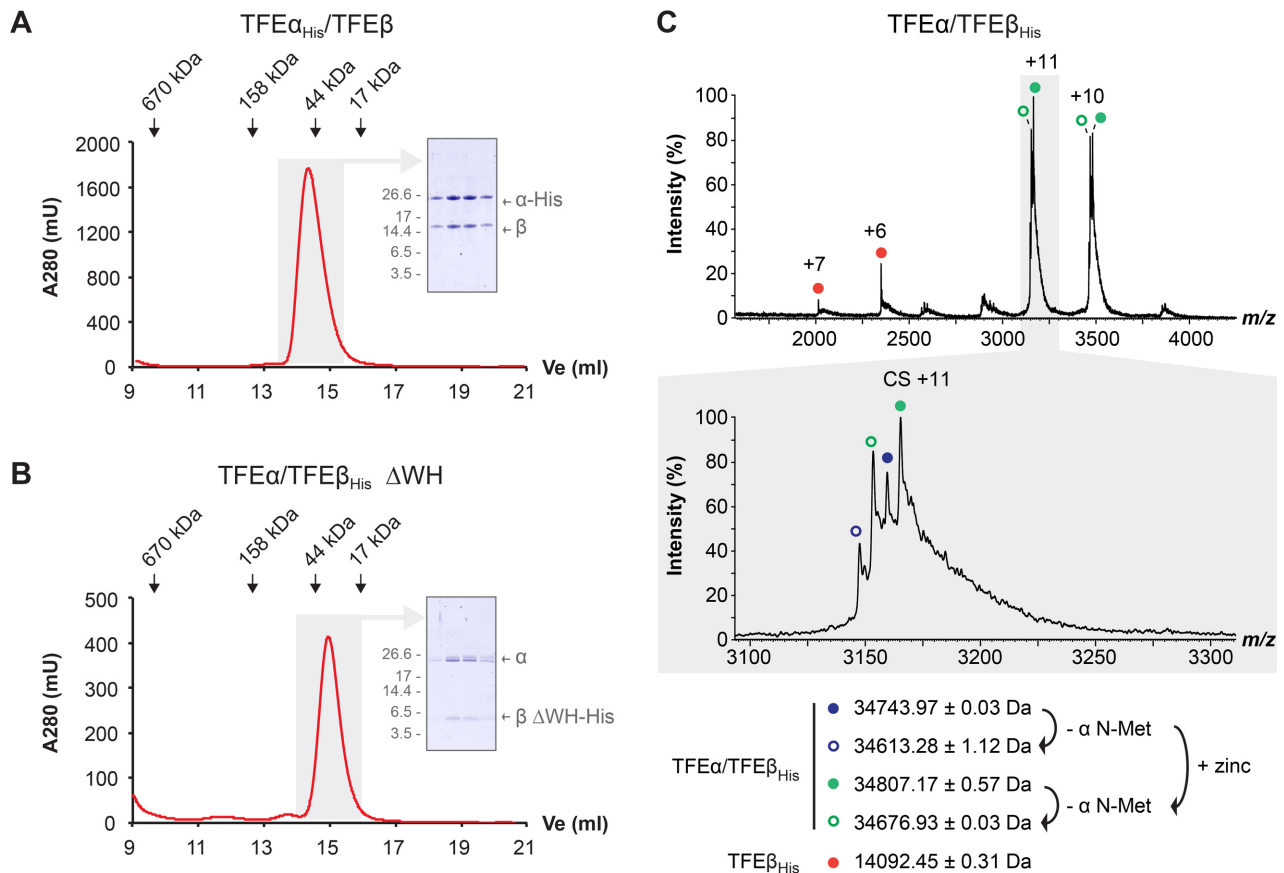


Figure 2. *Hvo* TFE α and TFE β form a dimeric complex in the absence of an FeS cluster. (A and B) Size exclusion chromatography of TFE α_{His} /TFE β (A) and TFE α /TFE β_{His} Δ WH (B) on a superose 12 h 10/30 column. Elution peaks of standard proteins are indicated. Peak fractions (highlighted in gray) were subjected to SDS-PAGE and Coomassie-staining (insert) to show symmetric elution of both subunits. (C) nESI mass spectrum of TFE α /TFE β_{His} . The different charge state series observed are labelled with red, blue and green filled circles. Charge state series derived from additional cleavage of TFE α N-Met are labelled with open circles. Highlighted in gray is a zoom over the region that includes charge state (CS) +11. All detected masses indicate the absence of a cubane 4Fe-4S cluster (352 Da).

In order to assess whether zinc binding increased the stability of TFE α_{TEV} , we carried out collision-induced unfolding (CIU) experiments. CIU combines activation of proteins by voltage to induce protein unfolding coupled with ion mobility mass spectrometry to detect the folding status (45). Ligand-free TFE α_{TEV} appeared in two populations, a faster travelling population (\sim 15 ms) corresponding to the native folded protein, and a slower travelling population (\sim 25 ms) equivalent to a more unfolded state triggered by a higher voltage (\sim 20 V, Figure 3C). To analyze the effect of zinc, we preincubated TFE α_{TEV} with 1 mM zinc acetate, isolated the corresponding peak (21 490.6 Da charge state +8) using the quadrupole mass analyzer, and subjected it to CIU using a voltage ramp in the trap collision cell prior to ion mobility separation. Interestingly a significantly higher voltage (\sim 25 V) was required to convert zinc-bound TFE α_{TEV} into the more unfolded state, indicating that zinc binding indeed exerted a stabilizing effect on TFE α . To gain further insight into the possible structural differences between WT TFE α_{TEV} and the C138S variant we recorded two-dimensional (2D) ^{15}N - ^1H Heteronuclear Single Quantum Coherence (HSQC) spectra. The HSQC

spectra of TFE α_{TEV} and the C138S variant showed good dispersion and superimposed well on each other with a single crosspeak shift between the two spectra that likely corresponds to C138 (Figure 4). These results demonstrate that (i) TFE α_{TEV} adopts a stable structure even in the absence of its interaction partner TFE β , and (ii) that the C138S mutation does not cause larger structural perturbation of TFE α_{TEV} .

The addition of zinc in equistoichiometric concentrations resulted in the disappearance of a number of residues in TFE α_{TEV} (Supplementary Figure S2A), which suggests that the structure of the protein is undergoing subtle changes upon zinc binding in agreement with CIU experiments. Importantly, the set of crosspeaks disappearing in WT TFE α_{TEV} contains the single crosspeak that is perturbed in the C138S variant. Several other crosspeaks disappearing upon zinc binding in the spectra of both WT TFE α_{TEV} and the C138S variant reflecting possibly additional residues involved in zinc binding (Supplementary Figure S2A and B). It should be noted that both CIU and NMR experiments are limited to salt concentrations that are far below the physiological conditions. However, the specific changes in

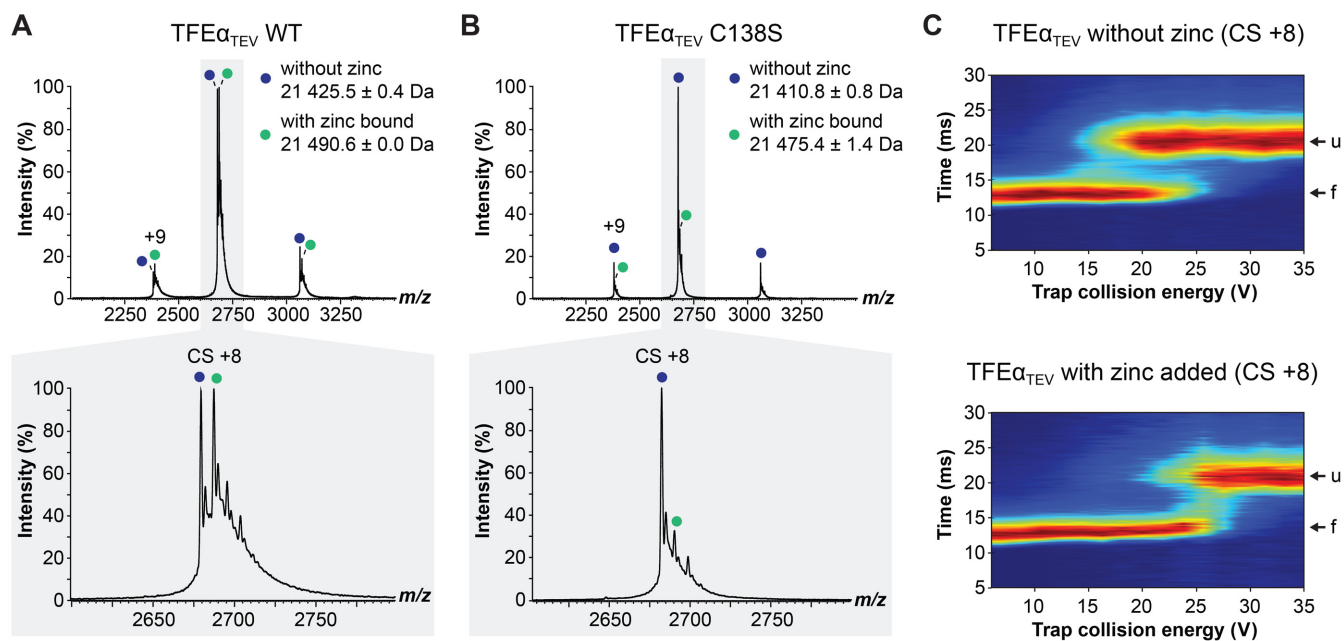


Figure 3. The degenerate zinc-ribbon domain of TFE α is capable of zinc binding. (A and B) nESI native mass spectrum of monomeric wild type TFE α_{TEV} (A) and TFE α_{TEV} C138S (B) after incubation with zinc. Charge state series of TFE α_{TEV} with and without zinc are labelled with green and blue filled circles, respectively. Highlighted in gray is a zoom over the region that includes charge state (CS) +8. (C) Collision induced unfolding experiment of TFE α_{TEV} with and without zinc addition. CS+8 was isolated using the quadrupole and activated in the trap collision cell using a CE ramp (6–36 V) prior to ion mobility separation. The ion mobility cell separated the protein into two populations corresponding to folded (f) and unfolded (u) states.

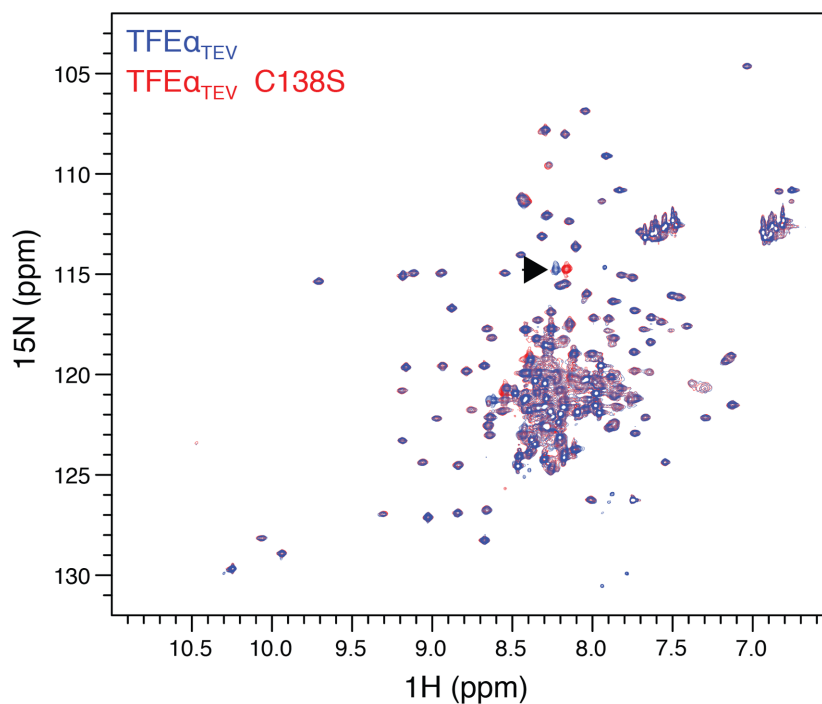


Figure 4. The C138S mutation does not cause larger structural perturbation of TFE α_{TEV} . Overlay 2D ^{15}N - 1H HSQC spectra of TFE α_{TEV} (blue) and C138S variant (red) without Zn^{2+} . The black arrow indicates the chemical shift perturbation caused by the C138S mutation. All data were recorded at 600 MHz, under the same conditions, number of scans and increments, at pH 7.5, 7% D_2O , 25°C.

the HSQC spectrum upon zinc binding and the 1:1 stoichiometry observed in the native mass spectra suggest that the zinc binding is specific.

Deletion of *tfeB* downregulates non-native genes differentially

In order to investigate the function of TFE *in vivo*, we attempted to delete the *tfeA* and *tfeB* genes in Hvo. One key feature of *H. volcanii* as archaeal model system is its genetic tractability, which includes a ‘pop-in/pop-out’ strategy for gene deletion using tryptophan and uracil auxotrophy (23). We were unable to delete *tfeA*, despite the use of a selectable *trpA*⁺ marker that significantly facilitates gene deletion (23), suggesting that *tfeA* is essential. However, the Hvo $\Delta tfeB$ strain H2644 proved to be viable but showed a severe growth phenotype, with a doubling time of 5.6 hrs compared to 2.2 h for the parental strain H53 (Figure 5A and B). Identical results were obtained with an independently-derived $\Delta tfeB$ strain H2645, and the growth defect in Hvo $\Delta tfeB$ H2644 was complemented by expression of the *tfeB* gene *in trans* (Figure 5C).

The Hvo $\Delta tfeB$ strains provided a unique opportunity to study a putative role of TFE β in the regulation of gene expression. In order to characterize the impact of the lack of TFE β on the haloarchaeal transcriptome, we isolated total RNA from the Hvo $\Delta tfeB$ H2644 and the wild-type H26 strain. H26 is *trpA*⁺ (23) and therefore its transcriptome will be more comparable to that of $\Delta tfeB::trpA$ ⁺ strain H2644 (Supplemental Table S3). RNA isolation was carried out during early exponential growth (OD₆₅₀ = 0.2) and subjected it to next generation RNA sequencing (RNA-seq). Consistent with the predicted role of TFE β as a basal transcription factor, the deletion of *tfeB* led to changes in relative mRNA abundance for 38% of all genes (1519 out of 3993 genes, $P_{\text{adj}} < 0.01$) (Supplemental Table S5). Approximately equal numbers of genes were up- or downregulated (781 and 738, respectively) with overall mild changes in relative transcript abundance (median values for log₂-fold change in abundance: 1.76 and -1.86 for up- and downregulated genes, respectively). Generally, slow growth is associated with reduced ribosome biosynthesis in pro- and eukaryotes (46). This holds true independent of whether slow growth results from genetic defects or poor carbon sources. We therefore analyzed the differential expression of the components of the basal translation machinery: ribosomal proteins, initiation and elongation factors. 42 out of 57 ribosomal proteins, three out of four elongation factors, and 4 out of 13 initiation factors showed differential expression ($P_{\text{adj}} < 0.01$). The majority of these genes show increased relative abundance (32 out of 49) (Figure 6A). Therefore, the changes in the transcriptome composition appear not to reflect a general slow growth phenotype. Instead these changes most likely reflect direct misregulation of gene expression as a result of *tfeB* deletion. In order to investigate the transcriptional response more systematically, we tested for the enrichment of specific functional classes of arCOGs (40) in the sets of differentially regulated genes. Genes involved in posttranslational modification, protein turnover and folding were enriched amongst the upregulated genes while genes involved in amino acid

transport and metabolism were underrepresented amongst downregulated genes (Fisher’s exact test, $P_{\text{adj}} < 0.05$) (Figure 6B and Supplemental Table S6). Interestingly, genes assigned to prophages and transposons were underrepresented among upregulated, but enriched among downregulated genes (Fisher’s exact test, $P_{\text{adj}} < 0.001$).

Hvo H26 harbours three megaplastids (pHV1, pHV3 and pHV4), with pHV4 stably integrated into the chromosome (19,47). Interestingly, the knockout of TFE β affects the genes residing on the megaplastids (in particular pHV1) more severely compared to genes encoded by the main chromosome (Table 1). Genes encoded by pHV1 differ significantly in their synonymous codon usage from the general codon usage in Hvo, suggesting that they are of non-native origin and have been acquired by lateral gene transfer (19). The notion that non-native genes are downregulated more severely than native ones is further supported by the predicted prophage residing in the main chromosome of Hvo. This region contains 43 open reading frames (ORFs HVO_2252 to HVO_2293), 32 of which have a decreased relative abundance in the $\Delta tfeB$ strain, while none of the remaining genes showed increased relative abundance. In order to test the effect of *tfeB* deletion on non-native genes more systematically, we plotted the rare codon frequency of genes (defined as codons with less than 10% synonymous codon usage genome-wide) against the log₂-fold change in relative transcript abundance (Figure 6C). Many genes with high frequency of rare codons appear to be downregulated and genes with a higher AT content than the genome average showed a similar correlation (Figure 6D). The systematic comparison of genes in the first quartile to those in the fourth quartile of rare codon frequency confirmed that genes with high frequency of rare codons show a greater tendency to be downregulated. The same result was obtained when genes were partitioned based on AT content (Fisher’s exact test $P < 0.001$ in both cases) (Supplementary Figure S3A).

To get further insight into how *tfeB* deletion globally affects the transcriptome, we calculated estimated transcript levels for the Hvo transcription units (Figure 6E). Downregulated genes appear to be associated with slightly higher transcript levels compared to upregulated ones (Wilcoxon rank sum test, $P < 0.001$). In summary, our results suggest that TFE β preferentially stimulates transcription of laterally transferred (non-native) genes.

In addition to downregulated genes, the deletion of TFE β also resulted in increased RNA expression levels that likely includes indirect effects such as a transcriptional response to suboptimal gene expression. For example, several genes encoding the basal transcription machinery including initiation factors and RNA polymerase subunits showed elevated RNA levels (Supplemental Table S5). Halophilic archaea utilize several TFB paralogues (20), which are divided into four evolutionary conserved clades (*Halobacterium* TfbA, TfbB/D/F, TfbC/G and TfbE) (48). Hvo possesses ten complete TFB paralogues. Three Hvo TFB paralogues belonging to the TfbB/D/F clade showed increased expression levels in the $\Delta tfeB$ strain, HVO_1052, HVO_1676 and HVO_B0285. Since TFE and TFB cooperate during OC formation, we surmise that Hvo compensates for inefficient transcription initiation due to the lack of TFE β by increas-

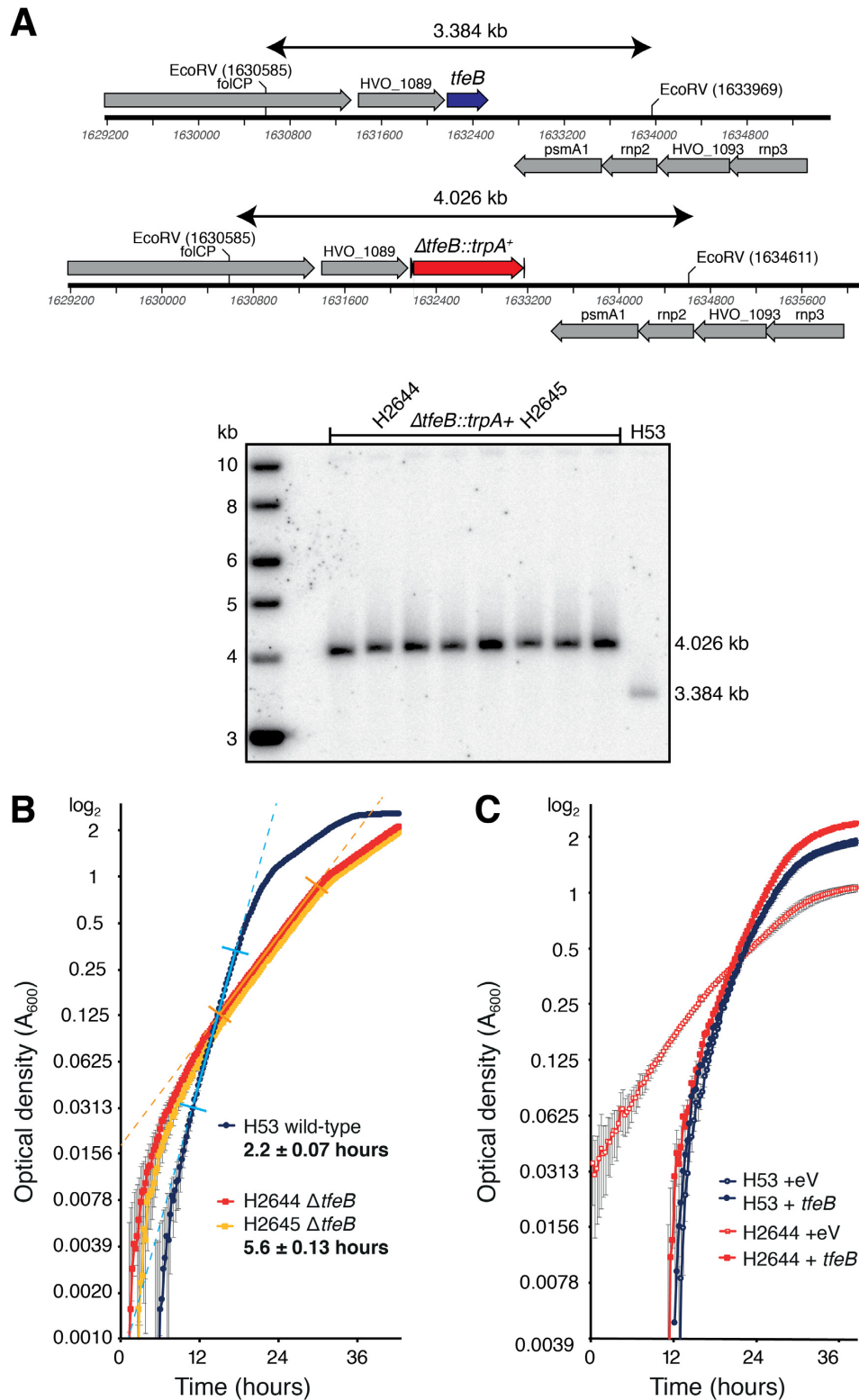


Figure 5. The *tfeB* gene encoding TFE β is non-essential in *H. volcanii*. (A) Southern blot of *EcoRV* digest confirming deletion of *tfeB* in H2644 and H2645. (B) Deletion of *tfeB* confers a slow-growth phenotype in H2644 and H2645, versus the parent strain H53. Data from the mean of four repeats (and standard error) is plotted on a \log_2 scale, the generation time in exponential phase is shown in bold (exponential growth phase is indicated by straight-line fit between perpendicular marks, $R^2 = 0.99$ for all strains). (C) The growth defect conferred by deletion of *tfeB* in H2644 is complemented by *in trans* expression of *tfeB* (eV, empty vector).

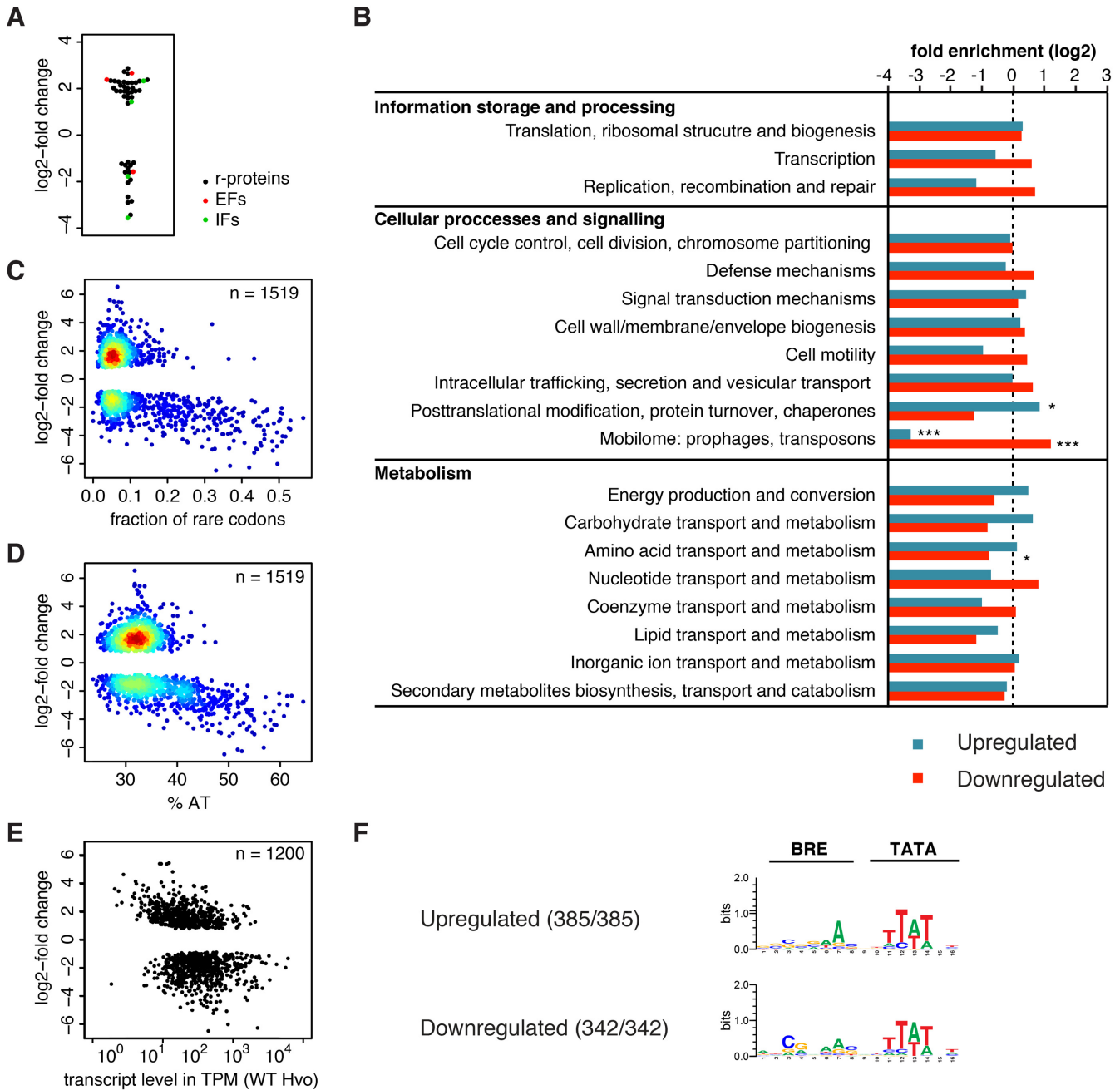


Figure 6. Deletion of *tfeB* leads to widespread misregulation of transcription. (A) log₂-fold change in relative transcript abundance ($P_{adj} < 0.01$) of genes coding for the basal translation machinery, ribosomal proteins (black), translation initiation factors (green) and elongation factors (red). (B) Analysis arCOG category distribution in significantly misregulated genes ($P_{adj} < 0.01$). Significant enrichment or underrepresentation is indicated (two-sided Fisher's exact test with Bonferroni correction for multiple testing) with * and *** denoting $P_{adj} < 0.05$ and 0.001 , respectively. (C and D) Heatscatter plot of rare codon frequency (defined as codons with less than 10% synonymous codon usage genome-wide) (C) and AT content of genes (D) against the log₂-fold change ($P_{adj} < 0.01$) in relative transcript abundance. (E) Plot showing the estimated transcript levels in parental strain H26 (transcripts per million, TPM, geometric mean for the two replicates) calculated for the transcription units of Hvo using RSEM (31) against the log₂-fold change in relative transcript abundance for the first cistron within the respective transcription unit ($P_{adj} < 0.01$). (F) BRE/TATA motif consensus in misregulated genes. In order to account for variable spacing between the TSS and BRE/TATA motifs we conducted a motif search using the MEME software version 4.11.4 (0 or 1 occurrence per sequence, 4 to 16 bp width, searching given strand only) (33). A background model based on the composition of the combined *H. volcanii* DS2 genome was employed. BRE/TATA motifs were discovered in all sequences.

Table 1. Changes in relative abundance in the $\Delta tfeB$ strain

	Total number of genes detected	Increased abundance	Decreased abundance
Main chromosome (excluding pHV4)*	2940	617	534
pHV1	75	1**	35**
pHV3	372	81	27**
pHV4*	606	82**	142

* Integrated into the main chromosome in laboratory strains, including H53 (47).

** $P < 0.001$ (Fisher's exact test compared to main chromosome).

ing the expression of a subset of TFB paralogues. Haloarchaeal TFB paralogues show preferences for different BRE motifs (22). A motif search did retrieve BRE/TATA motifs with slightly altered BRE consensus for up- and downregulated gene in the *tfeB* deletion strain, which might reflect the role of the upregulated TFB paralogues in the transcriptional response to *tfeB* deletion (Figure 6F and Supplementary Figure S3B).

Overall, our data suggest that *tfeB* deletion significantly alters the global transcriptome, particularly affecting non-native genes.

DISCUSSION

The identification and characterization of haloarchaeal TFE β variants provides us with a more complete picture of the evolution of TFE-like factors in archaea and eukaryotes. It corroborates the idea that dimeric TFE traces back to the last common ancestor of all archaea, and of eukaryotes. In *Euryarchaeota*, several lineages such as the *Thermococcales* and *Methanococcales* have lost the *tfeB* gene and this loss appears to coincide with a monomeric TFE α variant that seems to be functional on its own, as deduced from *in vitro* experiments (7–8,49). Other euryarchaeal lineages have retained the *tfeB* gene but we show that at least in Hvo it is non-essential. While basal transcription factors TBP and TFIIB facilitate RNAP function in eukaryotes, the archaeal homologues TBP and TFB have acquired a regulatory role, particularly in haloarchaea. We propose that the third archaeal initiation factor TFE similarly carries out a general function during OC formation and in addition stimulates transcription in a gene-specific fashion, in haloarchaea likely in conjunction with distinct TFB paralogues. The magnitude of TFE stimulation varies with the promoter context, and since the steady state levels of TFE β in Sso rapidly decrease in response to oxidative stress and upon entering stationary phase the genes controlled by promoters that respond stronger to TFE will be down regulated significantly more compared to genes under the control of relatively unresponsive promoters (2). Importantly, the structural FeS cluster of Sso TFE β that is crucial for all TFE functions is sensitive to oxidative damage, and its loss likely triggers the depletion of TFE β . Interestingly the Halobacteriales have lost the FeS cluster while maintaining the C-terminal domain, an event that is paralleled in *Saccharomyces* RNAPIII subunit C34 (2,13). The loss or degeneration of metal centres in Hvo TFE α/β may provide insight into the apparent loss of ZR and FeS cluster domains that occurred in the evolution of hRPC62 (C82) and TFIIE β , respectively. Similar to TFE β , the RNA polymerase subunit D in halophilic archaea appears to have lost

the structural FeS cluster (50), while other FeS cluster proteins such as XPD have retained them. What causes the loss of FeS clusters? It seems likely that general environmental factors could exert a selection pressure to drive the loss, including iron scarcity and -bioavailability in the high saline niche of haloarchaea, or elevated levels of oxidative stress in the environment. The functional adaptation of Hvo TFE β is also reflected in a remarkably acidic isoelectric point. Sso TFE β is a basic protein (predicted pI = 9.19) in line with the idea that its WH domain interacts with the non-template strand via electrostatic interactions on the RNAP surface during OC formation (2). In contrast, Hvo TFE β is acidic (predicted pI = 4.33). Under the physiological conditions of high salinity found in the cytoplasm of *H. volcanii*, electrostatic interactions with DNA are disfavoured. In addition, high salinity directly stabilizes dsDNA by counteracting repellent forces within the phosphate backbone of the two DNA strands. While the extreme natural environments of haloarchaea are likely to have driven the adaptation of haloarchaeal TFE β , different selective pressures might have led to the loss of the Fe-S cluster in its eukaryotic counterparts. This includes the emergence of different regulatory mechanisms for the RNAPII and III transcription machineries early in the evolution of eukaryotes. Another adaptation occurred in the Hvo TFE α subunit where the canonical ZR evolved into a degenerate form retaining only low affinity for zinc. This adaptation is specific to *Haloferax* and closely related species while the majority of haloarchaeal TFE α retained the four conserved cysteine residues forming canonical ZR domains. Nevertheless, it may be part of a general trend of reduced cysteine content in haloarchaeal proteomes (51,52).

The viability of the $\Delta tfeB$ deletion Hvo strain provided us with a unique opportunity to study the influence of TFE β on gene expression *in vivo*. In line with its function as basal transcription factor, approximately a third of all genes were misregulated in the $\Delta tfeB$ strain, notably non-native genes were affected more severely than native genes. Lateral gene transfer has played a key role in the evolution of Haloarchaea both at the root and during the subsequent diversification (29,32), and the Hvo genome harbours several large clusters of non-native genes (19). The impact of *tfeB* deletion on the expression of non-native genes shows that TFE β stimulates transcription of genes with promoters yet ill-adapted to the Hvo transcription machinery. The promoters of native genes coevolved with the host RNAP to form OCs readily including the region of the promoter that is initially melted during OC formation (IMR) and functionally interacts with TFE (2), while non-native promoters have not and apparently benefit to a greater extent from TFE β 's ability

to stimulate this process. However, this rationale is in apparent conflict with considerations concerning the negative selection pressure due to the potential harm arising from foreign genes, which are generally considered harmful (or neutral) for the organism (53). The greater dependency of these genes on TFE β could also be caused by different chromatinization of AT-rich genes posing a stronger barrier to transcription. While we cannot rule out that the different codon usage of foreign genes during translation feeds back into transcription initiation, there is no direct evidence for such a molecular mechanism.

Bacteria including *E. coli* are known to utilize the elongation factor NusG (Spt5 in archaea) and the termination factor Rho to suppress transcription of foreign DNA (54). A rigorous sequence analysis of the promoters that were misregulated in the $\Delta tfeB$ strain did not provide evidence for a specific DNA sequence bias in either up- nor downregulated promoters (Supplementary Figure S3B), congruent with the hypothesis that TFE β does not interact with DNA in a sequence specific manner. The underlying mechanisms for the promoter-specificity of TFE β likely involves additional factors including combinations of haloarchaeal TBP/TFB variants, as well as chromatin proteins and the topology of the promoters, all of which modulate the propensity of DNA strand separation during transcription initiation.

DATA AVAILABILITY

The accession number for the RNA-seq data reported in this paper is NCBI GEO: GSE101134.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

We thank Sunir Malla (Deep Seq, University of Nottingham NGS facility) for the DESeq analysis. We thank Thomas Fouqueau and Katherine Smollett (University College London) and David Grainger (University of Birmingham) for stimulating discussions. We would like to thank Jörg Soppa (Goethe University Frankfurt) for the kind gift of *H. volcanii* genomic DNA. We acknowledge the use of the ISMB Biological NMR Facility at UCL. The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

Author contributions: F.W., F.B., T.A. and D.A. conceived the study. F.B., N.Z. and M.Z. carried out the biochemical characterization of Hvo TFE; A.M.F. performed the NMR experiments; Z.S., T.P., K.T. performed the MS experiments; D.A. and T.A. generated the Hvo *tfeB* deletion strain and sequenced the transcriptome using NGS. F.B. and F.W. prepared the manuscript with input from all authors.

FUNDING

Research in the RNAP laboratory is funded by a Wellcome investigator award [WT096553MA to F.W.]; Research in TA laboratory is supported by the Biological Sciences Research Council (BBSRC) [BB/M001393/1]. Funding for open access charge: Wellcome Trust.

Conflict of interest statement. None declared.

REFERENCES

- Blombach, F., Smollett, K.L., Grohmann, D. and Werner, F. (2016) Molecular mechanisms of transcription initiation—structure, function, and evolution of TFE/TFIIE-like factors and open complex formation. *J. Mol. Biol.*, **428**, 2592–2606.
- Blombach, F., Salvadori, E., Fouqueau, T., Yan, J., Reimann, J., Sheppard, C., Smollett, K.L., Albers, S.V., Kay, C.W., Thalassinos, K. *et al.* (2015) Archaeal TFE α /beta is a hybrid of TFIIE and the RNA polymerase III subcomplex hRPC62/39. *Elife*, **4**, e08378.
- Meinhart, A., Blobel, J. and Cramer, P. (2003) An extended winged helix domain in general transcription factor E/TFIIE α . *J. Biol. Chem.*, **278**, 48267–48274.
- Miwa, K., Kojima, R., Obita, T., Ohkuma, Y., Tamura, Y. and Mizuguchi, M. (2016) Crystal structure of human general transcription factor TFIIE at atomic resolution. *J. Mol. Biol.*, **428**, 4258–4266.
- Plaschka, C., Hantsche, M., Dienemann, C., Burzinski, C., Pitzko, J. and Cramer, P. (2016) Transcription initiation complex structures elucidate DNA opening. *Nature*, **533**, 353–358.
- Blombach, F., Makarova, K.S., Marrero, J., Siebers, B., Koonin, E.V. and van der Oost, J. (2009) Identification of an ortholog of the eukaryotic RNA polymerase III subunit RPC34 in Crenarchaeota and Thaumarchaeota suggests specialization of RNA polymerases for coding and non-coding RNAs in Archaea. *Biol. Direct*, **4**, 39.
- Werner, F. and Weinzierl, R.O. (2005) Direct modulation of RNA polymerase core functions by basal transcription factors. *Mol. Cell. Biol.*, **25**, 8344–8355.
- Naji, S., Grünberg, S. and Thomm, M. (2007) The RPB7 orthologue E' is required for transcriptional activity of a reconstituted archaeal core enzyme at low temperatures and stimulates open complex formation. *J. Biol. Chem.*, **282**, 11047–11057.
- Feaver, W.J., Henry, N.L., Bushnell, D.A., Sayre, M.H., Brickner, J.H., Gileadi, O. and Kornberg, R.D. (1994) Yeast TFIIE. Cloning, expression, and homology to vertebrate proteins. *J. Biol. Chem.*, **269**, 27549–27553.
- Chiannilkulchai, N., Stalder, R., Riva, M., Carles, C., Werner, M. and Sentenac, A. (1992) RPC82 encodes the highly conserved, third-largest subunit of RNA polymerase C (III) from *Saccharomyces cerevisiae*. *Mol. Cell. Biol.*, **12**, 4433–4440.
- Stettler, S., Mariotte, S., Riva, M., Sentenac, A. and Thuriaux, P. (1992) An essential and specific subunit of RNA polymerase III (C) is encoded by gene RPC34 in *Saccharomyces cerevisiae*. *J. Biol. Chem.*, **267**, 21390–21395.
- Carter, R. and Drouin, G. (2010) The increase in the number of subunits in eukaryotic RNA polymerase III relative to RNA polymerase II is due to the permanent recruitment of general transcription factors. *Mol. Biol. Evol.*, **27**, 1035–1043.
- Hoffmann, N.A., Jakobi, A.J., Moreno-Morcillo, M., Glatt, S., Kosinski, J., Hagen, W.J., Sachse, C. and Muller, C.W. (2015) Molecular structures of unbound and transcribing RNA polymerase III. *Nature*, **528**, 231–236.
- Grohmann, D., Nagy, J., Chakraborty, A., Klose, D., Fielden, D., Ebricht, R.H., Michaelis, J. and Werner, F. (2011) The initiation factor TFE and the elongation factor Spt4/5 compete for the RNAP clamp during transcription initiation and elongation. *Mol. Cell*, **43**, 263–274.
- Schulz, S., Gietl, A., Smollett, K., Tinnfeld, P., Werner, F. and Grohmann, D. (2016) TFE and Spt4/5 open and close the RNA polymerase clamp during the transcription cycle. *Proc. Natl. Acad. Sci. U.S.A.*, **113**, E1816–E1825.
- He, Y., Fang, J., Taatjes, D.J. and Nogales, E. (2013) Structural visualization of key steps in human transcription initiation. *Nature*, **495**, 481–486.
- Williams, L.D. and Maher, L.J. 3rd (2000) Electrostatic mechanisms of DNA deformation. *Annu. Rev. Biophys. Biomol. Struct.*, **29**, 497–521.
- Holstege, F.C., Tantin, D., Carey, M., van der Vliet, P.C. and Timmers, H.T. (1995) The requirement for the basal transcription factor IIE is determined by the helical stability of promoter DNA. *EMBO J.*, **14**, 810–819.
- Hartman, A.L., Norais, C., Badger, J.H., Delmas, S., Haldenby, S., Madupu, R., Robinson, J., Khouri, H., Ren, Q., Lowe, T.M. *et al.* (2010)

- The complete genome sequence of *Haloflex volcanii* DS2, a model archaeon. *PLoS One*, **5**, e9605.
20. Facciotti, M.T., Reiss, D.J., Pan, M., Kaur, A., Vuthoori, M., Bonneau, R., Shannon, P., Srivastava, A., Donohoe, S.M., Hood, L.E. *et al.* (2007) General transcription factor specified global gene regulation in archaea. *Proc. Natl. Acad. Sci. U.S.A.*, **104**, 4630–4635.
 21. Coker, J.A. and DasSarma, S. (2007) Genetic and transcriptomic analysis of transcription factor genes in the model halophilic Archaeon: coordinate action of TbpD and TfbA. *BMC Genet.*, **8**, 61.
 22. Seitzer, P., Wilbanks, E.G., Larsen, D.J. and Facciotti, M.T. (2012) A Monte Carlo-based framework enhances the discovery and interpretation of regulatory sequence motifs. *BMC Bioinformatics*, **13**, 317.
 23. Allers, T., Ngo, H.P., Mevarech, M. and Lloyd, R.G. (2004) Development of additional selectable markers for the halophilic archaeon *Haloflex volcanii* based on the *leuB* and *trpA* genes. *Appl. Environ. Microbiol.*, **70**, 943–953.
 24. Norais, C., Hawkins, M., Hartman, A.L., Eisen, J.A., Myllykallio, H. and Allers, T. (2007) Genetic and physical mapping of DNA replication origins in *Haloflex volcanii*. *PLoS Genet.*, **3**, e77.
 25. Parkhomchuk, D., Borodina, T., Amstislavskiy, V., Banaru, M., Hallen, L., Krobitsch, S., Lehrach, H. and Soldatov, A. (2009) Transcriptome analysis by strand-specific sequencing of complementary DNA. *Nucleic Acids Res.*, **37**, e123.
 26. Bolger, A.M., Lohse, M. and Usadel, B. (2014) Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics*, **30**, 2114–2120.
 27. Trapnell, C., Pachter, L. and Salzberg, S.L. (2009) TopHat: discovering splice junctions with RNA-Seq. *Bioinformatics*, **25**, 1105–1111.
 28. Love, M.I., Huber, W. and Anders, S. (2014) Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biol.*, **15**, 550.
 29. Ammar, R., Torti, D., Tsui, K., Gebbia, M., Durbic, T., Bader, G.D., Giaever, G. and Nislow, C. (2012) Chromatin is an ancient innovation conserved between Archaea and Eukarya. *Elife*, **1**, e00078.
 30. Babski, J., Haas, K.A., Nather-Schindler, D., Pfeiffer, F., Forstner, K.U., Hammelmann, M., Hilker, R., Becker, A., Sharma, C.M., Marchfelder, A. *et al.* (2016) Genome-wide identification of transcriptional start sites in the haloarchaeon *Haloflex volcanii* based on differential RNA-Seq (dRNA-Seq). *BMC Genomics*, **17**, 629.
 31. Li, B. and Dewey, C.N. (2011) RSEM: accurate transcript quantification from RNA-Seq data with or without a reference genome. *BMC Bioinformatics*, **12**, 323.
 32. Quinlan, A.R. and Hall, I.M. (2010) BEDTools: a flexible suite of utilities for comparing genomic features. *Bioinformatics*, **26**, 841–842.
 33. Bailey, T.L., Boden, M., Buske, F.A., Frith, M., Grant, C.E., Clementi, L., Ren, J., Li, W.W. and Noble, W.S. (2009) MEME SUITE: tools for motif discovery and searching. *Nucleic Acids Res.*, **37**, W202–W208.
 34. Crooks, G.E., Hon, G., Chandonia, J.M. and Brenner, S.E. (2004) WebLogo: a sequence logo generator. *Genome Res.*, **14**, 1188–1190.
 35. Pringle, S.D., Giles, K., Wildgoose, J.L., Williams, J.P., Slade, S.E., Thalassinou, K., Bateman, R.H., Bowers, M.T. and Scrivens, J.H. (2007) An investigation of the mobility separation of some peptide and protein ions using a new hybrid quadrupole/travelling wave IMS/oa-ToF instrument. *Int. J. Mass Spectrom.*, **261**, 1–12.
 36. Hernandez, H. and Robinson, C.V. (2007) Determining the stoichiometry and interactions of macromolecular assemblies from mass spectrometry. *Nat. Protoc.*, **2**, 715–726.
 37. Sivalingam, G.N., Yan, J., Sahota, H. and Thalassinou, K. (2013) Amphitrite: A program for processing travelling wave ion mobility mass spectrometry data. *Int. J. Mass Spectrom.*, **345–347**, 54–62.
 38. Delaglio, F., Grzesiek, S., Vuister, G.W., Zhu, G., Pfeifer, J. and Bax, A. (1995) NMRPipe: a multidimensional spectral processing system based on UNIX pipes. *J. Biomol. NMR*, **6**, 277–293.
 39. Vranken, W.F., Boucher, W., Stevens, T.J., Fogh, R.H., Pajon, A., Llinas, M., Ulrich, E.L., Markley, J.L., Ionides, J. and Laue, E.D. (2005) The CCPN data model for NMR spectroscopy: development of a software pipeline. *Proteins*, **59**, 687–696.
 40. Makarova, K.S., Wolf, Y.I. and Koonin, E.V. (2015) Archaeal clusters of orthologous genes (arCOGs): an update and application for analysis of shared features between Thermococcales, Methanococcales, and Methanobacteriales. *Life (Basel)*, **5**, 818–840.
 41. Berkovitch, F., Nicolet, Y., Wan, J.T., Jarrett, J.T. and Drennan, C.L. (2004) Crystal structure of biotin synthase, an S-adenosylmethionine-dependent radical enzyme. *Science*, **303**, 76–79.
 42. Imlay, J.A. (2006) Iron-sulphur clusters and the problem with oxygen. *Mol. Microbiol.*, **59**, 1073–1082.
 43. Okuda, M., Tanaka, A., Arai, Y., Satoh, M., Okamura, H., Nagadoi, A., Hanaoka, F., Ohkuma, Y. and Nishimura, Y. (2004) A novel zinc finger structure in the large subunit of human general transcription factor TFIIE. *J. Biol. Chem.*, **279**, 51395–51403.
 44. Bell, S.D., Brinkman, A.B., van der Oost, J. and Jackson, S.P. (2001) The archaeal TFIIE α homologue facilitates transcription initiation by enhancing TATA-box recognition. *EMBO Rep.*, **2**, 133–138.
 45. Nyon, M.P., Prentice, T., Day, J., Kirkpatrick, J., Sivalingam, G.N., Levy, G., Haq, I., Irving, J.A., Lomas, D.A., Christodoulou, J. *et al.* (2015) An integrative approach combining ion mobility mass spectrometry, X-ray crystallography, and nuclear magnetic resonance spectroscopy to study the conformational dynamics of α 1-antitrypsin upon ligand binding. *Protein Sci.*, **24**, 1301–1312.
 46. Regenberg, B., Grotkjaer, T., Winther, O., Fausboll, A., Akesson, M., Bro, C., Hansen, L.K., Brunak, S. and Nielsen, J. (2006) Growth-rate regulated genes have profound impact on interpretation of transcriptome profiling in *Saccharomyces cerevisiae*. *Genome Biol.*, **7**, R107.
 47. Hawkins, M., Malla, S., Blythe, M.J., Nieduszynski, C.A. and Allers, T. (2013) Accelerated growth in the absence of DNA replication origins. *Nature*, **503**, 544–547.
 48. Becker, E.A., Seitzer, P.M., Tritt, A., Larsen, D., Krusor, M., Yao, A.I., Wu, D., Madern, D., Eisen, J.A., Darling, A.E. *et al.* (2014) Phylogenetically driven sequencing of extremely halophilic archaea reveals strategies for static and dynamic osmo-response. *PLoS Genet.*, **10**, e1004784.
 49. Hanzelka, B.L., Darcy, T.J. and Reeve, J.N. (2001) TFE, an archaeal transcription factor in *Methanobacterium thermoautotrophicum* related to eucaryal transcription factor TFIIE α . *J. Bacteriol.*, **183**, 1813–1818.
 50. Lessner, F.H., Jennings, M.E., Hirata, A., Duin, E.C. and Lessner, D.J. (2012) Subunit D of RNA polymerase from *Methanosarcina acetivorans* contains two oxygen-labile [4Fe-4S] clusters: implications for oxidant-dependent regulation of transcription. *J. Biol. Chem.*, **287**, 18510–18523.
 51. Paul, S., Bag, S.K., Das, S., Harvill, E.T. and Dutta, C. (2008) Molecular signature of hypersaline adaptation: insights from genome and proteome composition of halophilic prokaryotes. *Genome Biol.*, **9**, R70.
 52. Ebrahimie, E., Ebrahimie, M., Sarvestani, N.R. and Ebrahimie, M. (2011) Protein attributes contribute to halo-stability, bioinformatics approach. *Saline Syst.*, **7**, 1.
 53. Soucy, S.M., Huang, J. and Gogarten, J.P. (2015) Horizontal gene transfer: building the web of life. *Nat. Rev. Genet.*, **16**, 472–482.
 54. Cardinale, C.J., Washburn, R.S., Tadigotla, V.R., Brown, L.M., Gottesman, M.E. and Nudler, E. (2008) Termination factor Rho and its cofactors NusA and NusG silence foreign DNA in *E. coli*. *Science*, **320**, 935–938.
 55. Lefèvre, S., Dumay-Odelot, H., El-Ayoubi, L., Budd, A., Legrand, P., Pinaud, N., Teichmann, M. and Fribourg, S. (2011) Structure-function analysis of hRPC62 provides insights into RNA polymerase III transcription initiation. *Nat. Struct. Mol. Biol.*, **18**, 352–358.