RESEARCH ARTICLE

Metabolic and Chaperone Gene Loss Marks the Origin of Animals: Evidence for Hsp104 and Hsp78 Chaperones Sharing Mitochondrial Enzymes as Clients

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Abstract

The evolution of animals involved acquisition of an emergent gene repertoire for gastrulation. Whether loss of genes also co-evolved with this developmental reprogramming has not yet been addressed. Here, we identify twenty-four genetic functions that are retained in fungi and choanoflagellates but undetectable in animals. These lost genes encode: (i) sixteen distinct biosynthetic functions; (ii) the two ancestral eukaryotic ClpB disaggregases, Hsp78 and Hsp104, which function in the mitochondria and cytosol, respectively; and (iii) six other assorted functions. We present computational and experimental data that are consistent with a joint function for the differentially localized ClpB disaggregases, and with the possibility of a shared client/chaperone relationship between the mitochondrial Fe/S homoaconitase encoded by the lost LYS4 gene and the two ClpBs. Our analyses lead to the hypothesis that the evolution of gastrulation-based multicellularity in animals led to efficient extraction of nutrients from dietary sources, loss of natural selection for maintenance of energetically expensive biosynthetic pathways, and subsequent loss of their attendant ClpB chaperones.

Introduction

By the time of the last common ancestor of animals, the uniquely animal process of gastrulation had evolved. Gastrulation, while diverse in form, is a highly regulated re-arrangement and migration of cells producing a layered multicellular body with an internalized “endoderm”-derived digestion system. Many post-genomic studies have detailed the evolutionary expansion of eukaryotic transcription factor (TF) families and the emergence of new developmental genes during this major evolutionary transition [1–5]; these genes include those controlling the gastrulation program, which is shared from humans to sponges [6,7]. However, identification of pre-metazoan gene losses may be equally important because they may indicate the context, origins, and constraints associated with new transcription programs.

No study has yet focused on systematically identifying genes that are widely retained in the closest relatives of animals but undetectable in animals themselves. Such studies require

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resources and methodologies that have not been available or mature enough until recently. These include: (i) relevant genome assemblies; (ii) robust, accurate, orthology-calling pipelines based on reliable gene annotations; and (iii) powerful gene association engines and data sets from model genetic systems that would allow candidate losses to be placed into genetic pathways and regulatory programs. Now, assembled genomes are increasingly available from such early-branching animal lineages as sponges (e.g., *Amphimedon queenslandica* [4]), ctenophores (*Mnemiopsis leidyi* [8]), placozoans (*Trichoplax adhaerens* [9]), as well diverse bilaterians, including chordate and non-chordate deuterostomes and ecdysozoan and lophotrochozoan protostomes [10]. Also available are genomes of animal’s closest eukaryotic relatives, including its sister clade of choanoflagellates (*Monosiga brevicollis* [11] and *Salpingoeca rosetta* [12]), as well as filastereans (e.g., *Capsaspora owczarzaki* [13]), and diverse fungi from ascomycetes, basidiomycetes, and chytridiomycetes (e.g., *Batrachochytrium dendrobatidis*). In addition, Ensembl’s Maximum-Likelihood based orthology-calling pipeline EnsemblCompara [14] and BioMart query engine [15] can be used to identify candidate lost genes, which can then be investigated and confirmed with formal phylogenetic analysis on a gene by gene basis. Finally, powerful algorithms have been designed to query genetic, transcriptomic, proteomic, and sub-cellular localization data sets and to measure meaningful enrichment of functional categories [16,17]. Such association algorithms require large amounts of data and data types for sufficient power. Arguably, the most comprehensive data is available for the yeast, *Saccharomyces cerevisiae*. For this reason, this organism is the favored system for understanding at a quantitative and analytical level how genome-encoded functions interact to produce complex phenotypes [18,19].

Here we use comparative and integrative genomic approaches to identify and analyze a set of eukaryotic genes that are retained and detectable in the closest-living relatives of animals, but are no longer detectable in animals themselves. Interpreted parsimoniously, these undetectable genes correspond to genetic functions lost in the stem-metazoan lineage, which begins just after the latest common ancestor of animals and choanoflagellates and ends just before the latest common ancestor of animals. We find that the majority of these genes correspond to lost biosynthetic pathways, including biosynthesis of L-histidine, L-lysine, riboflavin, and L-tryptophan. Intriguingly, we find specific connections between the two lost lysine biosynthetic genes, *LYS4* and *LYS20*, and the lost eukaryotic ClpB chaperones encoded by *HSP78* and *HSP104*, which have roles in protein folding in the mitochondria and cytoplasm, respectively. Computational and experimental data suggest that the two ClpB chaperones may work together. In this study we investigate the hypothesis that, while *Hsp78* and *Hsp104* are differentially compartmentalized, they may share a subset of clients in nuclear-encoded, cytoplasmically-translated, mitochondrialy-imported proteins. Tandem affinity purification (TAP) of chaperones and identification of interacting proteins with mass spectrometry (MS) (TAP-MS) data in yeast reveal that the *Hsp104* and *Hsp78* interact with *Lys4* and *Lys20* but not with the proteins encoded by the other lost genes. Taken together our results are consistent with the hypothesis that *Lys4* and *Lys20* are prone to misfolding and require ClpB chaperones after thermal stress and suggest that some of the genes identified in our study do not necessarily represent independent losses but may instead reflect a series of dependent events. Our genomic approach and results are likely to improve our understanding of evolutionary origins and unique nature of animal metabolism.

**Results**

**Identification of twenty-five genes retained in choanoflagellates but not animals**

To identify protein-coding genes retained in the closest eukaryotic relatives of animals but which are not otherwise detectable in animals (henceforth the “lost genes”), we made two
methodological decisions. First, we chose *Saccharomyces cerevisiae* as a reference genome despite the possibility that we would not identify some lost genes associated with derived traits in the budding yeast. However, by using the budding yeast we avail ourselves of the large number of existing genetic, transcriptomic, proteomic, and sub-cellular localization data sets. Second, to avoid false-positive gene losses due to deficiencies in assembled genomes, we sampled diverse non-animal genomes without over-sampling any particular taxon, and focused on the fewest genomes possible without sacrificing taxonomic diversity. For Dikarya fungi we worked with the ascomycetes, *S. cerevisiae* and *Schizosaccharomyces pombe* genomes; for the basidiomycetes, *Cryptococcus neoformans* and *Puccinia graminis* (Ensembl BioMart); and for chytridiomycetes, *Batrachochytrium dendrobatidis* (JGI). For Filasterea, we chose the *Capsaspora owczarzaki* (Broad Institute) genome [13]. Last, for choanoflagellates we used *Monosiga brevicollis* (Joint Genome Institute, JGI) and the colonial species *Salpingoeca rosetta* (Broad Institute) [11,20].

To generate a manageable candidate gene list we used Ensembl’s BioMart query engine [14,15,21] to iteratively identify orthologs that are present in the four listed genomes of Dikarya (Fig. 1) but absent in the two bilaterians, *Drosophila melanogaster* and *Homo sapiens*, each of which has an excellent genome assembly with few gaps. We also accepted lineage-specific duplications of a single shared ancestral gene to avoid losing bona fide candidates arising from whole genome duplications in any one lineage. Of the 6,692 protein-coding genes in *S. cerevisiae* (Ensembl R64–1–1), 802 genes survived this initial screen. We then queried the Joint Genome Institute’s (JGI) BLASTP server (genome.jgi.doe.gov) with each yeast protein to identify those for which the predicted protein-coding sequences from the choanoflagellate *M. brevicollis* (Monbr1_best_proteins) matched the yeast proteins with a relatively stringent BLASTP expect value threshold (E-value lower (better) than 1.0E-30). This reduced our list to 210 yeast proteins. We used a threshold E-value of 1.0E-10 to remove yeast proteins that either had no matches or were not detectable for the colonial choanoflagellate *S. rosetta* and the filasterean *C. owczarzaki*. We also removed those that had E-values in animals smaller (i.e., better) than any of the three listed non-animal holozoans (the choanoflagellates and filasterean) (Fig. 1).

For animals, we used the sponge *Amphimedon queenslandica* (NCBI) [4], the ctenophore *Mnemiopsis leidyi* (NHGRI) [8], the placozoan *Trichoplax adhaerens* (NCBI) [9], and the lophotrochozoans *Lottia gigantea*, *Capitella teleta*, and *Helobdella robusta*, all of which have complete genome sequence assemblies [10]. With these genomic comparisons, we find that there are twenty-seven yeast genes that are largely retained outside of animals in the genomes we analyzed (Fig. 1, and S1 Table). Six of these genes (*TRR1/TRR2*, *LYS20/LYS21*, and *SCT1/GPT2*) correspond to lineage-specific duplications in yeast (connected genes in Fig. 1). Thus, we find twenty-four genetic functions that are definitively lost in animals or at least not retained at the same level of conservation as they are in non-metazoans.

A heat map of the E-values for the best BLASTP hits in each of the genomes shows that most “lost” candidate genes are undetectable in metazoans, but that a few have matches with poor E-values relative to the non-metazoans (Fig. 1B, S1 Table), or else could be shown to be environmental contaminants or horizontal transfers from distinct taxa (Fig. 1B, taxonomic names given). Some were found to be non-orthologous sequences that are members of other gene clades in a superfamilly defined by a common domain (Fig. 1B, gene names given). In only a single case (*HIS2*) was there a weak match from an animal genome (*Trichoplax adhaerens*) that was most closely-related to the orthologous gene from choanoflagellates or the filasterean *Capsaspora*. The single possible hit in *Nematostella vectensis* for *HIS7* (Fig. 1) corresponds to the proteobacterial version of this gene and not a gene related to opisthokont or eukaryotic *HIS7*, and similar results were found for *TRP5* and others as indicated (Fig. 1B). One gene labeled “clpB” in many animals and which matches yeast Hsp104 and Hsp78, is actually a fusion
**Fig 1. Twenty-five genetic functions undetectable in animals.** (A) Heat map of twenty-seven genes (first twenty-seven rows with gene names) from the budding yeast *Saccharomyces cerevisiae* (*S*. *cer*.) and their orthologs in related organisms with whole genome sequence assemblies and their closest matches in animals. Six of the twenty-seven yeast genes are the result of lineage-specific duplications in yeast (indicated by dark gray brackets, left of table), leaving twenty-four genetic functions either lost or not measurably conserved in metazoans. The heat map is based on the BLASTP expect values of the best match to the indicated yeast gene (see key, lower right of table). Species abbreviations: *S*. *cer*, *Saccharomyces cerevisiae*; *S*. *pom*, *Schizosaccharomyces pombe*; *C*. *neo*, *Cryptococcus neoformans*; *P*. *gra*, *Puccinia graminis*; *B*. *den*, *Batrachochytrium dendrobatidis*; *C*. *owc*, *Capsaspora owczarzaki*; *M*. *bre*, *Monosiga brevicollis*; *S*. *ros*, *Salpingoeca rosetta*; *A*. *que*, *Amphimedon queenslandica*; *T*. *adh*, *Trichoplax adhaerens*; *M*. *lei*, *Mnemiopsis leidyi*; *N*. *vec*, *Nematostella vectensis*; *L*. *gig*, *Lottia gigantea*; *C*. *tel*, *Capitella teleta*; *H*. *rob*, *Helobdella robusta*; *D*. *mel*, *Drosophila melanogaster*; and *H*. *sap*, *Homo sapiens*. Other abbreviations: Chytridio., Chytridiomycota; Placo., Placozoa; Ctenoph., Ctenophora; Ecdyso., Ecdysozoa; and Chor., Chordata. The last row is from an alternative screen for lost animal genes (see text) in which Amoebozoa PFAM domains that are absent in animals were identified (five genes already identified plus one new gene, *PURB*). (B) Close-up of the lost animal gene columns from panel A. This table shows either the taxonomic origin of a weak animal match, or the name of the gene if it is not directly related (i.e., not orthologous) to the named yeast gene (first column) for all matches with BLASTP expectation values < 1e-20. Most of the animal matches to the candidate lost genes correspond to lineage-specific horizontal gene transfers and/or environmental contaminants from unrelated (non-opisthokont) clades. Some matches correspond to non-orthologous genes such as *ANKCLP* in the case of the eukaryotic clpB genes *HSP78* and *HSP104*, or *ACO1* in the case of *LYS4* (Fig. 3). Only *HIS2* might have been lost soon after early animal radiation based on the presence of a weak match in *Trichoplax adhaerens*. This gene is most similar to the version from *Capsaspora* and not to either of the two choanoflagellates, so it is of uncertain origin.

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gene between an N-terminal ankyrin domain and a C-terminal partial sequence of a bacterial clp gene (S1 Fig.). Phylogenetic analysis of this “ANKCLP” fusion gene (S1 Fig.) indicates that it arose in the stem-holozoan lineage such that it still occurs alongside the HSP104 and HSP78 genes of both choanoflagellates.

We found that the closest matches in animal sequences to either Hsp104 or Hsp78 are partial fragments of bacterial ClpB in Amphimedon (Fig. 1) or the unrelated bacterial ClpC in Nematostella (S1 Fig.). We used Bayesian Inference and sampled mixed amino acid substitution models to infer a ClpB/Hsp78/Hsp104 phylogenetic tree consisting of the fungal and holozoan sequences as well as the best matches to the 456 amino acid Amphimedon sequence, which is only half as long as the proteins in the eukaryotic and bacterial ClpB clades (Fig. 2). Our results definitively place the predicted partial peptide sequence from Amphimedon in a ClpB clade alongside orthologs from γ-proteobacteria (Fig. 2). The sequences most closely related to the Amphimedon fragment are from bacterial endosymbionts of marine tube worms (Fig. 2). This suggests that this sequence is either an environmental contaminant or a partial horizontal gene transfer from endosymbiotic bacteria.

The second best match to Hsp104 in metazoans was an even shorter peptide sequence of 404 amino acids from Nematostella vectensis (S1 Fig.). This predicted peptide sequence most closely matches ClpC peptides from flavobacteria when the entire database is queried. Queries of Hsp104 sequences to the taxonomically unrestricted animal database reveal other spurious hits, which can be discounted for similar reasons (S1B Fig.). Thus, these results suggest that animals truly lost both Hsp104 and Hsp78.

Non-orthology for some weak matches in animal genomes is seen for the gene LYS4, which encodes mitochondrial homoaconitase. The closest match in animals to LYS4 is the ACO1 gene, which encodes the cytoplasmic aconitase of the tricarboxylic acid cycle and is broadly retained in both animals and non-animals as expected (Fig. 3A, Bayesian Inference; and Fig. 3B, Neighbor-Joining). For most lost genes, there are neither orthologs nor distantly related homologs in animals even when such genes are present in bacteria (e.g., Fig. 3C, LYS20).

To summarize the results of our gene loss approach, we find there are at least twenty-four highly retained genetic functions that are specifically lost in animals. To determine the extent to which this query strategy for lost animal genes is biased by the requirement for presence in fungi, we performed a second independent query focused on gene families not present in Saccharomyces cerevisiae, a key species in our first analysis. We identified 14 PFAM families present in the social slime mold Dictyostelium discoideum and the choanoflagellate Monosiga brevicollis, but not in animals (initially Homo sapiens, Drosophila melanogaster, Nematostella vectensis, Trichoplax adhaerens, and Amphimedon queenslandica) and the yeast S. cerevisiae. We used the Dictyostelium predicted protein sequences (19 peptide sequences) corresponding to these 14 PFAM domains as queries in a BLAST analysis to identify homologs in choanoflagellates. The Dictyostelium gene PURB (PF08238), which encodes an adenylosuccinate lyase (467 amino acids), is present in Monosiga brevicollis (2e-129), Salpingoeca rosetta (1e-134), and Capsaspora owczarkii (8e-169), but NOT in fungi or animals. A second Dictyostelium gene PHAZ (PF10503), which encodes a putative polyhydroxybutyrate depolymerase (341 amino acids), is present as two copies (paralogs) in each choanoflagellate genome (E-values range from 1e-87 to 4e-76), but we found this to have matches in the non-eumetazoan animal lineages. In summary, this alternate strategy identified six lost animal genes (GDH2, HSP104, HSP78, PAN6, PURB, TRR1/2) five of which were identified by the fungi-based screen (see Fig. 1A, right-hand column). In total, we found twenty-five distinct gene functions maintained in choanoflagellates (and in fungi or slime mold) but definitively lost in animals (S1 File).
Fig 2. Phylogenetic analyses confirming absence of intact clpB genes in animals. (A) Phylogenetic inference using Bayesian MCMC methods shows that the closest matching sequence in known animal genome sequences is a partial ClpB fragment in the sponge Amphimedon queenslandica. This fragment does not contain both nucleotide binding domains (NBDs) seen in bacterial or eukaryotic (Hsp78 and Hsp104) chaperone-client genes.
sequences, which are at least 800 amino acids long. The top three sequence hits in GenBank databases using the Amphimedon ClpB fragment as query are included. Two of these are intact ClpB protein sequences from bacterial (γ-proteobacteria) endosymbionts of marine tubeworms. Topological convergence was achieved after 300,000 generations with a relative burn-in phase of 25% (MrBayes) [80–82]. Mixed models were tested, but WAG [83] was the sole model favored upon completion. Average standard deviation of split frequencies was < 0.002. The tree was rooted between the bacterial clpB clade, including mitochondrial Hsp78, and the eukaryotic Hsp104 clade. (B) Maximum likelihood phylogenetic analysis of the ClpB family with choanoflagellates and chytridiomycetes (Batrachochytrium dendrobatidis and Gonapodya prolifera) representing fungal lineages shows that Hsp78 is the mitochondrially inherited clpB gene based on its sister grouping with α-proteobacteria. Dikarya sequences were omitted to simplify the tree. Numbers indicate bootstrap support from 500 replicates using MEGAS [85]. (C) A simple, unrooted, neighbor-joining tree of the highly-conserved clpB genes across all of life shows that the ClpB functions are maintained in all major domains of life and in all endosymbiotic eukaryotic compartments. The ubiquity of the clpB genes makes the loss in animals more striking.

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Animal-specific co-loss of the ClpB chaperones, Hsp78 and Hsp104

Both HSP104 and HSP78 are nuclear encoded eukaryotic orthologs of bacterial clpB, and encode AAA+ (ATPases Associated with diverse cellular Activities) family ATPases with two P-loop nucleotide-binding domains (NBDs). Together with an N terminal domain, the two NBDs form a three tier hexameric ring with a central pore through which misfolded, aggregated peptides are translocated prior to refolding [22,23]. Hsp78 localizes and functions in the mitochondria [24–26], while Hsp104 localizes to the cytoplasm and nucleus [27]. Because of Hsp104’s role in amyloid/cross-β disaggregation in yeast [28], the reason for the loss of the highly-conserved gene HSP104 in animals has been a source of much speculation [29].

In eukaryotes, the nuclear encoded HSP78 gene was acquired from the mitochondrial ancestor [30]. Our phylogenetic analyses repeatedly confirm that choanoflagellates, like fungi, have both HSP78 and HSP104, which correspond to the mitochondrial and eukaryotic clpB orthologs that are absent in animals (Fig. 2B; also see close relationship of HSP78 clade to α-proteobacterial clpB). The loss of these genes in the stem-metazoan lineage is more remarkable when considering that all organisms and organelles (e.g., eukaryotic mitochondrion, plant chloroplast), except those in animals, appear to retain ClpB functions, even when such genes are transferred to the nucleus (Fig. 2C).

HSP104 and HSP78 are co-regulated by a specialized cis-element, HSE4

To explore the possibility that joint loss of the ClpB orthologs reflects a joint requirement and/or a joint function for the proteins, we examined their regulation and, as described below, we also assessed their double mutant phenotype. To test the presence of shared clpB regulatory signatures, we initially focused on S. cerevisiae, for which predictions of co-regulation could be confirmed independently by gene expression meta-analyses of the large number of published transcriptome studies in yeast [31]. Regulatory motifs common to the upstream regions of the S. cerevisiae HSP78 and HSP104 genes were identified using BLASTN-based dot plot analysis. A comparison of 320 bp regions upstream of the HSP78 and HSP104 open reading frames (Fig. 4A) revealed three serially-aligned blocks of sequence whose length is consistent with their function as possible protein binding sites (Fig. 4B) and that are found in orthologous loci in other Saccharomyces species (Fig. 4C, track 6). These analyses led to the identification of three shared signature features (Fig. 4C): (i) an Hsf1 binding Element characterized by a series of four inverted 5’-nGAAn repeats [32–36] shown in blue (HSE4, track 3), and in red and orange (DP-1 = dot-plot motif 1, tracks 1 and 2); (ii) a triplet array of Stress-Response Element (STREs) that bind the paralogous zinc finger transcription factors, Msn2 and Msn4 [37,38]; and (iii) a core promoter element that resembles a TATA-box with additional sequences...
Fig 3. Phylogenetic analysis confirming absence of LYS4 and LYS20 in animals. (A) Phylogenetic inference using Bayesian MCMC methods shows that the closest sequence matches in animal genome sequences to the LYS4 genes are the ACO1 genes, which encodes the aconitase enzyme of the tricarboxylic acid (TCA) cycle. Choanoflagellates (triangles) have LYS4 and ACO1 but animals (squares) have only ACO1. The tree is rooted between the two gene clades for Lys4 and Aco1. (B) A representative neighbor-joining (NJ) tree shows the presence of LYS4 (blue) and ACO2 (pink) in choanoflagellates (triangles), but not in animals (squares), for which sequences could not be found. One related gene ACO1 (maroon), which encodes an enzymatic activity of the TCA cycle, is still present in animals, while LEU1 (mustard yellow), a second related gene, involved in the leucine biosynthesis as part of the branched chain amino acid pathway, is lost in both choanoflagellates and animals. ACO2 could not be found in the colonial choanoflagellate, Salpingoea rosetta. Numbers indicate bootstrap support from 500 replicates. The tree is rooted between the Leu1 outgroup clade and the Lys4/Aco1/Aco2 superclade. (C) A representative neighbor-joining tree shows the presence of LYS20/21 gene in choanoflagellates (triangles) and other organisms. LYS20 and LYS21 correspond to yeast specific duplications of this genetic function. No related homologs are found in animals (Fig. 1). Distantly-related bacterial outgroups do however possess these genes. The tree is rooted between the eukaryotic Lys20/21 clade and the bacterial (Deinococcus-Thermus) gene clade.

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immediately downstream (DP-III, track 5). Binding of Hsf1 to the HSE and binding of Msn2, and Msn4 to the STRE have been observed under oxidative stress, heat shock, and acid treatment as well as for growth in rich media (YPD) [39].

To determine whether the shared HSP104 and HSP78 motifs are in general use or specific to the clpB genes, we conducted whole-genome searches for each sequence. A search of the S. cerevisiae genome using an IUPAC consensus of the extended dot plot motif DP-I revealed that the HSP78 and HSP104 loci are the only two in S. cerevisiae with the DP-I motif (Table 1, Model 1). To identify highly similar variants, we implemented a position-weighted matrix (PWM) description of DP-I (Table 1, Model 2). This more permissive model identified 12 loci, half of which are related to a specific subset of proteins involved in various aspects of unfolded protein

Fig 4. Yeast HSP78 and HSP104 share a unique regulatory architecture. (A) A dot plot between 320 bp sequence blocks located -10 bp and -65 bp upstream of the S. cerevisiae HSP78 (vertical) and HSP104 (horizontal) open reading frames, respectively. These coordinates were chosen because it anchors their HSE4 elements at the same position (starting 80 bp from the left) for ease of comparison. This comparison reveals a serial ordered alignment corresponding to an HSE4 element (blue box), a short alignment partially overlapping a triple array of STRE elements (pink boxes), and a TATA-containing core promoter element (green box). (B) Graph of the relevant details of the HSP78 (top) and HSP104 (bottom) upstream regulatory sequences. Each track (numbered lines 1–6) provides locations of the dot plot motifs (DP-I and DP-II) and known binding motifs (colored boxes in lines 1–5). Labels above certain binding sites indicate binding of a known transcription factor at this position under the specific conditions listed [39,49]. Line 6 shows conserved non-coding sequences (CNS) as determined from comparisons of corresponding orthologs from other species of Saccharomyces. Dots and lines in the ruler mark off 10 bp and 100 bp intervals, respectively. The transcriptional arrows represent only approximate positions of the transcriptional +1 based on distance from the TATA-containing promoter motif. (C) Motif key for panel B. (D) Results using the Serial Pattern of Expression Levels Locator (SPELL [31]) algorithm/query engine to identify the closest regulated genes to HSP78 (left graph), HSP104 (center graph), or both genes combined (right graph). Plotted are the transcriptional correlation scores (SPELL, Adjusted Correlation Scores). The SPELL scores are based on pair-wise Pearson correlation co-efficients with Fisher z-transforms across 9,190 gene expression studies in yeast [31]. These graphs show that the two clpB genes are co-regulated with one another (red arrows) and to Hsp70/DnaK (K) and J-domain/DnaJ co-chaperones (J), which bind and recruit unfolded proteins to chaperone machinery in the cytoplasm (blue bars) and in the mitochondria (purple bars).

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Table 1. A perfect HSE4 element is specific to a clpB refolding regulon in S. cerevisiae.

<table>
<thead>
<tr>
<th>#</th>
<th>Model*</th>
<th>Model description*</th>
<th>Genomic distribution</th>
<th>Precision **</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>2-way DP-I</td>
<td>5’-TTTCCAGA\text{A}n\text{t}TTCTAGAAG</td>
<td>ClpB: HSP104, HSP78.</td>
<td>100% (2/2)</td>
</tr>
<tr>
<td>3</td>
<td>3-way PWM (Ψ\text{-}count: 0.2)</td>
<td>T:4660000000000266000000000, C:20006600020006000000, A:00000060660200060662.</td>
<td>ClpB: HSP104, HSP78. Hsp70: SSA1. DnaJ: SIST, APJ1. Other: BTN2, UBI4, ADD37.</td>
<td>78% (7/9)</td>
</tr>
<tr>
<td>4</td>
<td>HSE4</td>
<td>5’-TTCTnGAA\text{nn}TTCTnGAA</td>
<td>ClpB: HSP104, HSP78. Hsp70: SSA1. DnaJ: SIST. Other: BTN2, UBI4, HSP42.</td>
<td>78% (7/9)</td>
</tr>
<tr>
<td>5</td>
<td>Perfect HSE4</td>
<td>5’-TTCYRG\text{A}annTTCYRG\text{A}</td>
<td>ClpB: HSP104, HSP78. Hsp70: SSA1. DnaJ: SIST. Other: BTN2, UBI4.</td>
<td>100% (6/6)</td>
</tr>
<tr>
<td>6</td>
<td>Skipped HSE4 + 3x STREs</td>
<td>5’-TTCTnGAA\text{nnn}nnnGAA + 3x 5’-CCCCT</td>
<td>ClpB: HSP104, HSP78. Other: HSP42.</td>
<td>100% (3/3)</td>
</tr>
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Abbreviations: DP-I, dot plot motif I; IUPAC, International Union of Pure and Applied Chemistry; PWM, Position-Weighted Matrix; Ψ, pseudo-count correction.

* Models 1–3 describe the genomic distributions of the clpB-specific HSE4 element using different descriptions. IUPAC DNA codes are used as needed except ‘N’ is represented by a dot (‘.’), Y = pYrimidine (C or T), and R = puRine (A or G).

** Percentages refer to the fraction of all genomic loci matching the model signature that are known to be involved in prion homeostasis. All such loci are listed in bold in the penultimate column. Hits in CDS regions are not listed but are counted for the precision metric.

binding, disaggregation, and refolding. For example, one of the 12 loci corresponds to the promoter for SSA1, which encodes a member of the Hsp70/DnaK family of chaperones [40], which bind unfolded proteins to promote proper re-folding and prevent aggregation. In yeast, Ssa1 has known roles in disassembling misfolded protein aggregates together with Hsp104 [41], and in localizing mRNAs to the outer mitochondrial membrane for co-translational translocation and import into the mitochondria [42]. Hsp70-family proteins also interact with Hsp78. In conjunction with the mtHsp70 chaperone Ssc1, Hsp78 resolubilizes proteins from insoluble aggregates within mitochondria. Ssc1 is also involved in mitochondrial inner membrane import and refolding.

To improve our first PWM model, we constructed a second PWM descriptor based on the HSE4s upstream of the two clpB orthologs as well as SSA1 (Table 1, Model 3). With this modified PWM-based model, fewer matches in ORFs, which are likely spurious, and an additional match in the promoter for the ubiquitin-encoding locus, UBI4, were identified. Ubiquitin is conjugated to misfolded proteins, marking them for proteolysis rather than refolding. Thus, this PWM descriptor improves the precision of the model with 7 loci of 9 involved in the disaggregation of misfolded proteins, their refolding, or their proteolysis. Sequence alignments of these targets indicate that the clpB-type HSE4 defines a set of four inverted repeats with the sequence 5’-nGAAn, which can also be described as two pairs of the palindrome 5’-TTCYrGAA separated by 2 bp (IUPAC DNA code: y = C or T, and r = A or G, n = any letter).

To test the significance of the central “YR” in the HSE4 motif, we queried the genome with an HSE4 motif in which these residues are not specified (Table 1, Model 4). We identified only 12 loci, including one new proteostasis gene, HSP42, which encodes a small heat shock protein that plays an important role in aggregate sorting and mediating the retention of misfolded proteins including amyloids in a peripheral cellular compartment [43,44]. The idealized (perfectly palindromic) HSE4 element, 5’-TTCyRGAA\text{nn}TTCyRGAA, is found at only six loci (SSA1, HSP78, BTN2, UBI4, HSP104, SIS1) in the entire genome, all of which are located at promoters of genes involved in protein homeostasis, including both clpB orthologs (Table 1, Model 5). SIS1, like APJ1 (Table 1, Model 3) encodes a J-domain (DnaJ) family member that functions as...
co-chaperones to specific Hsp70 (DnaK) chaperones [45,46]. The BTN2 gene encodes a sorting factor that accumulates in stressed cells to promote the transit of misfolded proteins to specific protein deposition sites, which are determined by alternative complex formation between Btn2 and Hsp42 (Table 1, Model 4) or Sis1 [44,47,48]. The results of these analyses are consistent with the hypothesis that yeast HSP78 and HSP104 are co-expressed together with a small set of other protein folding genes. This suggests that the two ClpB chaperones may participate in a single process related to protein homeostasis.

To investigate whether yeast HSP78 and HSP104 are indeed co-regulated, we used the SPELL (Serial Pattern of Expression Levels Locator) algorithm and query engine to survey their transcriptional correlation across 9,190 gene expression microarray experiments [31]. We found that HSP78 was most significantly co-expressed with HSP104 and HSP42 (red arrow in Fig. 4D left panel). Similarly, HSP104 was most significantly co-expressed with HSP42 and HSP78 (red arrow in Fig. 4D center panel). Furthermore, HSP78 alone, HSP104 alone, and HSP78 plus HSP104 in combination are co-regulated with diverse J-domain proteins and Hsp70/DnaK co-chaperones, which are known to stabilize unfolded protein states, respectively (J and K in Fig. 4D). This Hsp70 chaperone machinery is also co-expressed with other HSE4-associated genes such as HSP42, BTN2, UBI4, supporting the functional reliability of the HSE4 signature (Fig. 4D).

HSE4 regulation of the clpB genes is conserved in choanoflagellates

To determine whether HSE4 regulation is also present in the choanoflagellates we looked for HSE4 signatures in the M. brevicollis and S. rosetta genomes, which maintain the clpB genes. Our analysis of the S. rosetta genes revealed the presence of a consensus HSE4 sequence upstream of the HSP78 gene and a near consensus HSE4 sequence upstream of HSP104. We identified a total of 5 additional HSE4 sequences in genome-wide searches for a consensus HSE4 (Table 2). One HSE4 is located upstream of UBA1. Three additional HSE4s are found upstream of unrelated genes, and one lies within an open reading frame. UBA1 encodes a ubiquitin-like modifier activating enzyme, which catalyzes the first step in ubiquitin conjugation, marking proteins for degradation. HSE4 regulation of UBA1 in the choanoflagellate is reminiscent of HSE4 regulation of ubiquitin (UBI4) in S. cerevisiae.

We also find more extended Hsf1 binding arrays, here “HSE4+” elements, at the clpB loci of the choanoflagellate M. brevicollis and chytrid fungus B. dendrobatidis. These match so-called extended “skipped HSE” motifs [32] in which the pattern consists of five or six inverted 5′-nGAAn repeats (HSE5, HSE6) with one or more missing internal triplet TTC’s or GAA’s. For these genomes, we can derive lineage-specific consensus signatures that are specific to a small clpB regulon. For example, in M. brevicollis we find a consensus skipped HSE at exactly 6 loci that include HSP104, HSP78, and an Hsp70 gene, while in the chytrid B. dendrobatidis we find only five loci, including HSP78 and HSP104, with either an HSE4 (5′-TTC..GAA..TTC..GAA) or a skipped HSE5 (5′-TTC..GAA..TTC..G..TTC). Last, we note that even in the yeast S. cerevisiae there is a single locus matching a skipped HSE5 motif, although in this case it is a perfect HSE5 (i.e., not skipped) and is associated with the SIS1 DnaJ-encoding locus that we already identified as having an HSE4 motif. Overall, these results show that the archetypical HSE4/HSE4+ elements are ancestral to a small regulon that includes HSP78 and HSP104 and that these genes evolve with only modest changes to the Hsf1 binding site motif.

ClpB promoter architecture is unique

To identify the extent to which the clpB promoter architecture is present in the S. cerevisiae genome, we searched for the combination of the HSE4 motif and the STRE cluster. We relaxed the stringency of the combined motifs until we found the closest matching sequences. For example,
a search for all 160 bp windows containing at least three STREs (5'-'CCCCT) and a match to either a skipped (5'-'TTCnnGAAnnnnnnnGAA) (Table 1, Model 6) or an end-clipped HSE4 (5'-'CnnGAAnnTTCnnG) identifies the same three loci: HSP42, HSP78, and HSP104. The spacing of the triplet cluster at HSP42 is not similar to that found at clpB promoters and the STRE elements at HSP42 are not conserved across the Saccharomyces genus [49], nor is the HSP42 family found in choanoflagellates. Thus, yeast HSP78 and HSP104 are uniquely associated with a shared promoter architecture, one element of which (HSE4s and skipped HSE5/6 arrays) is still uniquely associated with their orthologous promoters in choanoflagellate genomes.

Absence of Hsp78 exacerbates hsp104 mutant thermosensitivity

The co-regulation and co-expression of both mitochondrial and cytoplasmic ClpB functions is consistent with a potential joint function. To experimentally assess this hypothesis, we tested in S. cerevisiae whether a phenotypically normal hsp78 deletion had any effect on hsp104 mutant thermosensitivity. In these experiments, single and double mutants were subjected to extreme heat (50°C) in rich media (YPD) following pre-exposure to mild heat sufficient to induce HSP104 and HSP78 gene expression. As expected, survival of the hsp78 deletion mutant was indistinguishable from that of wild type (p = 0.9) and viability of the hsp104 mutant was reduced relative to that of wild type strains (Fig. 5A). This is consistent with previous reports showing that hsp104 deletion strains are measurably thermosensitive [50–52] while hsp78 deletion strains have more subtle effects when heat stressed [24–26,53–55]. However, we found that the hsp78 hsp104 double mutant was significantly more sensitive than hsp104 single mutant (~10-fold difference, p = 0.0005) suggesting that Hsp78 contributes to induced thermotolerance in the absence of Hsp104, although it has no appreciable effect in its presence (Fig. 5A). This revealed for the first time that survival in extreme heat depends on the presence of both compartmentalized ClpB functions. Together with the observation of co-loss and co-regulation of clpB orthologs, this result leads us to postulate a shared ClpB function.

Yeast interactome analysis suggests that the ClpB proteins share specific protein clients

Connections between the lost genetic functions can be inferred from gene association algorithms (e.g., GeneMania [16,17]) that identify physical or genetic interactions among the lost genes. We identified several links between genes in the different biosynthetic pathways (blue nodes in Fig. 6A) and other lost genes that may be informative upon further study. Here, we focus on several physical links connecting both of the ClpB chaperones (red nodes in Fig. 6A) with Lys4 and Lys20. Lys20 and Lys4 occupy early steps in the α-aminoadipate lysine biosynthesis pathway, which is found in a small group of thermophilic archaea and bacteria and in non-metazoan eukaryotes [56–58]. Lys20 is the mitochondrial isoform of homocitrate synthase
that exerts most of the flux control in lysine biosynthesis in yeast [59], while Lys4 encodes mitochondrial homoaconitase, a 4Fe-4S iron-sulfur protein [60].

The observation that Hsp78 and Hsp104 each make specific contacts with Lys4 and Lys20 (Fig. 6A) despite their compartmentalized roles, leads us to propose that clpB-encoded chaperones Hsp104 and Hsp78 were lost because of “lost shared clients” that included Lys4 and Lys20, but may also have included other nuclear-encoded, cytoplasmically-translated, and mitochondrially imported proteins. This idea is investigated further below.

Lys4 and Lys20 interact specifically with RAC and ClpB chaperone systems

If Lys4 and Lys20 are heat-labile clients specifically disaggregated by Hsp78 and Hsp104 after heat denaturation, they are likely also to require folding assistance de novo during translation and/or co-translationally with mitochondrial import. We examined whether Lys4 and Lys20 make specific contacts with the ribosome-associated chaperone (RAC), which is important for de novo folding during translation and leads to aggregation when compromised [61–63]. The RAC/Ssb system is also required during co-translational translocation into subcellular compartments [64]. RAC is composed of the Hsp70 family chaperone Ssz1 tightly associated with its J-domain co-chaperone Zuo1 and more loosely with a second Hsp70 family member, Ssb (Ssb1 or Ssb2). To investigate the role of RAC in Lys4 and Lys20 folding, we used experimental interaction data to construct the interactome among the Lys4 and Lys20 proteins, the RAC system and various Hsp70/DnaK and J-domain/DnaJ proteins that we identified in our gene regulatory analyses (Fig. 6B).

These analyses show that Lys4 and Lys20 interact with RAC machinery and the ClpBs but not with other Hsp70 chaperones and DnaJ co-chaperones. For example, Lys4 interacts
physically with both components of RAC (the Hsp70 Ssz1 and its dedicated J-domain co-chaperone Zuo1), and with the ribosomal Hsp70s Ssb1 and Ssb2, which work in conjunction with RAC (Fig. 6B). But Lys4 does not interact with diverse other cytoplasmic (Sse2, Ssa4, Ssa1) or mitochondrial (Ssc1) Hsp70s nor with J-domain family members (Sis1, Ydj1, Apj1, Mdj1), several of which (SSE2, SSA4, SSA1, SIS1, MDJ1) are co-expressed with the clpBs (Fig. 4D), or definitively co-regulated with the clpBs (SSA1, SIS1, APJ1) (Table 1) based on the presence of
HSE4 elements. A similar result was seen for Lys20. We thus propose that unfolded Lys20 and Lys4 are recognized by RAC during translation as nascent unfolded polypeptides and again by ClpBs after post-translational heat denaturation.

Other Fe/S proteins in the Lys4 superfamily are also lost in choanoflagellates and animals

If Lys4 and Lys20 were lost in animals due to their deleterious misfolding properties, then other proteins with similar structures may have had similar fates. Indeed, additional Lys4 superfamily proteins containing 4Fe-4S clusters (Fig. 7A) were also lost in the evolution of animals. We looked for examples in which other Lys4 superfamily members were lost in choanoflagellates and had therefore escaped our initial screen for losses in the stem-metazoan lineage. While Lys4 is absent in animals but present in choanoflagellates (Table 1 and S1 Table), we found that ACO2 is absent in animals and the colonial choanoflagellate S. rosetta, but present in the unicellular choanoflagellate, M. brevicollis, and that LEU1 and ILV3 are absent in both animals and choanoflagellates (Fig. 7B). Like Lys4, both Ilv3 and Aco2 receive their Fe/S clusters in the mitochondria [60] although Leu1 is a cytoplasmic enzyme [58]. Fe/S proteins have two special properties; first, the free iron that is presumably released upon stress-mediated denaturation can generate hydroxyl radicals, which are one type of insidious reactive oxygen species (ROS) [65,66]. Second, Fe/S cluster proteins are highly susceptible to inactivation by superoxide, which accumulates and contributes to the loss of viability in heat stressed cells [67–70]. Thus, Lys4, which is among the most highly conserved proteins in our list of twenty-four lost genetic functions (Fig. 1, S1 Table), may be especially heat labile and prone to ROS generation when misfolded in the presence of oxygen. The same may be true of other Lys4 relatives. We therefore speculate that this class of protein may require dedicated folding machinery.

Hydrophobic amino acids, which are normally packed into the interior of a peptide, are recognized by yeast Hsp104 when exposed in unfolded aggregates and when added to model clients increase their recognition after denaturation [71]. We therefore compared the hydrophobicity of the Fe/S coordination pocket of Lys4 with that of Aco1, which is maintained in animals. We first aligned the yeast and choanoflagellate peptide sequences from these proteins, including those from Aco2 and annotated the family-specific residues for Lys4 (cyan shading in Fig. 7C) and Aco1 (gray shading). We found that 41.4% (12/29) of the conserved Lys4-specific residues are hydrophobic compared with 22.8% of the conserved Aco1-specific residues (Fig. 7D). As the hydrophobic content of a protein is likely to be positively correlated with aggregation after heat denaturation, the two-fold enrichment in Lys4 relative to Aco1 is likely to pose a significant cellular challenge.

Discussion

Hsp78 and Hsp104 chaperones function coordinately and may have common clients in Lys4 and Lys20

Here we show for the first time that Hsp78 and Hsp104 (i) were both definitively lost in an ancestor of animals despite their presence in all non-animals; (ii) were uniquely co-regulated by a multimeric and ordered array of perfect binding sites for Heat Shock Factor 1 (Hsf1) in an ancestor of animals based on the shared presence and restricted use of this regulatory signature in fungal and choanoflagellate genomes; (iii) function together in thermotolerance (Fig. 5A); and (iv) make specific contacts with Lys4 and Lys20, two nuclear encoded, mitochondrial enzymes (Fig. 6A and 6B).
These results are consistent with the idea that specific mitochondrial biosynthetic enzymes are shared clients of both ClpBs and could be pertinent to their joint loss at the base of animals. We propose that cytoplasmic Hsp104 disaggregates and refolds nuclear-encoded, cytoplasmically translated mitochondrial enzymes such as Lys4 and Lys20 that are also refolded by Hsp78 in the mitochondria. However, how compartmentalized Hsp104 and Hsp78 chaperones could share mitochondrial clients is unclear. One idea is that there are parallel co-translational import and mitochondrial targeting pathways for getting Lys4 into the mitochondrial matrix. In this case, Lys4 apoprotein en route to the mitochondria might exist as a cytoplasmic pool that is prone to misfolding and aggregation. A second idea is that thermally denatured, incompletely...
translocated Lys4 enzymes might escape into the cytoplasm, which would create a need for maintaining a cytoplasmic ClpB activity. Finally, a third possibility is that Lys4 is able to escape via compromised mitochondrial membranes, which are thermosensitive. For example, even mild heat (37°C), sufficient to induce the ClpB chaperones, results in a 5-fold increase in mitochondrial leakage based on the transfer of mitochondrial DNA to the nucleus [72].

Despite our findings of a possible Lys4-ClpB client-chaperone relationship, it is not likely that Lys4 was the only client of the ClpBs in the stem-metazoan lineage. First, yeast data has implicated several clients of Hsp104 [73]. Second, ClpBs are conserved in plants and bacteria, but Lys4 is not, because plants and most bacteria utilize a lysine biosynthesis pathway different from the one in opisthokonts [56]. For these reasons, we suspect that the ClpBs in animals were lost not so much because a single dedicated client was lost, but because a threshold number of clients were lost.

A cascade of gene loss in the stem-metazoan lineage

Our results suggest that many metabolic genes were lost in the stem-metazoan lineage (Fig. 1). This may reflect wholesale reduction of biosynthetic capacity in favor of obligate high-throughput filter feeding and could be due to the trophic benefits of multicellular organization, allowing biosynthetic precursors to be derived as nutrients from dietary sources. For example, sponges, which have diverse complex multicellular chambers for pumping and filtering water, are known to take up and remove up to 99% of bacteria and 94% of small eukaryotes from the water that they filter [74,75]. Thus, we propose that the stem-metazoan innovation resulting in efficient dietary intake of picoplankton via grazing and/or filtering is likely to have resulted in loss of selection of certain biosynthetic pathways because it became more energetically efficient to obtain them from feeding rather than from biosynthesis. We suggest that this working hypothesis may lead to deeper insights into the evolutionary emergence of specialized-feeding surfaces and the forerunners of the “gastraea” [76].

Based on our finding of specific gene losses, we propose a two-step model in which loss of biosynthetic enzymes (step 1) is followed by secondary loss of attendant chaperones (step 2). This model is supported by several additional observations in choanoflagellates. First, we note that the TRP2 gene, which encodes the first enzyme in the energetically-costly tryptophan biosynthetic pathway is not as well conserved in choanoflagellates as the intermediate enzyme encoded by TRP3, and is more poorly conserved than TRP5, which encodes the final enzyme in tryptophan synthesis (Fig. 1 and S1 Table). Thus the extent of conservation is co-linear with position along the biosynthetic pathway, consistent with the possibility that choanocyte-type filter feeding [77] and possibly facultative coloniality in this lineage led to reduced selection of the early components of this pathway while the final biosynthetic enzymes were maintained for the ability to complete tryptophan biosynthesis using intermediate precursors provided by ingested microorganisms. Second, in the course of testing various computational approaches to gene loss, we discovered that some “lost” animal genes are present in the unicellular choanoflagellate *M. brevicollis* but not in the facultatively colonial [12] choanoflagellate *S. rosetta* or in animals (e.g., *ACO2* in Fig. 3B). Third, we did not find any of the genes encoding enzymes in the biosynthetic pathways for the branched chain amino acids, (leucine, isoleucine, and valine), in either animals or choanoflagellates. These amino acids are three of the ten essential amino acids animals must retrieve from their dietary intake. The pattern of metabolic gene loss in choanoflagellates suggests that (i) specific metabolic gene losses had already occurred in the stem-holozoan ancestor or were underway with holozoan divergence; (ii) additional metabolic gene losses were sustained during the divergence of choanoflagellates and animals (e.g., *ACO2*); (iii) gene losses in choanoflagellates began with genes encoding enzymes at the beginning of pathways (e.g., Trp enzymes); and (iv) metabolic genes were more completely lost with the
evolution of animals. These results are likely to prove fundamental for understanding formative events in the evolution of the metabolic physiology of the animal kingdom.

**Post-genomic forward genetics**

We find that phylogenomic-contextualized network analyses essentially function as a powerful post-genomic forward-genetics. While we have used this integrative systems approach to identify lost gene pathways, it could also be used to identify interacting pathways that originated together. These methods can lead to new hypotheses, informed by medium to high throughput experimental data sets. Thus, this post-genomic approach may be more economical and in principle more efficient because multiple families of hypotheses can be constructed and rejected *in silico* prior to selecting and experimentally testing the best, most specific, most well-informed, and experimentally informative hypotheses.

**Materials and Methods**

**Comparative genomics**

Ensembl’s BioMart tool [21] was used iteratively to query, sort, and identify orthology calls [78,79] for Ensembl Genes Release 75 (www.ensembl.org) and Ensembl Fungi Genes Release 22 (fungi.ensembl.org). The choanoflagellate genome of *Monosiga brevicollis* (v1.0) was queried from the Joint Genome Institute databases (genome.jgi.doe.gov), the *Salpingoeca rosetta* genome from Broad Institute databases (www.broadinstitute.org). Note that BioMart orthology calls are specific to a release set of genome comparisons and that orthology calls sometimes change with each Release. All genomes were also queried using taxonomic specification at NCBI’s BLASTP server, except for the ctenophore *Mnemiopsis leidyi*, which is available via National Human Genome Research Institute (NHGRI) server (research.nhgri.nih.gov/mnemiopsis). Homologs for fungal lineages outside of Ascomycetes and other non-fungal lineages were selected following BLASTP analysis of specific genomes using *S. cerevisiae* proteins as queries. Additional filtering using E-value thresholds is described in the text. Data from a Boolean-type PFAM taxonomy search (*Dictyostelium* protein domains absent in yeast and animals) was conducted at http://pfam.xfam.org. The identified PFAM domain IDs were then used to retrieve the protein sequences using BioMart [15,21]. These sequences were used in a BLASTP query of choanoflagellates. Orthologs of lost genes from *S. cerevisiae* or *D. discoideum* are listed in S1 File.

**Phylogenetic analyses**

Alignments were produced using ClustalW. Bayesian Markov Chain Monte Carlo analyses were conducted using MrBayes [80–82] and initial runs sampled diverse fixed amino acid rate matrices, but most data sets overwhelmingly supported the WAG [83] (ClpB trees) or Jones/JTT [84] (Lys4 trees) models after the 25% burn-in phase. Topological convergence was defined as having achieved an average standard deviation of split frequencies < 0.01, but we typically achieved < 0.002. Maximum Likelihood and exploratory Neighbor-Joining trees were conducted in MEGA5 and MEGA6 [85,86] typically using Jones/JTT matrices and a gamma distributions with shape parameters spanning 0.7 to 1.0 with replicate boot strap support based on 500 replicates. Alignment files and supporting Bayesian analysis files are archived in S2 File.

**Gene Association Analysis**

Gene association data pertaining to individual chaperones was retrieved from ChaperoneDB (chaperonedb.ccbr.utoronto.ca) and uploaded to GeneMania [16,17] (www.genemania.org). Evidence for physical interactions is represented by red edges, evidence for genetic interactions
is shown by green edges and predictions based on selected data imported from the global TAP tag MS dataset [73] is shown by gray edges.

**Regulatory Bioinformatics**

Identification of matching cis-regulatory patterns in whole-genome sequences were conducted by downloading text files of each respective genome, removing newlines, and searching for blocks of sequence containing the indicated patterns (Tables 1 and 2). These searches were conducted using regular expression patterning matching in simple UNIX shell scripts incorporating perl, grep, and standard shell commands. BLASTN-based dot plot analysis was conducted using a window size of 320 bp for sequences upstream of *S. cerevisiae* HSP78 (vertical) and HSP104 (horizontal) open reading frames. Known transcription factor binding sites at listed positions were retrieved using the UCSC genome browser [39,49]. Conserved non-coding sequences (CNSs) were determined from comparisons of corresponding orthologs. Transcriptional correlation scores correspond to the Adjusted Correlation Scores for 430 datasets representing 9190 total arrays from 286 published studies queried via SPELL [31] (Serial Pattern of Expression Levels Locator) and available through the *Saccharomyces* Genome Database (spell.yeastgenome.org).

**Yeast Strains and Survival Assay**

Yeast strains were constructed by mating and tetrade dissection of diploids starting with strains from the MATα haploid deletion collection (BY4742). Deletion collection strains were mated with BY4730 and diploids sporulated and dissected to generate LYS2+ derivatives of the *hsp78* and *hsp104* mutants (*hsp78*: JF2478, JF2479, JF2480 and *hsp104*: JF2473, JF2474 and JF2498). These strains were crossed to generate the double *hsp78 hsp104* mutants (JF2494, JF2495, JF2516). Strain genotypes are provided in Table A in S3 File. *clpB* genotypes were confirmed by colony PCR using a forward primer upstream of the deletion and a reverse primers within the kanMX cassette. Primer sequences are provided in Table B in S3 File.

Survival was assayed by colony forming unit (cfu) assays. Cultures were grown to saturation in YPD at 30°C, then subcultured and grown to log phase under the same conditions. Log phase cultures were shifted to 37°C for between 40–60 minutes and then diluted (1:1) into ice-cold water or diluted similarly into 50°C YPD media for 20 minutes. Cultures were then serially diluted and plated on YPD. Colony number was used to calculate the concentration of survivors for each strain and each treatment condition. Values for individual strains were averaged. SD, standard deviation.

**Supporting Information**

S1 Fig. Neighbor-Joining phylogenetic analysis of the closest related animal sequences to *clpB*. (A) Phylogenetic inference using Bayesian MCMC method, similar to Fig. 2A except a possible ortholog from the cnidarian *N. vectensis* is also included. This analysis shows that this sequence is correctly identified as a partial fragment of a *clpC* sequence, i.e., not a *clpB* gene. Topological convergence was achieved after 300,000 generations with a relative burn-in phase of 25% (MrBayes) [80–82]. Mixed models were tested, but WAG was the sole model favored upon completion. Average standard deviation of split frequencies was < 0.002. The tree was rooted between the entire *clpB* clade and the single *clpC* fragment. (B) The *M. brevicollis* Hsp104 sequence was used as a BLASTP query against the NCBI RefSeq database taxonomically restricted to animals. The highest scoring eight database hits were incorporated into the *clpB* phylogeny. High scoring bacterial sequences were also added to the phylogeny based on BLASTP analysis of each animal query against the RefSeq database taxonomically restricted to
bacteria. Five of the top animal hits are shown in this tree. All positions with less than 50% site coverage were eliminated. There were a total of 862 positions in the final dataset. Bootstrap values lower than 50 are not shown. Animal sequences included in the tree are XP_002944514.1, *Xenopus tropicalis*; XP_003391862.1, *Amphimedon queenslandica*; XP_003493100.1, *Bombus impatiens*; XP_002199578.2, *Taeniopygla guttata*; gb EDO25580.1, *Nematostella vectensis*; and gb AAA68910.1, *Bos taurus*.

**S2 Fig.** Phylogenetic analysis of the ANKCLP gene of animals and choanoflagellates. (A) Comparison of the ankyrin (ANK)–partial bacterial ClpB moiety to the actual ClpB domain structure, which contains two AAA+ domains. (B) Phylogenetic analysis by Bayesian MCMC (2,000,000 generations with average standard deviation of split frequencies < 0.0034 and with mixed amino acid models tested, 25% burn-in, and with WAG model having all posterior probability) showing that ANKCLP is found in holozoans and does not correspond to either bacterial clpB (*E. coli*) nor its eukaryotic orthologs HSP78 and HSP104. In the absence of HSP78 and HSP104 in animals, ANKCLP has been incorrectly annotated as "Casein lytic protease B, clpB". Posterior probabilities shown at nodes. (C) Neighbor-Joining analysis of the same sequence alignment used in (B) with support from 500 bootstrap replicates.

**S1 File.** FASTA file of list of 28 genes from **Fig. 1.**

**S2 File.** Zipped archive with all NEXUS alignment files and associated Bayesian analyses.

**S3 File.** Tables A and B. Table A, Yeast strains used in this study; Table B, Primer sequences used in this study.

**S1 Table.** List of twenty-four lost genetic functions.

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**Author Contributions**

Conceived and designed the experiments: AJE JSF. Performed the experiments: AJE JSF. Analyzed the data: AJE JSF. Contributed reagents/materials/analysis tools: AJE JSF. Wrote the paper: AJE JSF.

**References**


