

Evolvability of Chaperonin Substrate Proteins

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March 23rd, 2009

Abstract

Molecular chaperones ensure that their substrate proteins reach the functional native state, and prevent their aggregation. Recently, an additional function was proposed for molecular chaperones: they serve as buffers (capacitors) for evolution by permitting their substrate proteins to mutate and at the same time still allowing them to fold productively.

Using pairwise alignments of *E. coli* genes with genes from other gamma-proteobacteria, we showed that the described buffering effect cannot be observed among substrate proteins of GroEL, an essential chaperone in *E. coli*. Instead, we find that GroEL substrate proteins evolve less than other soluble *E. coli* proteins. We analyzed several specific structural and biophysical properties of proteins to assess their influence on protein evolution and to find out why specifically GroEL substrates do not show the expected higher divergence from their orthologs.

Our results culminate in four main findings: 1. We find little evidence that GroEL in *E. coli* acts as a capacitor for evolution *in vivo*. 2. GroEL substrates evolved less than other *E. coli* proteins. 3. Predominantly structural features appear to be a strong determinant of evolutionary rate. 4. Besides size, hydrophobicity is a criterion for exclusion for a protein as a chaperonin substrate.

Introduction

Molecular chaperones are proteins which assist newly synthesized polypeptide chains to fold and mature to functional proteins. Additionally, under cellular stress conditions such as heat shock they are markedly over-expressed, to help prevent the aggregation of unfolded proteins. More recently, it has been proposed that chaperones carry out yet another function: they possess a buffer capacity against detrimental mutations, thereby functioning as a capacitor for evolution.

It was shown for Hsp90 both in *Drosophila* (1) and *Arabidopsis thaliana* (2) that impairing Hsp90 levels either genetically or pharmacologically leads to the appearance of an array of phenotypes. This is attributed to the fact that detrimental genetic polymorphisms, cryptic under conditions with regular chaperone levels, are phenotypically expressed once Hsp90 function is affected. The observed effect can be explained assuming that chaperones have a certain buffering capacitance (thus allowing substrate proteins to accumulate mutations and still reach the native state) whereas chaperone-independent proteins are more likely to misfold when acquiring mutations.

So far, the evidence supporting the hypothesis that chaperones function as a buffer for evolution mainly came from phenotypic observations (1, 2). Since evolution is based on genetic variance, and high phenotypic variability can be based on pleiotropic effects of a few mutations, we decided to investigate the buffering effect of chaperones at the genetic level. However, although satisfying genome data is published for *Drosophila* (3) with 12 fully sequenced species, no satisfying, unbiased list of Hsp90 substrate proteins is available to allow the investigation of

the role played by chaperones for protein evolution in this model organism.

In fact, the concept of chaperones as buffers for evolution is not limited to Hsp90 alone. It was recently expanded to other classes of chaperones (4-6). In particular several biological and functional features of the bacterial GroEL/GroES chaperonin system closely resemble those of the eukaryotic Hsp90:

1. GroEL, just like Hsp90 has a discrete set of substrate proteins, making them both specialist rather than generalist chaperones (7).
2. Both chaperones are essential for survival in their respective environment (8).
3. Hsp90 and GroEL/ES are abundant cellular proteins. Their levels can be decreased significantly by depletion or pharmacological impairment without affecting viability under permissive conditions (2, 9).
4. Both GroEL and Hsp90 bind metastable folding intermediates rather than nascent polypeptide chains (10, 11).
5. As observed for Hsp90 in *Drosophila*, overexpression of GroEL/ES in *E. coli* buffers against a fitness loss caused by deleterious mutations (6).

We therefore decided to base our study on GroEL and its well described substrate proteins in *Escherichia coli* (7), measuring evolutionary distances between chaperone substrate proteins and their orthologs in related bacteria and comparing them to evolutionary distances determined for proteins folding independently of GroEL (Table 1).

If GroEL worked as an evolutionary capacitor, we would expect GroEL substrates from *E. coli* and their orthologous partners in related organisms to show a greater sequence divergence than orthologous pairs of proteins folding inde-

pendently of GroEL. Instead we find lower sequence divergence between GroEL substrate proteins and their orthologs. Our data allows to attribute this finding predominantly to the particular structural composition of the GroEL substrate-set. In general, we do not dismiss the intriguing hypothesis that chaperones function as evolutionary capacitors. However, our data shows, that *in vivo* GroEL substrate proteins in *E. coli* evolved less than proteins folding independently of GroEL. Additionally, our analysis establishes hydrophobicity as a criterion for exclusion for a protein to be a chaperonin substrate.

Results

Evolvability

We based our analysis on a modified list of GroEL substrate proteins published by Kerner et al. (7). The authors isolated stabilized GroEL-GroES-substrate complexes and identified GroEL interacting proteins by mass spectrometry. A quantitative MS approach allowed the identified GroEL substrates to be sorted according to their abundance in complex with the chaperone, relative to their native levels in an *E. coli* cell lysate. The presented data is based on 204 GroEL substrate proteins (Table S2). The Selection criteria for this set of proteins are explained in the Methods section.

As a measure for evolutionary distance, we compared pairwise divergence between genes coding for GroEL substrate proteins of *Escherichia coli* and their orthologs in eight other gamma proteobacteria (*Buchnera aphidicola*, *Haemophilus influenzae*, *Pasteurella multocida*, *Photorhabdus luminescens*, *Salmonella typhimurium*, *Shigella flexneri*, *Vibrio cholerae*, *Yersinia pestis*) (Figure 1).

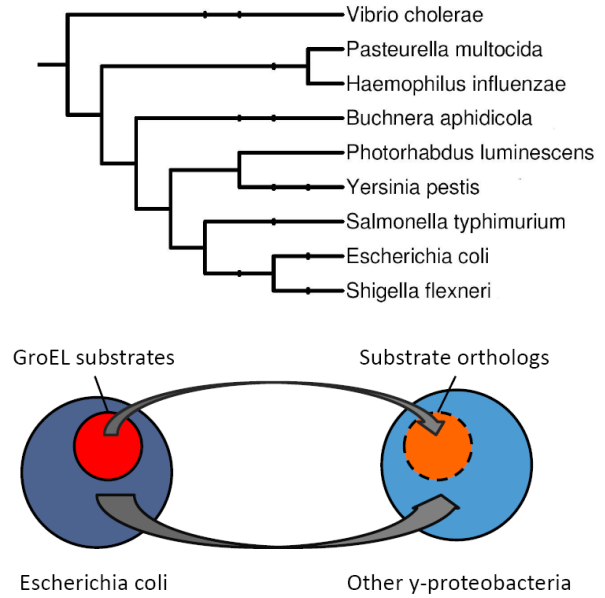


Figure 1. Evolutionary divergence between genes coding for GroEL substrate proteins and all mapped gene pairs in various gamma-proteobacteria. **A:** Organisms for which evolutionary distances to *E. coli* were calculated for this study. **B:** Cartoon depicting the approach for the calculation of evolutionary distances. Red/orange: GroEL substrate proteins and their orthologs. Blue/light blue: All orthologous protein pairs.

To assess the evolutionary buffering capacity of GroEL, the calculated distances between pairs of orthologous genes coding for substrate proteins of the chaperone were compared to evolutionary distances for all mapped gene pairs for the respective organism (Figure 1). Evolutionary proximity of the analyzed organisms ensures that gene pairs selected on the basis of high sequence similarity correspond to orthologous proteins. Only gene pairs with at least 40% sequence identity were considered (Table 1).

Where available, the assigned gene pairs were verified by confirming that they share the same classification in terms of KEGG orthology classes (12).

Organism	genome size	matched gene pairs	substrates compared	Evolutionary distance to E. coli	Ev. distance substrates to E. coli	Z-score	groEL similarity
Buchnera aphidicola	574	495	46	0.091	0.081	-1.7	94.50%
Haemophilus influenzae	1657	1069	91	0.12	0.094	-3.4	94.20%
Pasteurella multocida	2015	1255	107	0.13	0.099	-4.2	63.10%
Photobacterium luminescens	4683	1973	144	0.12	0.088	-5.4	95.40%
Salmonella typhimurium	4527	3187	184	0.074	0.05	-4.2	99.60%
Shigella flexneri	4445	3216	175	0.14	0.089	-2.9	75.20%
Vibrio cholerae	3835	1495	143	0.17	0.15	-4.1	92.50%
Yersinia pestis	4066	2310	159	0.12	0.093	-4.4	96.40%
Escherichia coli	4132	4132	204	0	0	0	100.00%

Table 1. Analysed gamma-proteobacteria and organism specific parameters. Note that Z-scores for all compared organism pairs are negative, indicating a lower evolutionary distance between GroEL proteins and their orthologs in the respective organism, as compared to all mapped protein pairs.

In accordance with the theory that chaperones serve as evolutionary buffers, chaperone dependence was expected to lead to a greater sequence divergence between orthologous genes coding for proteins stringently depending on GroEL for folding, as compared to orthologous gene pairs not coding for GroEL substrate proteins. Instead, we do not find evidence for higher evolutionary dynamics of GroEL substrates, but an opposite effect (Figure 2). In all eight analyzed pairs of organisms, genes coding for GroEL substrate proteins diverge less than their respective control sets of all mapped gene pairs for the tested organisms. (Table 1, Figure S1).

We reasoned that GroEL substrate proteins must possess specific properties, reversing the attributed buffering effect of the chaperone and hence accounting for the smaller genetic divergence. We analyzed parameters influencing the folding pathway (such as hydrophobicity) and structural properties of both GroEL substrates and proteins folding independently of GroEL. We also compared other possible determinants of evolutionary rate, namely expression level and essentiality (Table 2).

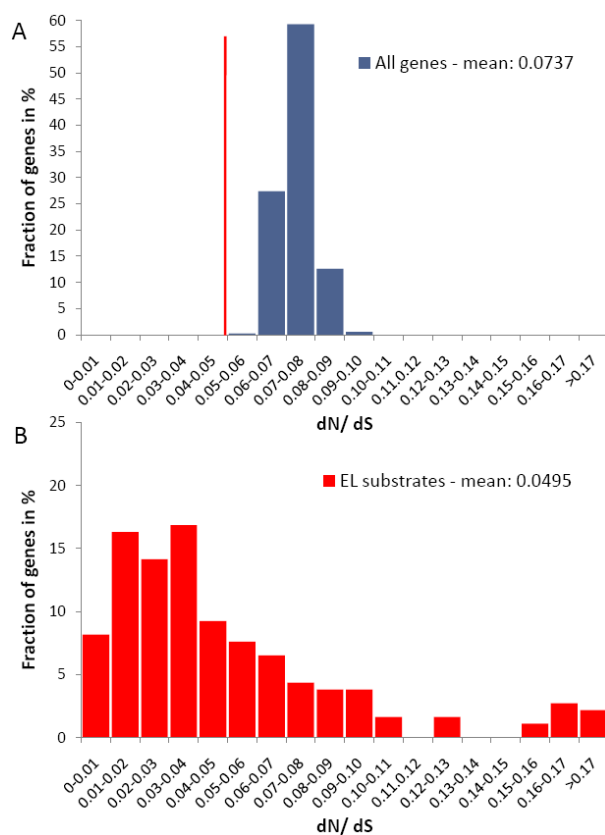


Figure 2. Genes coding for GroEL substrates evolve less than other *E. coli* genes. **A:** Distribution of evolutionary distances of 5000 random sub-sets of gene pairs between *E. coli* and *S. typhimurium*, each comprising 204 members. Red line: average evolutionary distance for GroEL substrate proteins. **B:** Distribution of evolutionary distances of the 204 GroEL substrate genes between *E. coli* and *S. typhimurium*.

We limited the analysis of these parameters to a comparison of *E. coli* with *S. typhimurium*. Several reasons suggested this pair as prime example of the analyzed gamma-proteobacteria:

1. Both organisms have comparable genome sizes (*E. coli*: 4132 genes, *S. typhimurium*: 4527 genes, Table 1)
2. Setting the sequence identity threshold to 90% still allowed us to confidently map 3316 gene pairs.
3. The respective groEL genes are 100% identical in their sequence (Table 1), suggesting that - although not a pre-requisite for this study - orthologous proteins to *E. coli* GroEL substrates also interact with the respective chaperonin of *S. typhimurium*.

The measured evolutionary distance (dN/dS) for the 204 mapped pairs of GroEL substrates between *E. coli* and *S. typhimurium* was 0.050 (Figure 2, Table 1). The average evolutionary distances of all mapped proteins pairs was 0.074 with a calculated Z-score of -4.2 (Figure 2, Table 1, Methods section).

Essentiality

6.4 % of all genes in *E. coli* have so far been characterized as essential, according to the Pec Plus database (<http://www.shigen.nig.ac.jp/ecoli/pecplus/index.jsp>) (13). The average evolutionary distance of mapped gene pairs coding for essential proteins between *E. coli* and *S. typhimurium* is 0.056 with a Z-score of -3.88, compared to 0.074 for all mapped gene pairs. Among the genes coding for GroEL substrate proteins, 14.7%, or 30 out of 204 genes encode essential proteins. The 30 essential proteins among the GroEL substrate proteins (Table S1) have an average evolutionary distance of 0.040 (Z-score: -2.33), as compared to 0.050 for all chaperonin substrates.

The finding that essential genes are more conserved than non essential genes is in agreement with published data, mainly on yeast (14-16), but also on higher eukaryotes (17, 18), although contradicting data has also been published (19). Correcting for the enrichment of essential genes among GroEL substrates did not significantly alter the observed bias in evolutionary distance between GroEL substrates and control proteins (0.052 and 0.075 for GroEL substrates and all proteins, respectively).

Expression level

Published data on yeast suggest a negative correlation between expression level and evolvability (19-21). We do not find a strong correlation between increased expression level and low evolvability in *E. coli* (Figure 3). The calculated Pearson coefficient is close to 0 (-0.055).

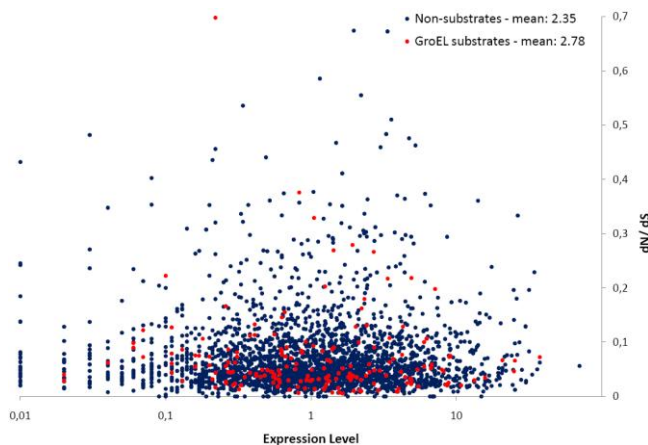


Figure 3. Expression level is not a major determinant for evolvability in *E. coli*. Genes coding for GroEL substrate proteins show slightly higher expression levels than genes coding for proteins not folding with GroEL. Red dots: GroEL substrate proteins. Blue dots: Proteins not interacting with GroEL for productive folding. For visibility reasons, the ordinate was shortened. Two data points for non-EL folders with evolvability values of 0.9365 and 1.303, respectively are missing in the graph.

We used the GEO (Gene Expression Omnibus) database to analyze expression levels of *E. coli* genes (22). We found genes coding for GroEL substrate proteins to be expressed to a higher level (mean: 2.78, standard deviation 9.11, Z-score: 3.3), as compared to the genome wide average (mean: 2.35, standard deviation 4.53) (Figure 3).

The five GroEL substrate proteins with highest expression are PepQ, a proline peptidase; XylA, xylose isomerase; RimJ, ribosomal protein alanine acetyltransferase; GatY and GatZ, two D-tagatose-1,6-bisphosphate aldolase subunits (Table S1).

To account for a potential bias for low abundant proteins to be detected to a lesser extent by mass spectrometry, we repeated the analysis using published data-sets of experimentally detected *E. coli* proteins in proteomic studies (23-25). The mean abundance for this subgroup of 2019 proteins was 2.57 with a standard deviation of 5.14, practically leaving no significant expression level differences between *E. coli* substrate proteins and the control set.

Protein Structure

Published data analyzing the correlation of protein structure and evolvability mainly focuses on the contact density, a measure for the designability of proteins. The contact density considers the fraction of buried amino acid residues in protein structures. It has been shown in some studies that evolvability correlates positively with the global contact density (26, 27). In contrast, other studies suggest that buried residues evolve less than amino acids exposed to the surface of a protein (28-30). We concluded that contact order is not a satisfying criterion to assess the relation of structure and evolution Protein structures can be categorized to higher de-

tail by assigning them to hierarchical SCOP classes (31, 32). For this study, we based the analysis of structural properties of *E. coli* proteins and evolvability on SCOP class assignments.

A detailed structural analysis of GroEL substrate proteins revealed a bias among proteins stringently depending on GroEL towards certain SCOP fold classes (7). We tested the hypothesis that the SCOP class bias observed among GroEL substrates can account for the observation that GroEL substrate proteins evolve less than all other *E. coli* proteins, which fold in a chaperonin-independent manner.

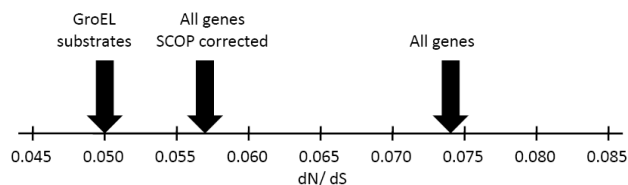


Figure 4. Predominantly structural reasons are responsible for the low evolvability of GroEL substrate proteins. Arrows indicate average evolutionary distances between *E. coli* and *S. typhimurium* for GroEL substrate proteins, all proteins, and all proteins with the same SCOP class distribution as for GroEL substrate proteins.

We repeated the pair-wise alignment of genes between *E. coli* and *S. typhimurium*, reflecting the SCOP class distribution of the GroEL substrates in the control sets. While the evolvability of the random sub-sets shows a mean of 0.074, the SCOP class correction leads to a mean of 0.057 (Z-score: -2.93). This is considerably closer to the calculated mean considering the chaperonin substrate set alone (0.050, Z-score: -4.14, Figure 4). We therefore reason that specific structural properties of the GroEL substrate set are mainly responsible for the observed lower evolvability.

GroEL substrate proteins are highly enriched on the TIM barrel fold (SCOP class c.1) (7). We find that 33 of the 204 GroEL substrate proteins (16.2 %) adopt this fold, as compared to all other protein coding genes where only 2.9% (114 of all 3928 SCOP-annotated proteins) fold to TIM barrels. This structural class is an example of a fold with high sequence divergence, meaning that many different sequences acquire the same fold as native structure (33, 34). It has been speculated that highly designable structures (that is structures which are encoded by many different sequences) evolve rapidly (27), hence TIM barrel proteins in different organisms should show a high evolutionary divergence. What we find is an opposite effect: the average evolutionary distance between TIM barrels of *E. coli* and *S. typhimurium* is 0.061 (Z-score: -2.24), whereas the average distance between all proteins is 0.074. The enrichment of the GroEL substrates in TIM barrels and the low evolutionary divergence between the TIM barrels folding with GroEL (0.054) contributes significantly to the observation that GroEL substrates evolve less when compared to non-substrates.

Hydrophobicity

We assessed the hydrophobicity of proteins using two algorithms: PEPWINDOW, which gives scores based on the Kyte-Doolittle index (35) and TANGO (36), measuring the aggregation propensity of proteins. For stringent GroEL substrates both algorithms independently showed a clear bias towards them being less hydrophobic than chaperonin-independent folders (proteins with predicted trans-membrane domains were excluded from the analysis). We further tested if the hydrophobicity scores of *E. coli* proteins are related to the evolvability of *E. coli* genes. The determined Pearson coefficient of 0.06 shows that there is no linear correlation between the

calculated evolutionary distance for matched gene pairs and the respective hydrophobicity value of the corresponding *E. coli* proteins (Figure 5).

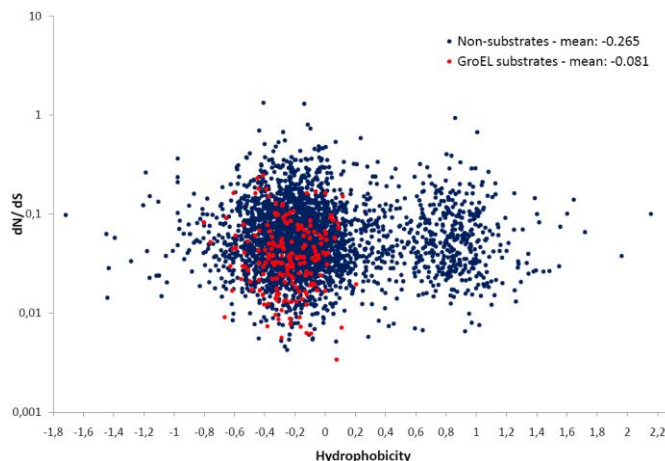


Figure 5. GroEL substrate proteins cluster at low hydrophobicity values and hydrophobicity does not correlate with evolvability in *E. coli*. Red dots: GroEL substrate proteins. Blue dots: Proteins not interacting with GroEL for productive folding.

Our results suggest that GroEL substrate proteins on average are significantly less hydrophobic than other *E. coli* proteins, with hydrophobicity averages of -0.265 and -0.081, respectively and a Z-score of -3.4 (Figure 5). The supports of the distributions of the hydrophobicity values are also markedly different between GroEL substrate proteins (0.18) and proteins folding independent of GroEL (0.47). Figure 5 shows a large group of outliers in the control set with hydrophobicity values of above 0.3 (546 proteins). None of the GroEL substrate proteins reaches this value, with only three proteins in the substrate set having a Kyte-Doolittle score above 0.1 (Figure 5, Table S1). Whereas GroEL substrates cluster in a relatively small window between -0.7 and +0.1, other *E. coli* proteins have Kyte-Doolittle scores between -1.5 and +2.0. To account for a possible mass spectrometry induced bias, we repeated the analysis, using only experimentally identified *E. coli* proteins in

the control group (23-25). Using only experimentally determined proteins showed an even stronger statistical significant difference between the average values of the two data-sets (Z-score: -5.1).

Discussion

We compared GroEL substrate proteins and chaperonin independent folders by evaluating biophysical, structural and physiological parameters with direct influence on evolution. Our analysis results in four findings (Table 2):

1. We find no evidence for GroEL functioning as capacitor for evolution in *E. coli in vivo*.
2. Instead, GroEL substrate proteins show a lower evolutionary distance to orthologous proteins, when compared to other proteins of *E. coli*.
3. Structural properties of GroEL substrate proteins account for their apparent lower evolvability.
4. Proteins folding with GroEL have closely clustered Kyte-Doolittle scores and are on average less hydrophobic than other cytoplasmic proteins of *E. coli*.

Parameter	Evolvability	Substrate specific
Essentiality	✓	○
Expression level	✗	○
Structural properties	✓	✓
Hydrophobicity	✗	✓

Table 2. Summary of the findings of the study. In *E. coli* essentiality correlates with low evolvability. There is a slight enrichment of essential proteins among GroEL substrates. Expression level does not correlate with evolvability. GroEL substrate proteins are slightly higher expressed than proteins not folding with GroEL. SCOP classes have different rates of evolution. GroEL substrates have a distinct distribution of SCOP classes. Hydrophobicity and evolvability are unrelated. GroEL substrates are less hydrophobic than proteins not folding with GroEL

Our data suggest that, although GroEL was shown to buffer against deleterious mutations when overexpressed (6), appear to have functioned as a capacitor for evolution *in vivo*. In fact, we observe that GroEL substrate proteins in different gamma- proteobacteria evolve less than chaperonin-independent folders. This finding does not necessarily contradict the current view of how chaperone interaction during protein folding conveys a higher tolerance for mutations. We attributed the finding that GroEL substrate proteins evolve less mainly to the specific structural composition of the GroEL substrate set (Figure 4). To a lesser extent, the observed difference in evolvability is also due to an enrichment in essential proteins among GroEL-dependent folders. Protein abundance is not significantly different between GroEL substrate proteins and proteins folding independently of GroEL. This, together with the finding that GroEL can be depleted in *E. coli* without affecting viability (9), suggest that GroEL at native levels is not saturated with substrate proteins, even though they are expressed to a level above average. We therefore suggest that GroEL, at native levels, offers a fast, initial response mechanism to cellular stresses such as heat shock, before sigma-32 mediated over-expression of GroEL leads to additional available chaperone to accommodate for folding stress.

We found that GroEL substrate proteins are significantly less hydrophobic than other *E. coli* proteins. Hydrophobicity was shown not to correlate with the evolvability of proteins (Figure 5). The hydrophobicity scores of GroEL substrate proteins cluster in a much smaller range than that observed for chaperonin-independent folders. We believe that proteins with low hydrophobicity might not expose enough hydrophobic residues during their folding process to be recognized by the apical domains of the GroEL te-

tridecamer as a substrate protein. Proteins with a very high content of hydrophobic residues might undergo an initial collapse during their folding process, burying hydrophobic residues in the core of the protein, thereby removing them effectively from the pool of potential GroEL substrates. This could explain that virtually no GroEL substrate was identified with hydrophobicity values larger than 0.1 on the Kyte-Doolittle scale. These findings establish hydrophobicity as a criterion to exclude *E. coli* proteins from folding with the help of GroEL, similar to the observed size cut-off for GroEL substrates due to the limited capacity of the GroEL cavity (7, 37).

An ongoing debate is addressing the influence of translation fidelity on evolvability, related to an organism specific codon bias (20, 38). Since GroEL recognizes and folds its substrates post-translationally (39), an introduced codon bias due to synonymous mutations was not considered relevant for the analysis of the evolvability of chaperonin substrates.

Even though, based on our study, the hypothesis that chaperones function as evolutionary buffers does not seem to hold for the *in vivo* GroEL substrate proteins in *E. coli*, it remains an intriguing theory with much supporting data from different organisms (1, 2, 6, 40, 41). We suggest an experimental approach to validate the hypothesis in bacteria *in vivo*, employing modern high-throughput DNA sequencing techniques and quantitative proteomics: engineered *E. coli* mutator strains with regulatable levels of GroEL/ES expression could be grown for many generations expressing different levels of GroEL and GroES. A comparison of both the DNA sequences of the respective genomes and the sets of isolated and quantified GroEL substrate proteins before and after the growth experiments would allow one to draw conclusions on the

effect chaperonin levels have on protein evolution. In addition, the analysis of acquired mutations of isolated and quantified GroEL substrates would potentially shed light on a yet unresolved question in biochemistry: what makes a protein a chaperone substrate. The publication of more complete chaperone substrate sets would allow additional bioinformatics-based evaluation of the buffering hypothesis for different organisms and different chaperone systems, including Hsp70 and Hsp90.

Methods

GroEL substrate-set

Kerner et al. distinguished three classes of GroEL substrate proteins and validated them experimentally (7). Class I proteins are abundant cellular proteins, which were also identified as GroEL interactors. Class II substrates are proteins which can use both GroEL and other chaperone systems for folding. Class III proteins stringently depend on GroEL for productive folding. For this study we excluded the identified substrate proteins belonging to class I, since we believe that these highly abundant cellular proteins do not represent typical GroEL substrates, but rather interact with GroEL on a stochastic basis. The presented data is hence based on 204 substrate proteins, comprising both class II and class III proteins. Taking into account only stringent GroEL substrates (84 proteins) does not significantly change the results of this study.

Statistical setup

To assess the statistical significance of the differences between the GroEL substrates and the entire proteome we consistently adopted the following procedure, which hinges on the fact that the substrates are nothing but a particular subset of the proteome of cardinality n , and is

independent of the specific feature we are looking at.

1. We form N groups of cardinality n , randomly extracting them from the proteome. We always take $N=5000$.
2. For each of the subsets i , we compute its mean m_i , $i=1 \dots 5000$. Notice that for large N we expect the m_i 's to be Gaussianly distributed, with the same average as the population's.
3. For each feature under examination, we compare the mean of the GroEL substrates m_{GroEL} with the mean m and the standard deviation σ of the m_i 's. It is then possible to give a Z-score equal to $(m - m_{\text{GroEL}}) / \sigma$.

Estimation of evolutionary distance

We downloaded the sequences of the proteins of all the organisms considered in this study from the KEGG database (<ftp://ftp.genome.jp/pub/kegg/genes/organisms>). For each transcribed gene of *E. coli* we then computed the corresponding closest gene in all other eight organisms. Closeness is defined by sequence identity calculated according to the Needleman-Wunsch algorithm (42) as implemented in NEEDLE (43). In this study we estimate the evolutionary distance as the ratio dN/dS .

To determine the evolutionary distance for each pair of genes assessed, we ran the software `yn00` from the package PAML (44). This in turn implied first aligning the amino acid sequence of each gene product (for which we used MUSCLE (45)) then using this result to align the nucleotide sequences via TRANALIGN (43). This step was performed to make sure that the procedure introduces only gaps in multiples of three nucleotides, and hence does not produce artifact stop codons in the final result. Note the `yn00`

program returns a number of possible distances; we always use the one described in (46).

We noticed that in all cases where the number of substrates pairs stay in a suitable proportion with respect to the number of protein pairs found after imposing the similarity thresholds, the Z-scores computed as in Section "Statistical setup" remain significant ($>3 \sigma$) and negative (Table 1), pointing towards a diminished evolutionary rate of the GroEL substrates.

Expression level

Expression data were taken from the database GEO (Gene Expression Omnibus, <http://www.ncbi.nlm.nih.gov/geo/> (22)) as our expression data source. Five data-sets with a sufficient number of expression profiles from wild type (WT) *E. coli* strains were randomly selected. The analyzed data-sets were: GDS680 (7 WT expression profiles, Z-score: 1.7), GDS1099 (all 15 expression profiles were considered here, since all were run on WT, although with different media/growing conditions, Z-score: 0.61), GDS2181 (6 WT expression profiles, Z-score: 3.3), GDS2768 (4 WT expression profiles, Z-score: 0.12) and GDS2825 (5 WT expression profiles, Z-score: 0.3). Expression values were averaged within each data-set, yielding five different expression estimates for each gene. The processed data-sets showed a very high correlation. The estimates obtained from data-set GDS2181 were used throughout this study.

The Z-scores computed as in Section "Statistical setup" point towards higher expression levels for GroEL substrates, although the significance of the result varies among the different datasets considered.

Hydrophobicity

The hydrophobicity of the proteins in this study was assessed with PEPWINDOW, a software which gives scores based on the Kyte-Doolittle index (35) and TANGO (36), which essentially analyses the aggregation propensity of proteins. Both algorithms gave comparable results. Not to introduce a bias, in this study membrane proteins were excluded for the analysis of protein hydrophobicity. For those results, the Z-score computed as in Section "Statistical setup" is -5.1

Protein structure

In the SCOP database version 1.73, 182 of the 204 GroEL substrate proteins had SCOP classes

assigned (and 2644 of the 3928 annotated *E.coli* proteins in total). They fall into the following classes: a: 10, b: 13, c: 109, d: 41, e: 6, f: 2, g: 1.

To assess the contribution of structural properties to the evolutionary rates of GroEL substrates, we extracted randomly N=1000 samples from the *E.coli* proteome. To build each sample, we choose randomly 10 proteins of class a, 13 of class b, etc... as to reflect the structural properties of the GroEL substrates. We then applied the statistical procedure described above to the sample distribution. We further extracted only the TIM barrel proteins, to compare them with the rest of the proteome as described in the main text.

Acknowledgements

TM is funded by an EMBO long-term fellowship. We would like to thank Ben Lehner for critically reading and correcting the manuscript.

Footnotes

ER and PR designed and performed the computational analysis and analyzed the data. LS contributed intellectually and corrected the manuscript. TM had the idea, planned the study, prepared the figures and wrote the manuscript.

The authors declare no conflict of interest in connection with the manuscript.

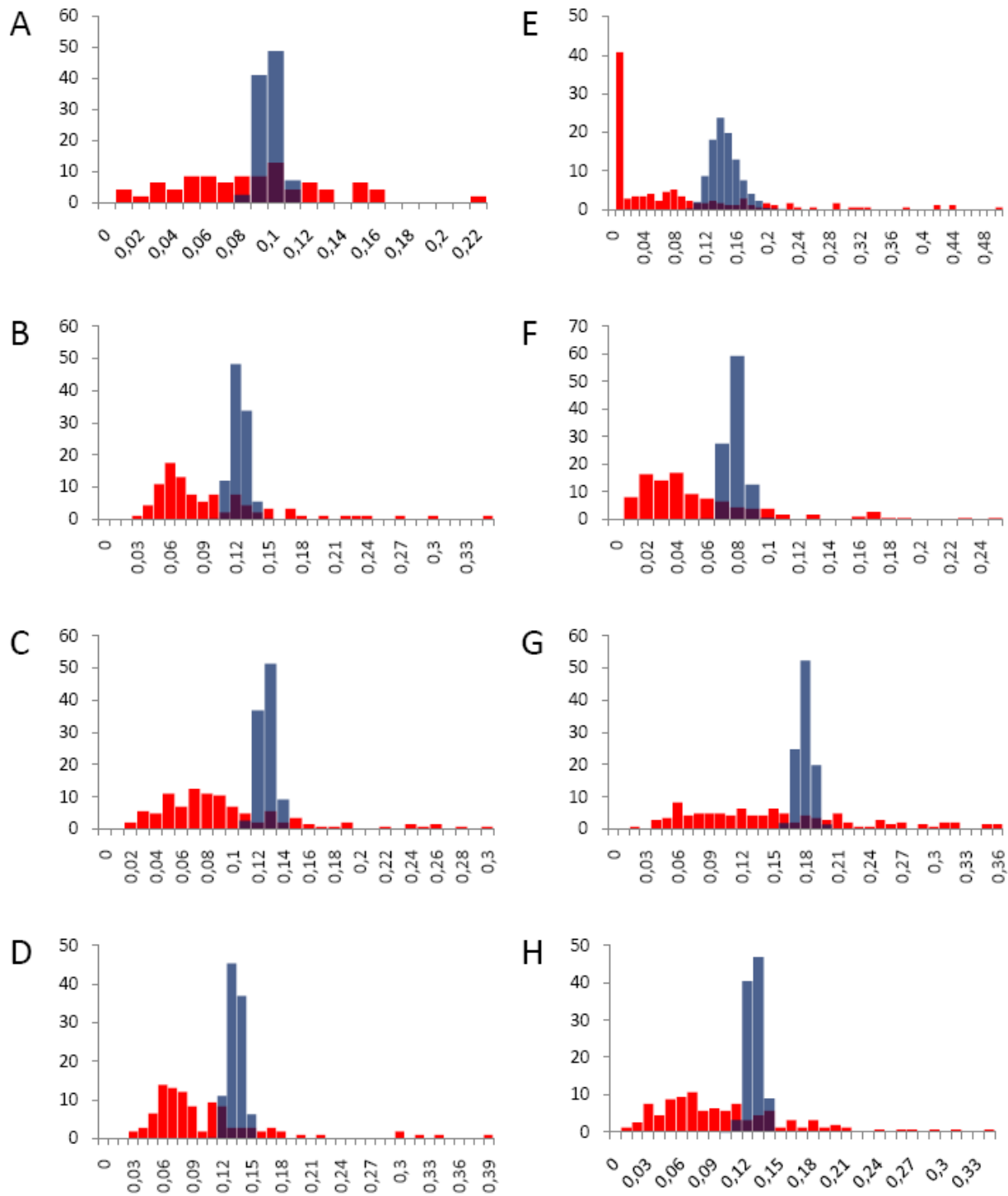
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Supplementary Figure 1. Histograms of evolutionary distances between GroEL substrates and all proteins for pairs of *E. coli* and different gamma-proteobacteria. Red bars: Distribution of evolutionary distances of GroEL substrate genes. Blue bars: Distribution of evolutionary distances of 5000 random sub-sets of gene pairs, each comprising 204 members. The abscissa shows dN/dS , the ordinate the fraction of genes of the data sets in % falling in each bin. **A:** *Buchnera aphidicola*; **B:** *Haemophilus influenzae*; **C:** *Photorhabdus luminescens*; **D:** *Pasteurella multocida*; **E:** *Shigella flexneri*; **F:** *Salmonella typhimurium*; **G:** *Vibrio cholerae*; **H:** *Yersinia pestis*. Averages for respective evolutionary distances are given in Table 1. Note that for illustrative purposes Figure 2 A/B and Figure S1 F are identical.

GroEL substrate proteins with high evolutionary distance			
Gene name	Protein name	Uniprot ID	Function
b0178	SKP	P0AEU7	Chaperone protein skp
b2095	GatZ	P0C8J8	D-tagatose-1,6-bisphosphate aldolase subunit
b0776	BioF	P12998	8-amino-7-oxononanoate synthase
b3092	Uxac	P0A8G3	Uronate isomerase
b1106	ThiK	P75948	Thiamine kinase
Essential GroEL substrate proteins			
Gene name	Protein name	Uniprot ID	Function
b0154	HemL	P23893	Glutamate-1-semialdehyde 2,1-aminomutase
b0181	LpxA	P0A722	Acyl-UDP-N-acetylglucosamine O-acyltransferase
b0185	AccA	P0ABD5	Acetyl-coA carboxylase carboxyl transferase alpha
b0369	HemB	P0ACB2	Delta-aminolevulinic acid dehydratase
b1093	FabG	P0AEK2	3-oxoacyl-[acyl-carrier-protein] reductase
b1204	Pth	P0A7D1	Peptidyl-tRNA hydrolase
b1207	KprS	P0A717	Ribose-phosphate pyrophosphokinase
b1215	KdsA	P0A715	2-dehydro-3-deoxyphosphooctonate aldolase
b1719	ThrS	P0A8M3	Threonyl-tRNA synthetase
b2153	FoIE	P0A6T5	GTP cyclohydrolase 1
b2231	GyrA	P0AES4	DNA gyrase subunit A
b2478	DapA	P0A6L2	Dihydrodipicolinate synthase
b2533	SuhB	P0ADG4	Inositol-1-monophosphatase
b2607	TrmD	P0A873	tRNA (guanine-N(1)-)-methyltransferase
b2608	RimM	P0A7X6	Ribosome maturation factor rimM
b2925	FbaA	P0AB71	Fructose-bisphosphate aldolase class 2
b2942	MetK	P0A817	S-adenosylmethionine synthetase
b3019	ParC	P0AFI2	DNA topoisomerase 4 subunit A
b3168	InfB	P0A705	Translation initiation factor IF-2
b3251	MreB	P0A9X4	Rod shape-determining protein mreB
b3256	AccC	P24182	Biotin carboxylase
b3433	Asd	P0A9Q9	Aspartate-semialdehyde dehydrogenase
b3463	FtsE	P0A9R7	Cell division ATP-binding protein ftsE
b3650	SpoT	P0AG24	Guanosine-3',5'-bis (di-P) 3'-pyrophosphohydrolase
b3783	Rho	P0AG30	Transcription termination factor rho
b3850	HemG	P0ACB4	Protoporphyrinogen oxidase
b3865	EngB	P0A6P7	Probable GTP-binding protein engB
b3982	NusG	P0AFG0	Transcription antitermination protein nusG
b3987	RpoB	P0A8V2	DNA-directed RNA polymerase subunit beta
b3988	RpoC	P0A8T7	DNA-directed RNA polymerase subunit beta'
Highly expressed GroEL substrate proteins			
Gene name	Protein name	Uniprot ID	Function
b3847	PepQ	P21165	Proline peptidase
b3565	XylA	P00944	Xylose isomerase
b1066	RimJ	P0A948	ribosomal protein alanine acetyltransferase
b2096	GatY	P0C8J6	D-tagatose-1,6-bisphosphate aldolase subunit
b2095	GatZ	P0C8J8	D-tagatose-1,6-bisphosphate aldolase subunit
Most hydrophobic GroEL substrate proteins			
Gene name	Protein name	Uniprot ID	Function
b0154	HemL	P23893	Glutamate-1-semialdehyde 2,1-aminomutase
b2091	GatD	P0A9S3	Galactitol-1-phosphate 5-dehydrogenase
b2107	KprS	P0A717	Ribose-phosphate pyrophosphokinase

Supplementary Table 1. Lists of GroEL substrate proteins and attributed properties: 1. High evolutionary distance 2. Essentiality; 3. High expression level; 4. High hydrophobicity.