Abstract

Proteins from thermophiles are generally more thermostable than their mesophilic homologs, but little is known about the evolutionary process driving these differences. Here we attempt to understand how the diverse thermostabilities of bacterial ribonuclease H1 (RNH) proteins evolved. RNH proteins from *Thermus thermophilus* (ttRNH) and *Escherichia coli* (ecRNH) share similar structures but differ in melting temperature (*Tₘ*) by 20°C. ttRNH’s greater stability is caused in part by the presence of residual structure in the unfolded state, which results in a low heat capacity of unfolding (∆C_p) relative to ecRNH. We first characterized RNH proteins from a variety of extant bacteria and found that *Tₘ* correlates with the species’ growth temperatures, consistent with environmental selection for stability. We then used ancestral sequence reconstruction to statistically infer evolutionary intermediates along lineages leading to ecRNH and ttRNH from their common ancestor, which existed approximately 3 billion years ago. Finally, we synthesized and experimentally characterized these intermediates. The shared ancestor has a melting temperature between those of ttRNH and ecRNH; the *Tₘ*S of intermediate ancestors along the ttRNH lineage increased gradually over time, while the ecRNH lineage exhibited an abrupt drop in *Tₘ* followed by relatively little change. To determine whether the underlying mechanisms for thermostability correlate with the changes in *Tₘ*, we measured the thermodynamic basis for stabilization—∆C_p and other thermodynamic parameters—for each of the ancestors. We observed that, while the *Tₘ* changes smoothly, the mechanistic basis for stability fluctuates over evolutionary time. Thus, even while overall stability appears to be strongly driven by selection, the proteins explored a wide variety of mechanisms of stabilization, a phenomenon we call “thermodynamic system drift.” This suggests that even on lineages with strong selection to increase stability, proteins have wide latitude to explore sequence space, generating biophysical diversity and potentially opening new evolutionary pathways.

Introduction

Protein thermostability is almost certainly tuned by natural selection. The fold of a protein is sensitive to denaturation at high temperatures: above the melting temperature (*Tₘ*) proteins lose structure, function, and become prone to aggregation. In laboratory evolution experiments, selection for growth at elevated temperatures leads to increases in *Tₘ* [1]. In natural systems, proteins from thermophilic organisms tend to have higher *Tₘ*s than homologs from their mesophilic counterparts [2]. Finally, there are good theoretical reasons to believe that natural selection, but not neutral drift, can lead to a sustained increase in *Tₘ*: because random amino acid substitutions tend to decrease protein stability, the final *Tₘ* of a protein is expected to be the result of a balance between selection to maintain adequate stability and mutational pressure that drives stability downward [3].

Given the functional importance of thermostability, as well as its utility in protein engineering, many studies have characterized the mechanisms by which stability is achieved [4,5]. Detailed comparisons of mesophilic and thermophilic homologs have revealed many differences that increase *Tₘ*, such as novel interactions in the folded state and residual structure in the unfolded state. These underlying biophysical differences, and the sequence differences that encode them, are usually interpreted as the direct product of selection during adaption to high-temperature environments [5,6]. Such narratives regarding natural selection, however, are essentially “just-so” stories with little or no empirical justification [7]: many of the mechanistic
Author Summary

The biophysical properties of proteins must adjust to accommodate environmental temperatures because of the narrow range over which any given protein sequence can remain folded and functional. We compared the evolution of homologous bacterial enzymes (ribonucleases H1) from two lineages: one from Escherichia coli, which live at moderate temperatures, the other from Thermus thermophilus, which live at extremely high temperatures. Our aim was to investigate how these structurally homologous proteins can have such different thermostabilities, unfolding at temperatures that are 20°C apart. We used bioinformatics to reconstruct the sequences of ancestral proteins along each lineage, synthesized the proteins in the lab, and experimentally traced the evolution of ribonuclease H1 stability. While thermostability appears to have been strongly shaped by selection, the biophysical mechanisms used to tune protein stability appear to have varied throughout evolutionary history; this suggests that proteins have wide latitude to explore different mechanisms of stabilization, generating biophysical diversity and opening up new evolutionary pathways.

Results

The Stabilities of Extant RNH Proteins Correlate with Environmental Temperature

To characterize the relationship between RNH thermostability and environmental temperature (T\textsubscript{eqv}), we measured the energetics of ten modern RNH proteins from organisms with a wide variety of optimal growth temperatures [20,21]. Growth temperatures were culled from the literature [22,23], with the exception of two mesophiles for which laboratory culture temperatures were used [24,25]. We determined the proteins’ melting temperatures by carrying out thermally induced denaturation studies monitoring the circular dichroism (CD) signal at 222 nm (see Materials and Methods) (Table S1). We observe a strong correlation (R\textsuperscript{2} = 0.84) between T\textsubscript{eqv} and T\textsubscript{m} (Figure 1; Table S1), despite uncertainty in the exact values of environmental growth temperature. Overall, for every 10°C increase in growth temperature, the T\textsubscript{m} increases by about 7–8°C. This trend remains even after removing the most thermostable protein, which is from T. thermophilus (Figure 1).

We also measured the global stabilities (∆G\textsubscript{mst})—the free energy of unfolding, which determines the ratio of unfolded and folded molecules at a given temperature—for a subset of these proteins over a range of temperatures. Unfolding free energies were measured by GdmCl-induced denaturation monitored by the change in CD signal at 222 nm (Table S1) and fit using a two-state linear extrapolation model. We found that ∆G\textsubscript{mst} was similar for all proteins at their environmental temperatures, ranging from 4.6–8.1 kcal mol\textsuperscript{-1} (average 6.28±1.5 kcal mol\textsuperscript{-1}). Together, these results suggest that RNH stability has evolved to accommodate diverse environmental temperatures across bacterial taxa.

Ancestral Sequence Reconstruction along Two RNH Lineages

We then used ASR to trace the divergence of the mesophilic ecRNH and thermophilic ttRNH from their common ancestor. Using 409 representative bacterial RNH protein sequences (Figure S1), we inferred the best-fit substitution model and maximum likelihood phylogeny. We then reconstructed the amino acid sequence at seven ancestral nodes along two lineages starting from their most recent common ancestor (Figure 2A). Anc1 represents the most recent common ancestor of ecRNH and ttRNH and is estimated to have existed approximately 3 billion years ago [26]. The other resurrected sequences are evolutionary intermediates at successive phylogenetic nodes along the lineage from Anc1 to ecRNH (AncA through AncD) and from Anc1 to ttRNH (Anc2 and Anc3). Intermediate nodes were chosen for their strong statistical support and for being spaced similarly along the two lineages: Anc2 and AncA each share 92% identity with Anc1, whereas Anc3 and AncB are 77% and 70% identical to Anc1, respectively (Figure S2). We performed the ASR using this unrooted bacterial phylogeny. To order the ancestral nodes in time, we rooted the tree by adding 43 archael RNH sequences to the bacterial alignment. The archael sequences formed a well-supported clade, providing an unambiguous root. The relative relationships of ecRNH, ttRNH, and their ancestors remained unchanged upon the addition of the archael sequences (Figure S2A).
The inferred ancestral sequences are well supported, with mean posterior probabilities per site ranging from 0.96 to 0.90 (Table S2). Most of the ambiguously reconstructed sites are in the C-terminus, which is poorly conserved among extant RNH proteins, varies in length, is largely unstructured or missing in crystal structures [27,28], and can be deleted from ecRNH without affecting function in vivo [29]. Most sites outside the C-terminus were reconstructed without ambiguity, and the handful of plausible alternative reconstructions (defined as those having posterior probability >0.3) were virtually all chemically similar to the maximum likelihood reconstruction.

As with all ASR studies, the reconstructed ancestral sequences are statistical approximations rather than certainties: the total posterior probability that the ancestral sequences are precisely correct ranged from 2.6 × 10⁻⁹ to 9.8 × 10⁻⁷. As discussed below, we addressed this uncertainty in two ways. First, we directly tested the impact of statistical uncertainty on our estimation of $T_m$ by experimentally characterizing ten alternative reconstructions for the deepest node, Anc1. Second, by resurrecting numerous sequences along two diverging lineages and focusing on broad trends rather than the properties of a single ancestor, we minimized the likelihood that statistical error would account for our overall observations.

**Resurrected RNH Proteins Are Folded and Functional**

Our first goal was to verify that the reconstructed ancestral proteins behaved like members of the RNH family by characterizing their structural and functional properties. Using far-UV CD, we found that all ancestors are folded at 25°C and exhibit secondary structure consistent with extant RNHs (Figure 2B). We determined the crystal structure of one representative ancestor, AncC, at 1.3 Å resolution (Table S3) and found that it adopts the canonical RNH fold, superimposing with the ecRNH structure with average C$_\alpha$ RMSD of 0.8 Å and with the ttRNH structure with an average C$_\alpha$ RMSD of 1.3 Å (Figure 2C). Finally, we assayed the reconstructed ancestral proteins for their ability to degrade RNA:DNA hybrids using a simple hyperchromic assay to follow RNA cleavage and nucleotide release [30,31]: all ancestors were active at 25°C, demonstrating that they are all functional ribonucleases H (Figure S3A).

**T. thermophilus and E. coli Lineages Exhibit Opposite Trends in Thermostability**

Trends in protein stability over the two evolutionary lineages were determined by measuring the $T_m$s of the ancestral proteins (Figure S3B). Anc1 has a $T_m$ of 77°C, which is intermediate between ecRNH (68°C) and ttRNH (38°C). Starting from Anc1 and proceeding toward the extant ecRNH and ttRNH, the thermophilic and mesophilic lineages exhibit opposite trends (Figure 3A). $T_m$s increase along the thermophilic branch and decrease along the mesophilic branch, with ttRNH showing the highest $T_m$ and ecRNH showing second lowest $T_m$. The temperature-induced unfolding of ancestors along the mesophilic lineage becomes irreversible starting with AncB and continuing through AncC, AncD, and ecRNH. For these proteins, the unfolding curve does not reflect a strictly equilibrium process, so the midpoint of the unfolding curve is therefore only an apparent $T_m$ (Table S1).

To verify that these trends were robust to uncertainty in the ASR, we experimentally characterized ten alternate reconstructions of Anc1. We generated these sequences computationally by randomly sampling at each site in the sequence an amino acid from the posterior distribution of states at that site, excluding implausible states with posterior probabilities <0.2. The resampled ancestors differed from each other and from the maximum likelihood sequence by two to 11 amino acids, with a mean of seven differences. We then synthesized coding DNAs, expressed the proteins, and measured the $T_m$s of each resampled ancestor. The $T_m$ for the ten alternate versions of Anc1 ranged from 75.6°C to 80.5°C with a mean of 78.2°C, comparable to the $T_m$ for the maximum likelihood Anc1 of 76.8°C (Table S1). The range of the phylogenetically plausible $T_m$s for Anc1 (+1.6°C standard deviation [SD]) is much smaller than the overall changes in $T_m$ along the ecRNH (−9°C) and ttRNH lineages (+12°C) (Figure S3A), indicating that the overall trends in stability are robust to uncertainty in the sequence of Anc1.

**Trends in $\Delta G_{\text{unfold}}$ at 25°C Mirror Changes in Thermostability**

To supplement the thermal melting data, we carried out GdnCl-induced denaturation studies to determine the $\Delta G_{\text{unfold}}$ at 25°C (Figure S3C; Table S4). All ancestors unfolded reversibly upon chemical denaturation and are well described using the two-state assumption, except AncB, which deviates from two-state behavior and was removed from all further analyses. The trends in $\Delta G_{\text{unfold}}$ mirror the trend in $T_m$: $\Delta G_{\text{unfold}}$ increases by 1.9 kcal mol$^{-1}$ along the thermophilic lineage and decreases by the same amount on the mesophilic lineage (Figure 3B).

**Analysis of the Global Stability Curve Reveals the Mechanism of Stabilization**

Our previous work revealed that $\Delta G_p$ of ttRNH is lower than that of ecRNH (1.9 versus 2.9 kcal mol$^{-1}$ K$^{-1}$) because it retains residual structure in the unfolded state [32,33]; this difference contributes 10.7°C to the observed difference in $T_m$ between the two proteins. This stabilization mechanism might be adaptive, allowing the protein to achieve a high $T_m$ while maintaining conformational flexibility and a moderate $\Delta G$ at its growth temperature; residual structure in the unfolded state could also help minimize aggregation under unfolding conditions [3,17,33]. We therefore hypothesized that the evolution of residual structure in the unfolded state was the mechanism by which adaptive thermostability evolved along the thermophilic lineage.

To test this hypothesis and elucidate the mechanisms by which stability was tuned along these lineages, we determined the global
protein stability curve for each ancestor (Figure S4). This curve, a plot of $\Delta G_{\text{unf}}$ against temperature, can be determined by performing chemical denaturation experiments across a wide range of temperatures. It exhibits a characteristic parabolic shape that is described by the Gibbs-Helmholtz equation:

$$\Delta G_{\text{unf}}(T) = \Delta H_s + \Delta C_p \left( T - T_s - T \ln \left( \frac{T}{T_s} \right) \right)$$  \hspace{1cm} (1)

This equation defines the temperature-dependent relationship between the global stability of a protein ($\Delta G_{\text{unf}}$) and three key thermodynamic parameters: the temperature of maximum stability ($T_s$), the change in enthalpy upon unfolding at $T_s$ ($\Delta H_s$), and the change in heat capacity upon unfolding ($\Delta C_p$). The $T_m$, or thermal melting temperature, is the right-most $x$-intercept of this curve. By fitting the parameters of the equation to our data, we can extract detailed thermodynamic and mechanistic information about each
Thermodynamic System Drift in Protein Evolution

Different Ancestors Use Different Mechanisms of Stabilization

All of the ancestors have \( \Delta G_p \)'s falling within a narrow range (Figure 4C). To test the hypothesis that changes in the heat capacity of unfolding caused the gradual increase of \( T_m \) along the thermophile lineage, we carried out two different analyses. First, we analyzed the results from our fits of the stability curve for each ancestor (Figure 5A; Table S5) to quantify the relative contribution of \( \Delta C_p \), \( T_m \), and \( \Delta H_i \) to changes in \( T_m \) relative to Anc1. We assessed the change in \( T_m \) predicted when the measured parameters for each extant or ancestral RNH protein were individually substituted into the Gibbs-Helmholtz equation with all other parameters from Anc1. This analysis uses Anc1 as a reference state and assumes that the effects of each parameter can be assessed independently.

Our hypothesis predicts that \( \Delta C_p \) should decline consistently along the thermophilic lineage from Anc1 to ttRNH in a fashion correlated with the evolutionary increase in \( T_m \). As expected, \( \Delta C_p \) of ttRNH is lower than that of Anc1 (2.28±0.2 and 1.91±0.3 kcal mol\(^{-1}\) K\(^{-1}\), respectively). The elevated \( T_m \) of ttRNH, however, is caused not only by changes in \( \Delta C_p \) but also by substantial changes in \( T_m \) and \( \Delta H_i \); whereas changes in \( \Delta C_p \) caused an increase in \( T_m \) of 5.4°C, changes in \( \Delta H_i \) and \( T_m \) caused additional increases of 3.6°C and 3.4°C, respectively (Figure 5A).

Contrary to our prediction, however, the observed trends in \( T_m \) are not mirrored by trends in \( \Delta C_p \) (Figure 5A), and the underpinnings of the \( T_m \) of each ancestor are different. The \( T_m \) of ttRNH is 12°C above Anc1’s because of contributions from \( \Delta C_p \) (+5.4°C), \( \Delta H_i \) (+3.6°C), and \( T_m \) (+3.4°C). In contrast, although the \( T_m \) of Anc3 is 8°C higher than Anc1, \( \Delta C_p \) makes no contribution at all to its elevated thermostability, which is instead driven by \( \Delta H_i \) (+5.5°C) and \( T_m \) (+3.5°C). Anc2 exhibits yet another pattern: its \( T_m \) is 1°C higher than Anc1 due to a strong contribution by \( T_m \) (3.7°C), which offsets a destabilizing effect of \( \Delta C_p \) (−2.7°C), while \( \Delta H_i \) makes virtually no contribution. Thus, although \( T_m \) consistently increases along the thermophilic lineage, our analysis suggests that different stabilizing mechanisms are utilized between each ancestor.

The mechanisms altering thermostability fluctuate along the mesophilic lineage, as well (Figure 5B). The \( T_m \) of the modern mesophilic ecRNH is 11°C lower than that of Anc1. This reflects major unfavorable contributions from \( \Delta H_i \) (−6.1°C) and \( \Delta C_p \) (−6.2°C), with a minor stabilizing effect by \( T_m \) (+1.1°C). As with the thermophilic lineage, however, the contributions of changes in \( \Delta C_p \), \( \Delta H_i \), and \( T_m \) fluctuate over evolutionary time. AncC and AncD have nearly identical thermostabilities, with \( T_m \) of AncA is 5°C lower than Anc1, but \( \Delta C_p \) adding favorable interactions in the folded state. Such changes in enthalpy are a common basis for change in stability upon mutation, as small changes in side-chain functional groups can result in large changes in native-state interactions. Second, depressing \( \Delta C_p \) broadens the stability curve, increasing \( \Delta G \) at temperatures above or below \( T_m \). \( \Delta C_p \) depends largely on the difference in protein-water interactions between the unfolded and folded states; a variant that creates residual structure in the unfolded state will decrease the difference in protein-water interactions between the unfolded and folded states and thereby decrease \( \Delta C_p \). Third, increasing \( T_m \) moves the curve to the right along the x-axis. \( T_m \) is relatively insensitive to mutations and is therefore not considered a common mechanism of stabilization [34].

Figure 3. Thermophilic and mesophilic lineages exhibit opposite stability trends. Starting with the shared ancestor, Anc1, stabilities increase along the thermophilic lineage and decrease along the mesophilic lineage. (A) Melting temperature of the maximum likelihood ancestors and the ten alternate reconstructions of Anc1 as a function of evolutionary distance from the last common ancestor, Anc1. Distances are calculated as the sum of the branch lengths connecting Anc1 to the protein of interest. The grey region defines the range of \( T_m \)'s measured for the Anc1 alternates, which appear individually as grey data points. The error bars are one standard deviation. (B) Average \( \Delta G \)'s at 25°C as a function of evolutionary distance from Anc1. Error bars are one standard deviation. See also Table S1 and Dataset S2.

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(−4.2°C) and $T_s$ (+1.9°C) have now switched roles, with $\Delta H$, (−3.3°C) continuing to be destabilizing.

We statistically analyzed whether the apparent fluctuations in the mechanistic underpinnings of stability over evolutionary time were robust to uncertainty in the estimates of the parameters of the Gibbs-Helmholtz equation. We used a bootstrap and refitting approach to identify for each ancestor a cloud of plausible parameter values that are consistent with the experimental data (Figures S6 and S7). We found that the thermodynamic underpinnings of the ancestral proteins’ stabilities are indeed distinct from each other (Figure S7; Table S6). We then asked about the possible evolutionary trajectories along phylogenetic lineages given uncertainty in the parameter estimates. We discretized the parameter space for each ancestral protein and calculated from the refitted bootstrap parameter estimates the probability that the ancestral protein was in each region of this parameter space (see Materials and Methods). We then calculated the probability of each possible pathway through this space that could be taken by lineages of successive ancestors on the phylogenetic tree, given the bootstrap-derived probability that each ancestor’s parameters were in each parameter region. We found that, along both lineages, the most probable path was circuitous, with values of $T_m$ and $\Delta C_p$ in particular fluctuating across the evolutionary intervals (Figure 6A and 6B). This stands in contrast with the value of $T_m$, which increases smoothly on the thermophile lineage and, after an early drop, remains essentially constant along the mesophilic lineage (Figure 3A). We also investigated the distribution of alternative possible paths (Figure 4).

Figure 4. All possible mechanisms for manipulating $T_m$ are represented in RNH’s evolutionary history. (A) Thermodynamic strategies for increasing $T_m$. Relative to the reference state (blue line), the stability curve can be upshifted (solid black line), broadened (dashed line), or right-shifted (dotted line). (B) Superimposed stability curve fits for ecRNH (blue solid circles), ttRNH (red solid circles), Anc1 (black solid circles), ancestors from the mesophilic lineage (AncA, blue Xs; AncC, blue solid squares; AncD blue open circles), and ancestors from the thermophilic lineage (Anc2, red solid squares; Anc3, red open circles). (C) $\Delta C_p$ as function of evolutionary distance from Anc1. See also Dataset S2.

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Thus, our observed experimental data and the bootstrapped refitting both indicate that the mechanistic underpinnings of stability did not follow the smooth evolutionary trajectories followed by stability itself but instead fluctuated dramatically during the evolution of thermophilic and mesophilic RNH proteins. The parameter that appears to be most consistently associated with changes in \( T_m \) along both lineages is \( \Delta H_s \) with the noted exception of ttRNH (Figure 6C). Changes in \( \Delta H_s \) can be achieved simply by manipulating side-chain interactions in the folded state. This is the most commonly observed mechanism in thermophilic proteins [5] and is the one most often exploited for engineering protein stability [35]. Despite this general correlation, however, it is clear that no singular mechanism dominates along either lineage.

**Discussion**

**Current Mechanisms for Thermostability Differ from Historical Ones**

In this study, we used ASR to study the evolution of thermostability of RNH enzymes along mesophilic (ecRNH) and thermophilic (ttRNH) lineages. We observed that, while \( T_m \) changed smoothly over both lineages, the mechanism of stabilization followed a more tortuous path. This was particularly surprising for \( \Delta C_p \), which is the most striking thermodynamic difference between ecRNH and ttRNH, accounting for a full 10.7°C difference in \( T_m \). As such, we hypothesized that a depressed \( \Delta C_p \) would be the primary mechanism of stabilization. The fluctuations observed in the mechanistic underpinnings of function along both lineages indicate that this parameter, and the changes in residual unfolded structure that produce it, are not driven by the same selective pressures imposed on \( T_m \). Further evidence for the de-coupling of \( \Delta C_p \) and \( T_m \), coming from the modern RNH homolog from *Chlorobium tepidum*, which has a mesophile-like \( T_m \) of 66.5°C and a thermophile-like \( \Delta C_p \) of 1.7 kcal mol\(^{-1}\) K\(^{-1}\) [21].

**Implications for Inferences of Ancient Environmental Temperature**

Our findings concerning trends in \( T_m \)s bear on recent efforts to infer long-term changes in global environmental temperature based on studies of resurrected ancestral proteins. We observed that the RNH of the ancestor of thermophilic and mesophilic bacteria had a \( T_m \) intermediate between those of extant members of the two groups, with a gradual progression to higher \( T_m \)s along the thermophilic lineage and an initially abrupt evolution to lower \( T_m \)s on the mesophilic lineage followed by little long-term change. These results differ from studies of several other proteins, which observed monotonic increases in \( T_m \) as one goes back in time and interpreted this result as evidence for an ancient, global trend from higher to lower environmental temperatures [8–10,12,13]. Our oldest ancestor, Anc1, has a \( T_m \) of 10°C higher than extant mesophiles and 10°C lower than extant thermophiles, in contrast to similarly aged thioredoxins and \( \beta\)-lactamases, which were found to have \( T_m \)s 25–30°C higher than both their mesophilic and thermophilic descendants [12,13]. Our findings are more similar to those of a study showing that ancestral EF-Tu proteins have thermostabilities between those of extant thermophiles and mesophiles, a result interpreted as providing evidence for a relatively hot ancient global environment [9]. By tracing changes in \( T_m \) along multiple taxonomic lineages, we found patterns that are inconsistent with any gradual, long-term trend in global environmental temperatures: for example, Anc2, which is estimated to have existed approximately 2 billion years ago [26],...
has a melting temperature only 2°C higher than that of present-day mesophilic ecRNH. Taken together, our observations suggest that particular proteins in particular lineages undergo their own paths to accommodate the local environments they colonize and the functions they perform. This view is consistent with the wide variety of temperature niches populated by both ancient and modern bacteria. It is also consistent with recent findings that thermophilicity evolved in parallel numerous times over a period of just 3 million years in a family of enzymes involved in leucine biosynthesis [11]. We therefore suggest that the tracking the $T_m$ of any individual protein or lineage is an unreliable way to estimate long-term trends in global environmental temperatures.

RNH Exhibits Thermodynamic Systems Drift

Our work reveals that RNH thermostability is evolving in a regime analogous to “developmental systems drift” (DSD). In DSD, a developmental process or outcome is conserved by selection, but the underlying genetic or molecular mechanisms shift during evolution [36]. By analogy, RNH exhibits thermodynamic systems drift, in which the $T_m$ is under selection, but the mechanisms by which it is achieved vary over evolutionary time. “Drift,” in this usage, does not refer to the evolutionary process of neutral drift, but rather to changes in mechanism uncorrelated with changes in phenotype.

Although our data do not directly reveal the evolutionary forces that shape the mechanism of stabilization, the most parsimonious explanation of our observations is a neutral evolutionary process. A protein’s stability must be above a given threshold to maintain protein function and prevent the accumulation of misfolded protein [37,38], but there is little evidence that selection can “sense” the underlying mechanism of stabilization, implying that neutral evolution would be free to alter stabilization mechanisms while maintaining the required stability. Further, invoking selection to explain the fluctuations we observed in the thermodynamic parameters that determine stability requires that the each ancestor experienced a unique selective regime and that this regime was different from that driving stability itself (Figure 1). We believe that this scenario, although formally possible, is very unlikely.

Thermodynamic systems drift has some important implications. First, it highlights the pitfalls of proposing an adaptive trajectory based on comparisons of modern proteins. Almost all studies of thermostability have compared mesophilic and thermophilic homologs [5]; our work indicates that mechanistic differences between any two such proteins—even those that strongly correlate with stability—may reflect exploration of alternate stabilization mechanisms rather than the initial adaptive stabilization mechanism. A second implication is that studying multiple thermophilic homologs—or an evolutionary lineage leading to a thermophilic protein—has the capacity to reveal multiple methods of stabilization for a given protein, thus providing insight for engineers trying to identify stabilizing mutations by studying protein diversity. Finally, thermodynamic systems drift implies that evolving proteins can efficiently explore sequence space, even when selection establishes a threshold for stability [39]. As proteins explore sequence space, their evolutionary potential changes,

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**Figure 6.** The thermodynamic underpinnings of RNH $T_m$ fluctuated over evolutionary time. Estimated $\Delta C_p$, $T_s$, and $\Delta H_s$ over the evolution of ttRNH (red) and ecRNH (blue). Maximum likelihood path is shown in bold. The shortest (dashed line) and longest (solid line) plausible alternate trajectories (95% confidence interval) are shown in faded blue and pink. See also Dataset S2.

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Materials and Methods

Ancestral Sequence Reconstruction

Bacterial and archael RNH sequences were identified by BLAST against the NCBI non-redundant protein database using RNH from *E. coli* and *T. thermophila* sequences as seed sequences [43,44]. Redundant sequences were removed using cdhit 4.6 [45]. In total, 409 sequences were kept for further analysis. Sequences were aligned using MUSCLE 3.8.31, followed by manual refinement using Mesquite 2.75 (Maddison and Maddison). Alignment quality was verified by checking for alignment of three universally conserved acidic residues that compose the RNH active site (Figure S1). The final alignment is available in Dataset S1. The maximum likelihood phylogenetic tree was constructed using the JTT+Γ6 substitution model—identified by ProtTest [46] —and SPR moves as implemented in PhyML 3.0 [47,48]. Branch supports were estimated using the approximate likelihood ratio test [49]. Maximum likelihood ancestral RNH sequences were reconstructed with the maximum likelihood topology, branch lengths, and phylogenetic model using PAML 3.14 [50,51].

Expression and Purification

Genes encoding the ancestral proteins were codon optimized for expression in *E. coli* and synthesized by GENEART. The genes were subcloned using NdeI and HindIII restriction sites into the multiple cloning site of a pET27 vector (Life Technologies). Other site-specific variants were constructed via site-directed mutagenesis and verified by sequencing.

Plasmids were transformed into BL21 (DE3) pLysS cells for expression. Cells were induced with 1 mM IPTG at OD = 0.6 and grown at 37°C for 3 hours before harvesting. Cells were lysed in buffer via sonication. All ancestors expressed predominantly in the opening and closing pathways to new functions or properties [39–42].

Denaturant-Induced and Thermal Denaturation

Thermal and chemical denaturation melts were performed by monitoring the CD signal at 222. CD melts were carried out in a 1-cm pathlength cuvette (50 μg/ml protein, 20 mM NaOAc [pH 5.5], 50 mM KCl, and 1 mM TCEP). Samples were prepared individually, equilibrated overnight or longer for temperatures below 25°C, and allowed to stir in the instrument for 1–2 minutes before data collection. For higher temperatures, proteins were equilibrated overnight only if solubility was not compromised. Otherwise, shorter equilibration times were used, typically 2–3 hours for melts performed manually and 5–15 minutes for melts performed using an automated titrator. Data from titration experiments were only used if the measured Cm was within 0.1 M of the value collected manually at the same temperature.

To measure CD signal at 222 nm as a function of temperature, samples were allowed to equilibrate for five minutes at each temperature and data were collected every 3°C. Spectra were taken at 25°C before and after the thermal melt to test for reversibility. Reversibility was defined as recovery of 80% of CD signal at 222 nm. Temperature melts were fit to a two-state model using the Gibbs-Helmholtz relationship (Equation 1).

Denaturation and Stability Curve Data Analysis

To generate stability curves, global stabilities derived from GdmCl-induced denaturation melts were plotted as a function of temperature. Tm,s extracted from thermal denaturation experiments were used as single points at ΔG = 0 for trRNH, AncA, Anc1, Anc2, and Anc3, which all unfold reversibly. Only data collected at temperatures above 15°C were included in fits due to deviation from two-state behavior at lower temperatures (Figure S5). Data were fit to the Gibbs-Helmholtz equation to extract all thermodynamic parameters (Equation 1).
To characterize the uncertainty in our estimates of the thermodynamic parameters, we generated 10,000 bootstrap pseudo-datasets for each protein—sampling three times with replacement from the measured ΔG values at each temperature—and then refit our model to each dataset (Figure S6). This generated a distribution of parameter values consistent with the experimental measurements taken for each protein. To characterize uncertainty in the trends that parameters followed along phylogenetic lineages, we sampled possible pathways between the bootstrapped parameter values of successive ancestors along the tree and then characterized the lengths and probabilities of these pathways. We first placed all three parameters on the same scale by normalizing their values to the interval [0, 1]. Next, we discretized the parameter space into a 15×15×15 grid, reducing the 10,000 unique parameter estimates for each protein to ~35 unique estimates. We calculated the relative probability of each grid cell, given our experimental data, as the proportion of all pseudo-datasets with parameter estimates in that cell. Finally, we exhaustively sampled trajectories among successive ancestral proteins through this parameter space. For each trajectory, we calculated the path length (the sum of Euclidean distances between successive ancestor’s parameter values) and its probability (determined by multiplying the probabilities of each bin in the path). Statistical analyses were done in the R 3.1.0 statistical environment [39]. Fittings was done using the L-BFGS-B algorithm as implemented in the optim function [60].

Supporting Information

Figure S1 WebLogo representation of the RNH multiple sequence alignment [61]. Conservation is reflected by the overall height of the stack at each position. Height of individual letters within the stack indicates the relative frequency of a residue at the position. Numbering and secondary structure elements are based on ecRNH. Active site residues are starred. See also Dataset S1. (TIF)

Figure S2 RNH phylogenetic tree and sequence comparisons. (A) Rooting does not change the relative relationships between ecRNH, trRNH, and the ancestors, ASR was performed using an unrooted tree built from an alignment of 409 RNH sequences. An additional 45 archaean RNH sequences were used to create the rooted tree, which allows ordering of the ancestors in time. Branch length reflects sequence distance, as indicated by the scale bar, in average number of substitutions per position. Resurrected nodes are starred (see Table S2). Branch supports for the trees are labeled. (B) Alignment of ancestors with ecRNH and trRNH. Secondary structure elements are based on ecRNH. (C) Sequence identity matrix for ancestors, ecRNH, and trRNH. Ancestors that are analogously spaced along the thermophilic and mesophilic lineages appear in the same color. See also Datasets S3 and S4. (TIF)

Figure S3 Measuring activity and stability of ancestors, ecRNH, and ttRNH. (A) Activity at 25°C in 10 mM Tris (pH 8), 50 mM NaCl, 10 mM MgCl2, 1 mM TCEP, and 16.7 μg/ml poly-rA:dT20 substrate for ecRNH (blue solid circles), trRNH (red solid circles), Anc1 (black solid circles), ancestors from the mesophilic lineage (AncA, blue Xs; AncB, blue open squares; AncC, blue solid squares; AncD blue open circles), and ancestors from the thermophilic lineage (Anc2, red solid squares; Anc3, red open circles). (B) Chemical denaturation as probed by CD signal at 222 nm. (C) Chemical denaturation at 25°C as monitored by CD at 222 nm. See also Dataset S2. (TIF)

Figure S4 Stability curves. (A) Anc1, (B) Anc2, (C) Anc3, (D) AncA, (E) AncC, (F) AncD, (G) ecRNH, (H) trRNH. Average ΔG values measured at 15°C or higher were used for the fits, and errors are standard deviations from fits of replicate experiments. See also Dataset S2. (TIF)

Figure S5 Deviation from two-state behavior at low temperatures. (A) ΔG values of ecRNH unexpectedly plateau below 15°C. Data at 5°C and 10°C reflect the averages of 12 and seven independent experiments, respectively. The displayed fit does not include data below 15°C. (B) Stability curve fit from the cysteine-free variant ecRNH C13A/C63A/C133A (black curve) superimposed with data from single cysteine variants (ecRNH C13A/C133A, orange closed circles; ecRNH C13A/C63A, orange open circles; ecRNH C133A/C63A, orange Xs). Asymmetry in ecRNH stability data is due to C63. See also Dataset S2. (TIF)

Figure S6 Global fits to bootstrap samples of stability versus temperature curves of different RNH proteins. Points show ΔG_ref values measured for each protein as a function of temperature. Lines show a random sample of 100 re-fits of these data generated by bootstrap sampling. Colors denote different proteins: Anc1 (black), Anc2 (wheat), Anc3 (orange), trRNH (red), AncA (light blue), AncC (dark green), AncD (slate), and ecRNH (blue). Arrows show the maximum likelihood trajectory through this space, starting from Anc1 and going to trRNH (red path) or starting from Anc1 and going to ecRNH (blue). (TIF)

Figure S7 No smooth pathway exists through possible ancestral parameter space. Plots projection of 3D plot of ΔGo versus T, versus ΔH, for each protein. Points show fit parameters extracted from bootstrap replicates. Colors denote different proteins: Anc1 (black), Anc2 (wheat), Anc3 (orange), trRNH (red), AncA (light blue), AncC (dark green), AncD (slate), and ecRNH (blue). Arrows show the maximum likelihood trajectory through this space, starting from Anc1 and going to trRNH (red path) or starting from Anc1 and going to ecRNH (blue). (TIF)

Figure S8 Distribution of all possible path lengths reveals that 95% of possible paths are nearly as long as the ML path. Histograms of possible path lengths weighted by path likelihoods, extracted from explicit enumeration of pathways through the parameter space. Path lengths are normalized to the maximum likelihood path. Red and blue curves denote the thermophilic and mesophilic lineages, respectively. Dashed lines indicate 95% cutoff. (TIF)

Table S1 Growth temperatures, T_m, and ΔGs at T_m, for extant and ancestral RNH proteins. * Extracted from stability curve fits for two-state proteins. † Errors reported are standard deviations from replicate experiments. ‡ Taken from reference [20]. § Taken from reference [21]. † † Not determined. (DOCX)

Table S2 Statistical support for resurrected ancestors. (DOCX)
Table S3 Data collection and refinement statistics for AncC. *Values in parentheses are for highest-resolution shell. (DOCX)

Table S4 ΔGs and m-values at 25°C. *Errors reported are standard deviations from replicate experiments. (DOCX)

Table S5 Thermodynamic parameters from stability curve fits. *Errors from fit. †Extracted from thermal melt fit. ‡Extracted from stability curve fit. (DOCX)

Table S6 Individual proteins populate discrete regions of parameter space. (DOCX)

Dataset S1 RNH sequence alignment. (DOCX)

Dataset S2 Numerical data underlying main text and supplemental figures. (XLSX)

Dataset S3 Phylogenetic tree file used to generate the unrooted tree in Figure S2. Tree files are in the standard “newick” ascii text format and can be opened via a wide variety of freely available and commercial tree-viewing programs. Two free programs are FigTree (http://tree.bio.ed.ac.uk/software/figtree/) and archaceopteryx (https://sites.google.com/site/czmsmasek/home/software/archaceopteryx). (TXT)

Dataset S4 Phylogenetic tree file used to generate the rooted tree in Figure S2. Tree files are in the standard “newick” ascii text format and can be opened via a wide variety of freely available and commercial tree-viewing programs. Two free programs are FigTree (http://tree.bio.ed.ac.uk/software/figtree/) and archaceopteryx (https://sites.google.com/site/czmsmasek/home/software/archaceopteryx). (TXT)

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Author Contributions

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