Adaptive Threonine Increase in Transmembrane Regions of Mitochondrial Proteins in Higher Primates

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Abstract

Background: The mitochondrial (mt) gene tree of placental mammals reveals a very strong acceleration of the amino acid (AA) replacement rate and a change in AA compositional bias in the lineage leading to the higher primates (simians), in contrast to the nuclear gene tree. Whether this acceleration and compositional bias were caused by adaptive evolution at the AA level or directional mutation pressure at the DNA level has been vigorously debated.

Methodology/Principal Findings: Our phylogenetic analysis indicates that the rate acceleration in the simian lineage is accompanied by a marked increase in threonine (Thr) residues in the transmembrane helix regions of mt DNA-encoded proteins. This Thr increase involved the replacement of hydrophobic AAs in the membrane interior. Even after accounting for lack of independence due to phylogeny, a regression analysis reveals a statistical significant positive correlation between Thr composition and longevity in primates.

Conclusion/Significance: Because crucial roles of Thr and Ser in membrane proteins have been proposed to be the formation of hydrogen bonds enhancing helix-helix interactions, the Thr increase detected in the higher primates might be adaptive by serving to reinforce stability of mt proteins in the inner membrane. The correlation between Thr composition in the membrane interior and the longevity of animals is striking, especially because some mt functions are thought to be involved in aging.

Introduction

Mitochondria supply most cellular energy and influence cell growth, human disease, and probably aging [1]. A better understanding of mt protein functions may be possible by detecting adaptive evolution that occurred in specific lineages of the mt gene tree of the placental mammals. Interestingly, the mt gene tree displays elevated rates of AA replacement in lineages leading to the simians, rodents, and hedgehogs (Figure 1) [2]. Although the nuclear gene tree also shows rate accelerations in the lineages of rodents and hedgehogs, no significant acceleration is observed in the primate lineage [3,4] whereas the mt branch lengths of the simians (relative to the total branch lengths) are twice as long as those on the nuclear gene tree. This feature of the mt gene tree suggests that the simian lineage experienced a particularly unusual event in molecular evolution. One hypothesis is that an episode of adaptive evolution occurred in the simian lineage [5–9]. Another hypothesis is that the acceleration and compositional bias were caused by a higher rate of directional mutation [10,11]. The adaptive hypothesis is supported by a highly significant acceleration of nonsynonymous changes, in contrast to a rather homogeneous rate of synonymous changes. The mutation bias explanation is supported by the existence of directional nucleotide mutation pressure at the fourfold-degenerate third nucleotide sites (FD3rd) in codons [12–14], and also by the duration of the single-stranded state of the 12 mt protein-coding genes (except for NADH dehydrogenase subunit 6) during replication [12].

Here, we find that the rate acceleration in the simian lineage is accompanied by a marked increase in Thr residues within the transmembrane region of mt proteins. This Thr increase may be adaptive because it is correlated with the protein structure. An essential role of Thr residues is to stabilize mt proteins by forming helix-helix interactions [15–19]. We also find an intriguing evolutionary correlation between the increase in maximum life span and the increase in Thr composition of membrane interiors.

Results

Decrease in hydrophobic AAs and increase in Thr residues on the placental mammal tree

We first report that the 12 mt protein-coding genes underwent, at an early stage of simian evolution, a net flow of AA
Fin whale
Blue whale
Sperm whale
Hippopotamus
Cow
Sheep
Pig
Alpaca
Horse
Donkey
I. rhinoceros
W. rhinoceros
Harbor seal
Gray seal
Polar bear
Dog
Cat
Ryukyu flying fox
Little flying fox
Horseshoe bat
Formosan bat
Jamaican fruitbat
J. pipistrelle
Long-tailed bat
European mole
J. mole
J. shrew-mole
L. c. shrew
Formosan shrew
Hedgehog
L. hedgehog
Moonrat
Chimpanzee
Bonobo
Human
Gorilla
B. orangutan
S. orangutan
Gibbon
Baboon
Macaque
Capuchin
Slow-loris
Tree shrew
Mouse
Rat
Vole
Guinea pig
Cane rat
Fat dormouse
E. red squirrel
Rabbit
American pika
African elephant
Dugon
Cape hyrax
Aardvark
Tenrec1
Tenrec2
Elephant shrew
Cape golden mole
Armadillo

Adaptive Mt-Thr Increase

Supraprimates

Afrotheria

Laurasiatheria

Xenarthra

0.1
replacements from hydrophobic AAs (Hoa) to Thr. We refer to this compositional change as the Hoa→Thr flow. For this flow analysis, we classify eight amino acid types (Ala, Cys, Ile, Leu, Met, Phe, Pro and Val) as hydrophobic (Table S1). The Hoa→Thr flow violates the “detailed balance” condition that would have the numbers of AA replacements from Hoa to Thr be about equal to that from Thr to Hoa. Figure 1 plots the numbers of Hoa and Thr residues (relative to those of the root sequence) in each inferred node sequence on a placental mammal tree [2]. The Hoa→Thr flow appeared specifically in the long branches leading to the simians, as well as those leading to the sperm whale and elephant. However, the flow did not occur in the long branches of the mt or nuclear gene trees leading to the rodents or hedgehogs.

Hoa→Thr flow in the higher primate lineage

By chronologically tracing the greatest flow along the simian lineage, we detect a strong correlation between the decrease in Hoa and the increase in Thr (Figure 2A and B). The tree shrew lineage, with a very short branch (Figure 1), had a minimal flow, and the loris lineage, with a moderately long branch, had a small flow. We also observe that the Hoa→Thr flow is associated with an accelerated AA evolutionary rate (Figure 2C) in the early stages of primate evolution previously reported [20]. This observation seems to be consistent with the higher rates of nonsynonymous relative to synonymous substitutions that occurred in complexes IV (cytochrome c oxidase subunits) [5–9]. By applying the flow analysis to each protein, we found that the Hoa→Thr flow was generally large in complexes I (NADH dehydrogenase subunits 1, 2, 4 & 5; ND1, ND2, ND4 & ND5) and III (cytochrome b; Cytb), and rather small in complexes IV (cytochrome c oxidase subunits) and V (ATPase subunits). We note that ND2 displayed a Hoa→Ser flow in addition to the Hoa→Thr flow.

Hydrophobicity dependence of Hoa→Thr flow

As described in Materials and Methods, we analyzed in more detail the Hoa→Thr flow in the three lineages of simians and also sperm whale and elephant, by locally estimating the node sequences of the three lineages at the DNA level. The analysis displayed both substantial increases in Thr/Leu residues and reductions in Ile/Met residues (Figure 3A). The increase in hydrophobic Leu residues accompanied a compensatory decline in other hydrophobic residues. Consequently, only the Hoa→Thr flow appeared. We next performed a protein structure analysis [21–23], by introducing a moving average, S, of local hydrophobic scores around each AA site (Materials and Methods), and expressed the local hydrophobic environment of the root sequences using an AA scale model (Table S1) gives the hydrophobic score of each AA with an orthodox model [24]. We find that the Hoa→Thr flow occurred especially in the hydrophobic region (S>0.6) of mt proteins (the frequency distribution of S-values gives the maximum at S=0.6, whereas the remaining 11 AAs (Rma) except for Hoa and Thr had no flow (Figure 3B and Table 1A). This hydrophobicity dependence of the Hoa→Thr flow was also observed in the sperm whale and elephant lineages (Table 1B and 1C). To investigate this dependence in detail, we estimated the spatial distribution of the Thr residues around the membrane interior using a transmembrane analysis to predict the helical regions of the AA sequences [22,23]. Here, the distribution was estimated in the simian and root sequences. It was measured from the AA coordinates of the membrane surface on the matrix side by splitting an AA sequence into fragments, each of which was embedded only once in the membrane interior. Figure 4 demonstrates that the Thr increase in the lineage leading to the simians took place preferentially in the membrane-spanning (helix) regions. Because the maximum parsimony reconstruction of the root sequence may be biased toward the consensus state [25] and this could produce an artifact that incorrectly suggests an AA composition flow, we conducted an additional analysis with the simian lineage. We defined the root of the simians as the most recent common ancestor of the tree shrew and the primates. Then, the root sequence estimated could be approximately substituted by the real sequence of the tree shrew, because the branch length between the root and the tree shrew is very short, as seen in Figure 1. This substitution was useful to confirm that the estimated root sequence is not biased, and provided a similar result of the Hoa→Thr flow to that given by the estimated root sequence (Tables 1A versus 1D). Table S2 lists the AA positions of mt proteins at which the Hoa sites of the primate root sequence in the hydrophobic region (S>0.6) changed to Thr in chimpanzee or human.

Discrimination between adaptive evolution and mutation bias at the DNA level

By estimating the node sequences at the nucleotide level, we calculated the numbers of nonsynonymous changes at the first and second codon sites, and also the number of synonymous changes at the FD3rd codon sites, between the root sequence and the terminal sequences. A predominant T→C flow (T, thymine; C, cytosine) at the second codon site was inferred in the three lineages of simians, sperm whale, and elephant (Tables 2A–2C). Furthermore, this flow occurred preferentially in the highly hydrophobic regions, and produced the main part (the Ile/Met→Thr flow) of the Hoa→Thr flow. We also note that the T→C flow direction was reversed at the FD3rd codon sites of the sperm whale and elephant lineages, with no hydrophobicity dependence. Conversely, the predominant A→G flow (A, adenine) was inferred at the FD3rd codon sites, without hydrophobicity dependence in the simian and sperm whale lineages. The A→C flow also occurred at the first codon sites and yielded intralouds (the Ile/Met→Leu) among the Hoa, but the increase in Leu residues showed no hydrophobicity dependence, in contrast to the increase in Thr residues (Figure 3B). Consequently, we conclude that the T→C flow was induced by AA adaptive evolution, whereas the A→G C flow was caused by nucleotide mutation bias. This conclusion was supported by using again the tree shrew instead of the primate root (Tables 2A versus 2D), and by estimating the significance of differences between two nucleotide flows in the hydrophobic regions of S<0.6 and S>0.6. The T→C flow for adaptive evolution in the simian lineage showed a significant hydrophobicity dependence with p = 1.123×10⁻³.

Relationship between Thr increase and longevity in primates

We showed that the frequency of Thr residues greatly increased in the highly hydrophobic region (S>0.6) of mt proteins in the
Figure 2. Hoa→Thr flow along the primate lineage. Figures A, B and C represent, respectively, the increased number in Thr, the decreased number in Hoa (relative to the Thr/Hoa numbers of the root sequence) and the evolutionary rate, as functions of the divergence times of primates. The evolutionary rate was traced along the human lineage. The values of the times and rates were estimated using a harmonic mean method which is robust against strong evolutionary rate changes [20]. Here, we used the root and node AA sequences which were obtained in Figure 1.

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Figure 3. AA transition net flows in the three lineages of simians, sperm whale, and elephant, and hydrophobicity dependence.

Figure A represents the net flow of each AA in the three lineages of the primate, the sperm whale and the elephant. The net flow of an AA is the site number of this AA which increased or decreased in terminal sequences, relative to a root sequence. The roots of the primates, the sperm whale and the elephant were defined as the most recent common ancestors of the tree shrew and primates, the sperm whale and hippopotamus, and the elephant and golden mole, respectively (Figure 1). The bottom lines denote the 20 AAs and the nucleotides at the first and second codon sites. Figure B represents the hydrophobicity dependence of the AA net flows in the two regions $S<0.6$ and $S>0.6$ of mt proteins, for the simian lineage (Methods and Materials) and shows that the Hoa $\rightarrow$ Thr flow appears in only the AA sites with $S>0.6$ while the increases of the Leu residues exists independent of the $S$ values because of the directional mutation pressure causing the T $\rightarrow$ C flow. $S$ expresses the local hydrophobic environment of AA sequences and is given by a moving average of local hydrophobic scores around each AA site (Materials and Methods).

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**Table 1.** Hoa→Thr flow in the three lineages of simians, sperm whale, and elephant, and hydrophobicity dependence.

|                | S<0.6 |  | S>0.6 |  | Flow |
|----------------|-------|---|-------|---|------|
|                | In    | Out | In    | Out |      |
| **A. Simians** |       |     |       |     |      |
| Hoa            | 83    | 75  | 33    | 102 | −61  |
| Thr            | 70    | 67  | 76    | 24  | 55   |
| Rma            | 68    | 79  | 39    | 22  | 6    |
| **B. Sperm whale** |     |     |       |     |      |
| Hoa            | 35    | 44  | 19    | 49  | −39  |
| Thr            | 40    | 25  | 45    | 16  | 44   |
| Rma            | 26    | 32  | 13    | 12  | −5   |
| **C. Elephant** |       |     |       |     |      |
| Hoa            | 75    | 66  | 33    | 93  | −51  |
| Thr            | 59    | 60  | 75    | 23  | 51   |
| Rma            | 69    | 76  | 33    | 26  | 0    |
| **D. Simians (tree shrew)** | | | | |      |
| Hoa            | 95    | 104 | 40    | 98  | −66  |
| Thr            | 105   | 90  | 79    | 33  | 61   |
| Rma            | 93    | 100 | 38    | 26  | 5    |

Table gives the incoming (In) and outgoing (Out) AA numbers for the three AA groups of Hoa, Thr, and Rma (the remaining AAs except for Hoa and Thr), between the root and terminal sequences. For example, in the case A, the In and Out numbers denote the average site numbers of Thr residues which appeared and disappeared in the simian terminal sequences, respectively. These numbers were estimated in the two AA site domains, S<0.6 and S>0.6, of the local hydrophobic environment values (Methods and Materials). The flow denotes the differences of the AA numbers between In and Out, which become quite large in the high S-value domain of the AA sites. A, B and C represent the three cases of simians, sperm whale, and elephant, respectively. D gives the case of the real sequence of the tree shrew with a short branch from the root, to confirm that the estimation of the root sequence is not biased. The roots of the simians, the sperm whale and the elephant were defined as the most recent common ancestors of the tree shrew and primates, the sperm whale and hippopotamus, and the elephant and golden mole, respectively (Figure 1).

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**Figure 4.** Thr distribution measured from the membrane surface on the matrix side. Figure estimates the spatial distribution of Thr residues in the 12 mt proteins. The x-values stand for the AA site numbers measured from the inner membrane surface on the matrix side. The inner and outer surface positions (vertical lines) of the helices in each protein AA sequence were inferred from a secondary structure analysis of the mt proteins [21–23]. The black and red circles show the Thr compositional distribution (%) in the root sequence (the most recent common ancestor of the tree shrew and primates) of the primates and that (the averaged value) in the simian terminal sequences, respectively. The blue circles represent the increased fraction (gain) of Thr residues in the simians.

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Adaptive Mt-Thr Increase

Table 2. Nucleotide substitution flows in the three lineages of simians, sperm whale, and elephant.

|                | 1st     | 2nd     | FD3rd   |
|----------------|---------|---------|---------|
| A. Simians     |         |         |         |
| T→C            | 17 (10.7)| 67 (6.61)| 19 (10.9)|
| T→A            | 6 (5.1) | 6 (2.8) | −51 (−28, −23)|
| T→G            | −1 (2.3) | 0 (0.0) | 2 (2.0) | |
| A→C            | 48 (25.23) | 3 (4.1) | 196 (99.97)|
| G→C            | 4 (1.3) | 8 (3.5) | 7 (4.3) | |
| G→A            | 28 (10.18) | 4 (1.3) | −41 (−20, −21)|
| B. Sperm whale |         |         |         |
| T→C            | 0 (−1.1) | 37 (12.25) | −35 (−20, −15)|
| T→A            | −3 (2.5) | 0 (−4.4) | −56 (−25, −31)|
| T→G            | −2 (−3.1)| 0 (1, −1)| 5 (3.2) | |
| A→C            | 27 (14.3)| 3 (4, −1)| 103 (38.65)|
| G→C            | 11 (6.6) | 5 (1.4) | 0 (−1) | |
| G→A            | 9 (5.4) | 2 (1.1) | −50 (−25, −25)|
| C. Elephant    |         |         |         |
| T→C            | 18 (13.5)| 29 (−7.36) | −19 (−7, −12)|
| T→A            | −1 (−3.2)| 7 (4.3) | −37 (−13, −24)|
| T→G            | −6 (−2, −4) | 4 (1.3) | 2 (2.0) | |
| A→C            | 10 (1.9)| −6 (−4, −2) | 21 (21.0)|
| G→C            | −8 (−5, −3)| −3 (−1, −2) | −8 (−6, −2)|
| G→A            | −5 (−16.11)| −2 (−2.0)| −31 (−16, −15)|
| D. Simians (tree shrew) |         |         |         |
| T→C            | 20 (12.8)| 49 (6.63) | 33 (12.21)|
| T→A            | 18 (10.8)| −1 (−7.7) | 2 (−7.9) | |
| T→G            | −8 (−4, −4) | 0 (0.0) | −4 (−4.0)|
| A→C            | 48 (21.27)| −3 (0, −1)| 134 (72.62)|
| G→C            | 14 (7.7)| 13 (4.9) | 23 (12.11)|
| G→A            | 71 (31.40)| 12 (7.5)| 3 (−2) | |

Table gives the nucleotide substitution flows of nonsynonymous changes at the first and second codon sites, and those of synonymous changes at the FD3rd sites. The quantity X→Y denotes the net flow from X to Y. The values u and v in parentheses [u, v] denote the fractions in the two score regions S=0.6 and S=0.6, respectively. A, B, and C represent the three cases of simians, sperm whale, and elephant, respectively. D indicates the case of the real sequence of the tree shrew with a short branch instead of the root sequence of the primates, to confirm that the estimation of the root sequence is not very biased. doi:10.1371/journal.pone.0003343.t002

Table 3. Thr & Cys-MLS correlations in 20 primates and tree shrew.

|                | 12 proteins | 4 proteins |
|----------------|-------------|------------|
|                | AA sites (all) | AA sites (S>0.6) | AA sites (S>0.6) |
|                | LS PGLS | LS PGLS | LS PGLS |
| Thr            | 0.431 | 0.492 | 0.597 | 0.439 | 0.568 | 0.533 |
| (0.051) | (0.025) | (0.004) | (0.047) | (0.007) | (0.013) |
| Cys            | 0.575 | 0.458 | 0.619 | 0.467 | 0.523 | 0.53 |
| (0.006) | (0.038) | (0.002) | (0.034) | (0.016) | (0.013) |
| Thr+Cys        | 0.607 | 0.552 | 0.740 | 0.535 | 0.650 | 0.656 |
| (0.015) | (0.030) | (0.001) | (0.035) | (0.007) | (0.007) |

We estimated the Thr-Log(MLS), Cys-Log(MLS) and Thr/Cys-Log(MLS) correlation coefficients (R) by using the multivariate regression analysis. The result of least square (LS) which does not take account of the tree structure is compared with that of the phylogenetic generalized least-squares (PGLS) which takes account of it (Materials and Methods). The parentheses give the p-values for significance. The correlations were more explicit in the hydrophobic region (S>0.6) of the AA sites in the 4 proteins (ND1, ND2, ND4 and Cytb). doi:10.1371/journal.pone.0003343.t003

Table gives the values of Thr/Cys compositions and Log (MLS) in primates and the accession number of mt DNA sequences used in this analysis.

Discussion

The compositional changes that we have detected in the membrane regions may be of importance to those who are using mitochondrial DNA sequences for phylogenetic studies, because standard models of sequence analysis do not permit or predict these changes. The correlation at the AA level between the Thr composition in the hydrophobic region and the MLS in primates is intriguing, since many other factors to regulate longevity may be considered at a macroscopic level [32] and a recent study reports that the base composition of mtDNA, especially the G or GC content could undergo co-evolution with the mammalian longevity [33].

Schmitz et al. pointed out a compositional nucleotide shift from T→G and A→C in the higher primates, and demonstrated the impact of these changes on the AA composition [13]. Furthermore, Gibson et al. suggested that the increase in Thr residues in 12 mt proteins correlated with the increase in the C content [14]. These authors argued that the variations in the AA compositions among species were caused by nucleotide mutation bias. However, those studies were mainly based on inclusive compositional regression analysis of the terminal sequences. We investigated the nucleotide changes along the evolutionary pathway of the simian lineage, and observed that the T→G flow was dependent on the local hydrophobic environment of the root AA sequence, while the A→C flow was less dependent on it.

A specific increase of Thr residues in the membrane interior must be presumably connected to some mt functions. Recent studies elucidated that membrane proteins undergo large movements during functions [15,16]. A trade-off between mobility and stability of proteins may exist [16]. We note that helix–helix interactions play an essential role in the stabilization of membrane

simians (Figure 3B and Table 1A). We speculate that this threonine increase may be related to life span via an increased stability of these proteins. Recently, Moosmann and Behl noted that Cys composition is negatively correlated with life span in a wide range of animals [26]. They argue that Cys depletion may render mt proteins more resistant to oxidative attack. By adding several taxa recently stored in the NCBI database [http://www.ncbi.nlm.nih.gov/Genbank/] and then calculating the Cys and Thr compositions (%) of primate AA sequences, we characterized the correlations between these AA compositions and the logarithm of maximum life spans (MLS). This was done by using phylogenetic generalized least-squares [27] because it takes into account the effect of tree structure within a multivariate regression framework (Materials and Methods; Table 3). A significant relationship between Thr composition and the logarithm of MLS appeared in primates where a predominant Hoa→Thr flow is observed (Figure 5A). Interestingly, the magnitude of this correlation was compatible with that of the Cys-Log (MLS) negative correlation (Figure 5B). The correlation was stronger in the hydrophobic region (S>0.6) of 4 proteins (ND1, ND2, ND4 and Cytb). Table S3 gives the values of Thr/Cys compositions and Log (MLS) in primates and the accession number of mt DNA sequences used in this analysis.
proteins [16–19] and about half of all helix pairs are formed by classical hydrogen bonds [18,19]. Interestingly, most of these bonds are found in motifs involving medium-polar residues, such as Thr or Ser. Thr/Ser motifs can also drive the association of model transmembrane helices [28]. Thus, Thr/Ser residues tend to face the membrane interior, the location of which is important for helix–helix interactions, and the methyl (CH$_3$) group in Thr is especially suited to the hydrophobic environment of the membrane interior [17]. Consequently, a plausible adaptive explanation for the increase in Thr in the membrane interior is that added Thr reinforced the stability of mt proteins. This Thr increase in adaptive evolution may be an important step toward a solution to the membrane-protein-folding problem [29].

We searched for AA positions at which putative helix-helix interactions could be formed by the change from Hoa to Thr, on the basis of the bovine crystal structure of respiratory complex III (cytochrome bc1 complex; PDB ID: 1QCR reported by Xia et al. [30]). We first looked at the AA position 78 in Cytb which gives Hoa (such as Ile, Leu, Met, and Val) in most of the 62 taxa of Figure 1. Among the 62 taxa, this position has Thr only in gorilla, baboon,
and whether such a flow has a connection to longevity.

Ser flow has occurred in a wider range of evolutionary lineages than in controls [31]. This Ile78Thr replacement in human individuals with extreme longevity may contribute to the stability of the respiratory complex by enhancing the helix-helix interaction within the molecule. The second example is the AA position 180 of Cytb which gives Thr in simians and sperm whale, and Ala in the remaining taxa except for cat with Gly. When Ala180 in helix D of bovine is replaced by Thr, this Thr180 can form a hydrogen bond with the side chain of Ala302 in helix F of bovine Cytb.

Materials and Methods

Supporting Information

Table S1  Hydrophobicity scores of AAs. Our primary structure analysis used a model of Cowan and Whittaker for hydrophobicity
indices of AAs [24]. The * and # symbols denote the hydrophobic AAs (Hoa) defined in this paper and threonine, respectively. Although Trp is hydrophobic, it was not included in Hoa since it did not indicate any flow (Figure 3).

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Table S2 The AA sites of mt proteins which changed from Hoa to Thr. This Table lists the AA sites of proteins which changed from the Hoa in the root sequence of primates to the Thr in the chimpanzee or human sequence. These changes took place in the hydrophobic region of S>0.6. The root was defined as the most recent common ancestor of the tree shrew and the primates.

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Table S3 The Thr/Cys composition and Log (MLS) values in primates. The second and third columns give the common names of primates and the accession numbers of the DNA sequences in the NCBI database, respectively. The fourth and fifth columns give the Thr and Cys compositions (%) in all AA sites of 12 mt proteins, respectively. The sixth and seventh columns give those in the hydrophobic sites (S>0.6), respectively. The last column gives the Log [MLS (years)] values.

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Author Contributions

Analyzed the data: YK NN. Contributed reagents/materials/analysis tools: YK HK MH NN JLT MT. Wrote the paper: YK HK MH JLT MT.