

Response to Comment on “Positive Selection of Tyrosine Loss in Metazoan Evolution”

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Su *et al.* claim guanine-cytosine (GC) content variation can largely explain the observed tyrosine frequency variation, independent of adaptive evolution of cell-signaling complexity. We found that GC content variation, in the absence of selection for amino acid changes, can only maximally account for 38% of the observed tyrosine frequency variation. We also uncovered other mechanisms acting to reduce tyrosine phosphorylation that further support our previous proposal.

The expression of Src tyrosine kinase in the unicellular yeast *Schizosaccharomyces pombe* and *Saccharomyces cerevisiae*, whose genomes encode no tyrosine kinase, is toxic due to deleterious tyrosine phosphorylation (1, 2). This raises the question of how metazoans have evolved to accommodate the numerous tyrosine kinases encoded in their genomes (e.g., 90 tyrosine kinases in humans). We proposed that an observed tyrosine depletion (OTD) in metazoan proteins—in addition to tyrosine phosphatase and spatiotemporal/contextual regulation of tyrosine kinases—helps reduce deleterious tyrosine phosphorylation (3, 4). We further proposed that this accounts, at least in part, for the observed negative correlation of genomic tyrosine frequency ($Freq_{Tyr}$) with cell type number and tyrosine kinase number (Num_{TyrKin}) across different species (3). The mutational forces changing GC content, briefly mentioned in our original paper and highlighted by Su *et al.* (5), might have facilitated the OTD. However, we disagree with Su *et al.* that “biased nucleotide substitutions (A/T → G/C) removed spurious tyrosine phosphorylation sites, a random genomic dynamics independent of the adaptive evolution of cell-signaling complexity” and that GC content variation can explain most of the observed variation in $Freq_{Tyr}$.

Genetic mutations generate phenotypic variation for selection forces to act upon. Hence, the observed high GC content in metazoans does not nullify our proposal. The concern is whether the OTD has been passively driven, in the absence of

selection for amino acid changes, by (i) the mutational forces behind high GC content and (ii) the selection for their effects at the nucleotide level (e.g., transcription and translation) that we collectively termed GC directional force (GC force). We computed a GC-content measure minimally influenced by amino acid selection: GC content at the third position of all four-fold degenerate codons (GC4). We correlated $Freq_{Tyr}$ with GC4 content to quantify how much OTD could be passively driven by GC force on protein-coding regions that can directly affect amino acid changes, in contrast to Su *et al.* who focus on GC content in noncoding regions. There are eight amino acids encoded by four-fold degenerate codons. The GC content at the third position for each of them (GC3) and the computed GC4 showed strong correlation with each other (see Fig. 1A). Thus, we conclude that GC4 content is a robust readout of global GC directional forces minimally influenced by amino acid selection. We also computed GC content at all codon positions (GC_{123}) for comparison.

We found that variation in GC4, GC_{123} , and Num_{TyrKin} can individually account for up to 37.6, 56.2, and 73.4% of variation in $Freq_{Tyr}$, respectively (R^2) (Fig. 1B). Adopting the approach by Su *et al.* to correct for phylogenetic relationship of species analyzed (gradual Brownian model), the statistical significance for the negative correlation of $Freq_{Tyr}$ with GC4, GC_{123} , and Num_{TyrKin} is $P = 8.5 \times 10^{-3}$, 1.6×10^{-3} , and 1.8×10^{-4} (subroutine test in R software), respectively (6). The GC4 content of a subset of species does correlate negatively with $Freq_{Tyr}$ (Fig. 1B, yellow ellipse), but the trend is reversed in several other lineages (Fig. 1B, blue ellipse). In contrast, $Freq_{Tyr}$ covaries more consistently with Num_{TyrKin} across different lineages (Fig. 1B). Performing an orthogonal analysis using a generalized additive linear model (7, 8), which models the effects of multiple parameters on a variable simultaneously, we also observed that $Freq_{Tyr}$ correlates better with Num_{TyrKin} than with GC4 (Table 1). These results invalidate Su *et al.*'s claim that GC content variation alone can explain most

of the observed variation in $Freq_{Tyr}$. As GC directional forces on coding regions directly influence amino acid frequency in the absence of selection forces, we question why Su *et al.* chose to emphasize the correlation of $Freq_{Tyr}$ with GC content variation in noncoding regions while not commenting on the results of GC4 variation in coding regions. As we are studying protein evolution, it is more relevant to focus on protein-coding DNA regions.

Although we previously acknowledged that other mechanisms may have contributed to the OTD, we supported our proposal with empirical data that human proteins with no detectable phosphotyrosines (non-pTyr proteins) have lost considerably more tyrosines than known tyrosine-phosphorylated proteins (pTyr proteins), as compared with their *S. cerevisiae* orthologs (3). Here, we extended this analysis to *S. pombe* and reached the same conclusion (Fig. 2A). Yeasts are compared because they are the known eukaryotes phylogenetically closest to human that lack a dedicated phosphotyrosine signaling system, and we implicitly assume both species are informative of the ancestral tyrosine frequency. Our conclusion is supported by the observation that phenylalanine is not lost preferentially in either protein group (Fig. 2A). This is crucial because phenylalanine and tyrosine are structurally identical except for a phosphorylatable hydroxyl group on tyrosine, and are each encoded by two AT-rich codons (Phe: TTT and TTC; Tyr: TAT and TAC), and thus should be subjected to a similar degree of GC force. Hence, the observed preferential loss of tyrosine is due to the presence of its hydroxyl group and strongly supports the argument that signaling fidelity is a driving force behind OTD.

During evolution, phenylalanine and tyrosine are commonly substituted for each other because of their similar physicochemical properties and encoding codons [see BLOSUM matrices, for example (9)]. We investigated their substitution pattern from yeast to human. As expected, tyrosines in yeast are most frequently substituted with phenylalanine (~35%) in human, whereas phenylalanine and tryptophan in yeast are frequently substituted with tyrosine in human (Fig. 2B). However, we observe that the substitution of phenylalanine and tryptophan in yeast by tyrosine in human is significantly underrepresented in non-pTyr proteins compared with pTyr proteins (Fig. 2B) ($P < 10^{-7}$, Fisher's exact test, one-tailed). This phenomenon is minimally influenced by GC force because only T ↔ A substitution is required to directly switch between phenylalanine and tyrosine. We detected no statistical difference ($P < 0.01$, Fisher's exact test, two-tailed) in the substitution of tyrosine in yeast with phenylalanine and tryptophan in human between the two protein groups. Thus, constrained substitution of phenylalanine and tryptophan with tyrosine is possibly another mechanism contributing to OTD and supports our observation that there is selection pressure to remove tyrosines for signaling fidelity.

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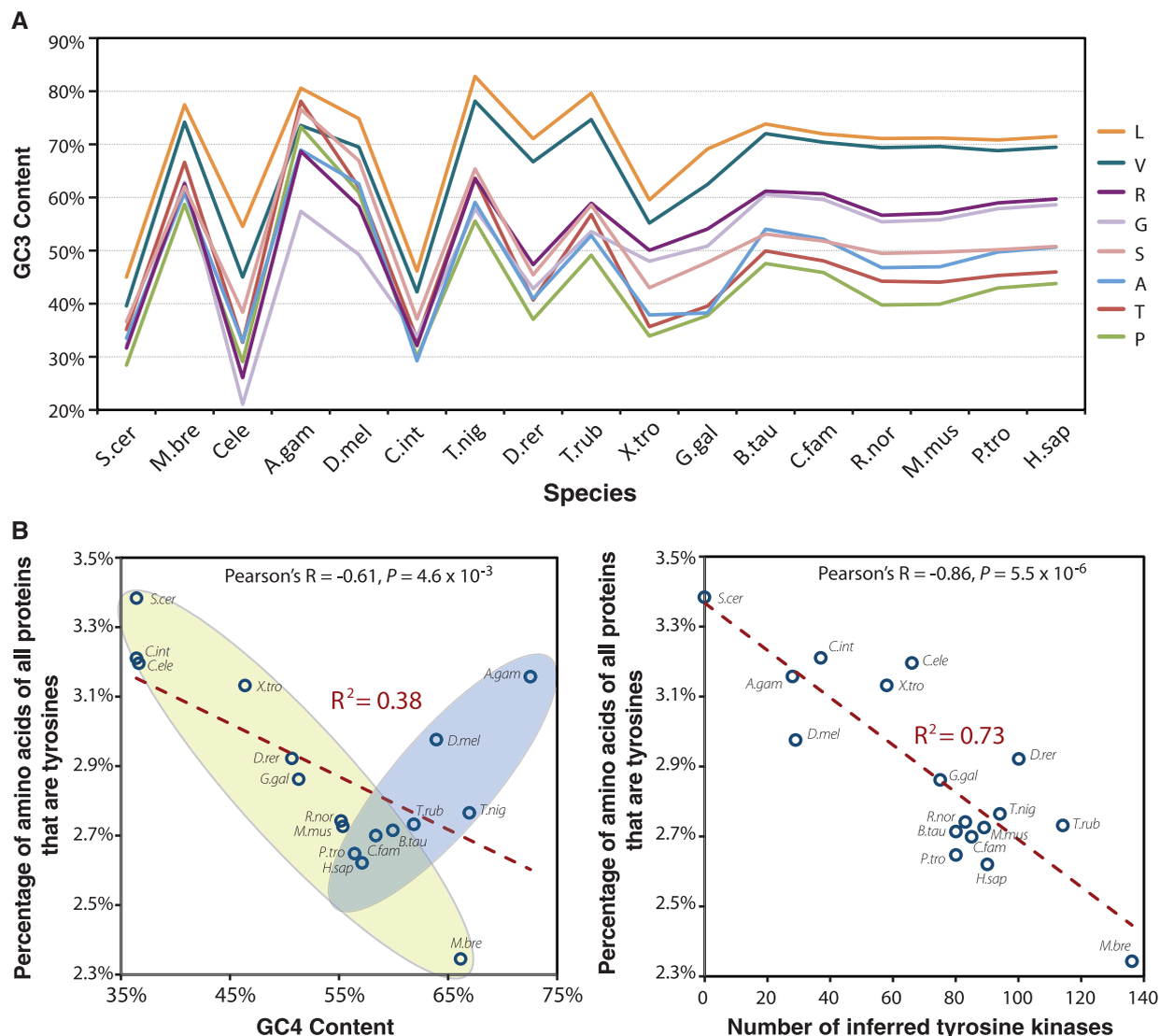


Fig. 1. (A) GC content at the third codon position (GC3) for each set of four-fold degenerate codons in all coding sequences in different species. The four-fold degenerate codon set of each amino acid are GCN (alanine), CGN (arginine), GGN (glycine), CTN (leucine), CCN (proline), TCN (serine), ACN (threonine), and GTN (valine). The species are sorted with decreasing evolutionary distance from human. The GC3 content for these eight amino acids are highly correlated with each other (average Pearson's correlation = 0.85, SD = 0.11). GC3 content for all sets of four-fold degenerate codons are summed to derive the GC4 content for each species. The GC4 content is strongly correlated with GC3 content of these eight amino acids across the species analyzed (average Pearson's correlation = 0.96, SD = 0.036). **(B)** Correlation of genomic tyrosine frequency ($Freq_{Tyr}$) with GC4 content and inferred number of tyrosine kinases (Num_{TyrKin}) in species analyzed. Num_{TyrKin} are inferred as previously described (3) except for *Monosiga brevicollis*, which is based on (14). The low

$Freq_{Tyr}$ and high Num_{TyrKin} observed in *M. brevicollis* is consistent with our proposed evolutionary model (3). The species analyzed are budding yeast (*S. cerevisiae*), choanoflagellate (*M. brevicollis*), worm (*Caenorhabditis elegans*), sea squirt (*Ciona intestinalis*), fly (*Drosophila melanogaster*), mosquito (*Anopheles gambiae*), zebrafish (*Danio rerio*), tetraodon pufferfish (*Tetraodon nigroviridis*), Japanese pufferfish (*Takifugu rubripes*), frog (*Xenopus tropicalis*), chicken (*Gallus gallus*), dog (*Canis familiaris*), cow (*Bos taurus*), mouse (*Mus musculus*), rat (*Rattus norvegicus*), chimpanzee (*Pan troglodytes*), and human (*Homo sapiens*). Looking at the closest pairs of species in five lineages (*R.nori/M.mus*, *P.tro/H.sap*, *C.fam/B.tau*, *T.rub/T.nig*, and *D.mel/A.gam*), a negative relationship between $Freq_{Tyr}$ and Num_{TyrKin} is observed for all five species pairs. Contradicting Su *et al.*'s claim, a positive correlation between $Freq_{Tyr}$ and GC4 is observed for three species pairs (*C.fam/B.tau*, *T.rub/T.nig*, and *D.mel/A.gam*).

Next, for the set of tyrosines that are conserved between human and *S. cerevisiae* as identified from sequence alignments, we applied the NetPhorest algorithm (10) to investigate their propensity to be phosphorylated by human tyrosine kinases that are influenced by residues flanking the tyrosines on primary sequences. In general, we found that tyrosines from human are less phosphorylatable by human tyrosine kinases than

the corresponding tyrosines from *S. cerevisiae* (Fig. 2C) ($P = 5.34 \times 10^{-5}$, Wilcoxon Test). This suggests that there are selection forces favoring mutations flanking tyrosines that reduce tyrosine phosphorylation.

In summary, we contend that our analysis invalidates Su *et al.*'s claims. Furthermore, we uncovered multiple plausible mechanisms acting to reduce tyrosine phosphorylation. Hence, we

suggest that using GC content variation as an Occam's Razor, without considering the physicochemical properties of amino acids in relation to the study of biological systems, can compromise our understanding of similar phenomena. Su *et al.* concur with us that low $Freq_{Tyr}$ permitted the expansion of tyrosine kinases through reducing deleterious phosphotyrosines. Reciprocally, an increased Num_{TyrKin} , other than promoting

Fig. 2. (A) Frequency of tyrosine, phenylalanine, and tryptophan in human tyrosine-phosphorylated (pTyr) proteins, human non-tyrosine-phosphorylated (non-pTyr) proteins, and their protein orthologs in budding yeast (*S. cerevisiae*) and fission yeast (*S. pombe*). Tyrosine, but not phenylalanine and tryptophan, is preferentially depleted in non-pTyr proteins over pTyr proteins. Classification of pTyr and non-pTyr human proteins is based on empirical data [as previously described (3)], which are available in (15). Human protein orthologs in budding yeast and fission yeast are inferred using Inparanoid as described in (3), and only proteins with an inferred one-to-one orthologous relationship are used. Phenylalanine and tyrosine are structurally identical except for the hydroxyl group on tyrosine that renders the residue phosphorylatable. Tryptophan is also included for comparison because it is physico-chemically similar to phenylalanine and tyrosine. Statistical significance indicated is for differences in the distribution between pTyr and Non-pTyr protein sets for observed differences in amino acid frequency of human-yeast orthologous protein pairs, computed with Mann-Whitney test (one-tailed) as previously described (3) using R (16). **(B)** Observed substitution of phenylalanine and tryptophan by tyrosine between human and yeasts. Pairwise sequence alignments between orthologous protein pairs are performed using MAFFT with default parameters (17). We then computed the frequency of mutated phenylalanine and tryptophan being substituted with tyrosine from yeast to human and vice versa. To reduce error due to faulty alignments, residues with fewer than five identical aligned flanking residues out of 10 positions (five on each side) are excluded. We observed that the substitution rate of phenylalanine by tyrosine from yeast to human for pTyr proteins is similar to the rate for human to yeast. However, the rate is significantly lower for non-pTyr from yeast to human. Statistical significance of observed differences is computed with Fisher's exact test (one-tailed) using R. **(C)** Lower phosphorylation propensity of tyrosines in human compared to *S. cerevisiae*. Tyrosines conserved between human and *S. cerevisiae*, as identified from pairwise sequence alignments, are tested for their phosphorylation propensity by human tyrosine kinases using the NetPhorest algorithm (10). Proteins with experimentally observed phospho-tyrosines (pTyr) are tested separately from proteins that have no experimentally observed phospho-tyrosines (non-pTyr). The median probabilities are labeled for each data set, and the indicated *P* values were calculated using the Wilcoxon test.

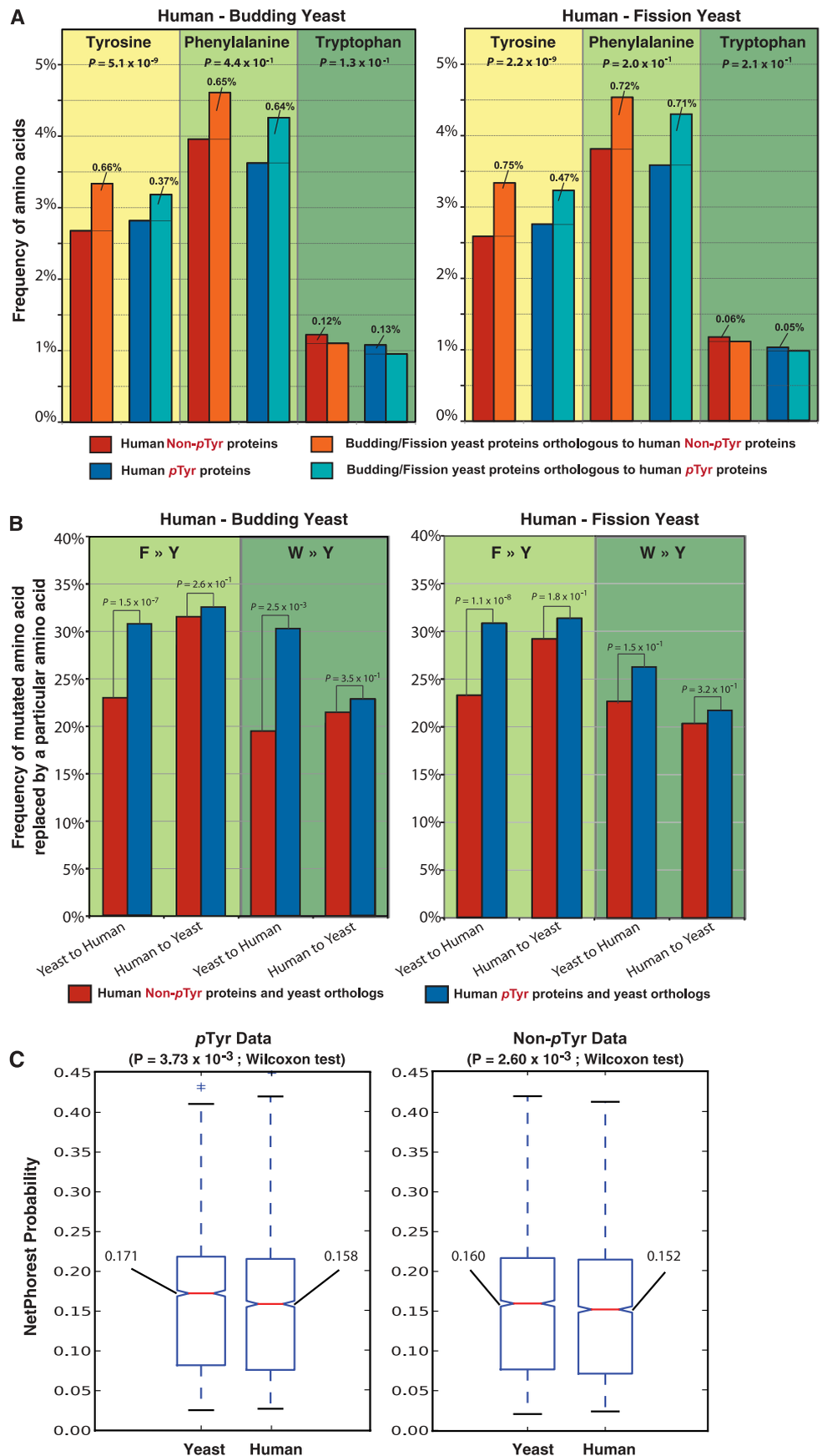


Table 1. Statistical linear models for correlation between tyrosine frequency ($Freq_{Tyr}$) and one or more of the variables GC4 content (GC4) and tyrosine kinase number (Num_{TyrKin}). Results are produced with R (16). AIC, Akaike

information criterion (lower values indicate a better model). Pr, probability. The last column is a visual indicator of the significance of the estimated parameters: *** $P < 0.001$; ** $P < 0.01$; * $P < 0.05$.

Statistical linear model	AIC	R^2	R^2 adjusted	Variables	Estimate	Std. Error	t	Pr ($> t $)	P
$Freq_{Tyr} \sim GC4$	0.80	0.363	0.32	(Intercept)	3.63	0.265	13.71	6.83×10^{-10}	***
				GC4	-0.0147	0.00505	-2.92	1.05×10^{-2}	*
$Freq_{Tyr} \sim Num_{TyrKin}$	-14.13	0.735	0.72	(Intercept)	3.37	0.0840	40.11	$<2 \times 10^{-16}$	***
				Num_{TyrKin}	-0.00675	0.00105	-6.45	1.09×10^{-5}	***
$Freq_{Tyr} \sim GC4 + Num_{TyrKin}$	-19.58	0.829	0.80	(Intercept)	3.71	0.143	26.05	2.92×10^{-13}	***
				GC4	-0.00808	0.00291	-2.78	1.49×10^{-2}	*
				Num_{TyrKin}	-0.00579	0.00094	-6.18	2.39×10^{-5}	***

tyrosine-removing mutations, could also constrain new tyrosine appearance to favor $Freq_{Tyr}$ change unidirectionally. We found it interesting, based on Su *et al.*'s result, that $Freq_{Tyr}$ negatively correlates better with GC content in flanking non-coding regions than with GC4 content and wonder whether CpG dinucleotide site and transcription initiation are attributing factors. If increased GC content does associate with increased gene and protein expression (e.g., abundance and multi-tissue expression) reported in studies (11–13), we speculate that it helps to remove more tyrosines to counter increased encounters of cellular proteins with tyrosine kinases.

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6. To address Su *et al.*'s criticism directly, we applied the gradual Brownian model as described in the supporting online material (SOM) of Su *et al.*, using the species tree inferred by them. The contrast program in PHYLIP software suite computed $R = -0.66$ and $R = -0.83$ for the correlation of $Freq_{Tyr}$ with GC4 and Num_{TyrKin} , respectively. Even without this sophisticated analysis, we already observed that $Freq_{Tyr}$ shows a stronger covariance with Num_{TyrKin} than with GC4 among pairs of closely related species in a phylogenetic-aware manner (see Fig. 1B). The computed contrast values are in the SOM for this paper.
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References

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