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1 Random peptides rich in small and disorder-

² promoting amino acids are less likely to be

3 harmful

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19 Abstract

20	Proteins are the workhorses of the cell, yet they carry great potential for harm via misfolding
21	and aggregation. Despite the dangers, proteins are sometimes born de novo from non-coding
22	DNA. Proteins are more likely to be born from non-coding regions that produce peptides that
23	do little to no harm when translated than from regions that produce harmful peptides. To
24	investigate which newborn proteins are most likely to "first, do no harm", we estimate fitnesses
25	from an experiment that competed Escherichia coli lineages that each expressed a unique
26	random peptide. A variety of peptide metrics significantly predict lineage fitness, but this
27	predictive power stems from simple amino acid frequencies rather than the ordering of amino
28	acids. Amino acids that are smaller and that promote intrinsic structural disorder have more
29	benign fitness effects. We validate that the amino acids that indicate benign effects in random
30	peptides expressed in <i>E. coli</i> also do so in an independent dataset of random N-terminal tags in
31	which it is possible to control for expression level. The same amino acids are also enriched in
32	young animal proteins.

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34 Introduction

35 Proteins are the workhorses of the cell, but they are dangerous. For example, the polypeptide 36 backbone is the key structural feature of amyloids, putting all proteins at risk of forming insoluble aggregates (Chiti and Dobson 2017), and most proteins are expressed at or just 37 38 beyond their solubility limits (Vecchi, et al. 2020). Despite these dangers, new protein-coding 39 genes are nevertheless born de novo from essentially random sequences (McLysaght and 40 Guerzoni 2015; Van Oss and Carvunis 2019; Vakirlis, Carvunis, et al. 2020). To be beneficial 41 enough for *de novo* birth, a random peptide must first do no serious harm, i.e. it must not be detrimental to the basic functioning of a cell. Here we quantify the degree to which, and the 42 43 summary statistics via which, a random peptide's propensity for harm can be predicted.

44 Neme et al. (2017) competed over 2 million Escherichia coli lineages, each containing a plasmid designed to express a unique random peptide, and tracked lineage frequencies over 45 46 four days using deep DNA sequencing. This study has been criticized for providing too little 47 support for the beneficial nature of the top candidates (Weisman and Eddy 2017; Knopp and 48 Andersson 2018). But these criticisms do not detract from using the dataset to identify 49 statistical predictors of serious harm versus relatively benign effect. Neme et al. (2017) used a 50 strong promoter, so evaluation is of tolerance to high expression. Some fitness differences 51 might be due to variation in expression e.g. due to auto-downregulation at the RNA level 52 (Knopp and Andersson 2018) - we will return to this point in the last portion of the Results. 53 Here we pursue analyses based on the hypothesis that the properties of the peptides 54 contribute to variation in fitness among lineages.

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55	Conveniently, computational predictors from peptide sequences alone are available for
56	some properties, such as intrinsic structural disorder (ISD) and aggregation propensity. Because
57	insoluble proteins have been implicated in toxicity and disease (Chiti and Dobson 2017) and
58	peptides with high ISD are less prone to forming insoluble aggregates (Linding, et al. 2004;
59	Angyan, et al. 2012), we hypothesize that highly disordered peptides are least likely to be
60	strongly deleterious. Random sequences with high predicted disorder are well-tolerated in vivo
61	(Tretyachenko, et al. 2017). Existing mouse (Wilson, et al. 2017) and Drosophila (Heames, et al.
62	2020) proteins, which are the product of evolution, are predicted from their amino acid
63	sequences to be more disordered than what would be translated from intergenic controls.
64	Younger protein-coding sequences should be particularly constrained to first do no
65	harm, as they have had little time to evolve more sophisticated harm-avoidance strategies (Foy,
66	et al. 2019). In support of the idea that high ISD is an accessible way to avoid harm, young
67	animal and fungal domains (James, et al. 2021) and genes (Wilson, et al. 2017; Foy, et al. 2019;
68	James, et al. 2021), and novel overprinted viral genes (Willis and Masel 2018) have higher
69	predicted disorder than their older counterparts. Some studies have found that putative de
70	novo protein candidates in Saccharomyces yeasts have lower rather than higher ISD (Carvunis,
71	et al. 2012; Basile, et al. 2017; Vakirlis, et al. 2018), but this could be an artifact of
72	proportionately greater inclusion of non-genes within the younger age classes. When Wilson et
73	al. (2017) reanalyzed Carvunis et al.'s (2012) "proto-genes" of different ages, using more
74	rigorous criteria to exclude non-genes from the data, the direction of the ISD trend was
75	reversed. The same reversal of trend following a quality filter was also found by Vakirlis et al.
76	(2018).

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77	How much do amino acid frequencies matter compared to the order in which those
78	amino acids are arranged? Prior research on young genes has suggested that high predicted ISD
79	in that context is driven primarily by amino acid frequencies, with amino acid order playing a
80	more minor role (Wilson, et al. 2017). Fortunately, the dataset of Neme et al. (2017) is large
81	enough to look at the frequencies of each amino acid as predictors, rather than assume that
82	existing prediction programs such as IUPred (Dosztányi, et al. 2005; Meszaros, et al. 2018) or
83	Tango (Fernandez-Escamilla, et al. 2004; Linding, et al. 2004; Rousseau, et al. 2006) integrate all
84	information about both amino acid frequencies and ordering in the best possible way. We can
85	then test whether such programs have additional ability to predict peptide fitness, above and
86	beyond the influence of amino acid frequencies. In doing so, we can estimate the relative roles
87	of amino acid frequencies versus amino acid ordering in predicting fitness, as well as determine
88	which amino acids have which effects.
89	Here we investigate the degree to which amino acid frequencies and amino acid

90 ordering can predict the fitness effects of random peptides, and if so, which properties are

91 most predictive. We also investigate whether the properties that help random peptides avoid

harm in *E. coli* are also enriched in young eukaryotic proteins. With our work, we hope to

93 further our understanding of how peptides avoid harm.

94 Methods

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95 Data retrieval

96	Neme et al. (2017) performed seven experiments where <i>E. coli</i> lineages, each with a plasmid
97	containing a unique random peptide, were grown and tracked using deep DNA sequencing. We
98	downloaded sequencing counts from Dryad at <u>http://dx.doi.org/10.5061/dryad.6f356</u> , and
99	obtained amino acid and nucleotide sequences directly from Rafik Neme. Experiment 7 was by
100	far the largest with over 4 million reads, more than five times larger than the 2 nd largest
101	experiment and over 1.2 million reads more than all other experiments combined. Experiment
102	7 contained all the peptides that the other six experiments classified as "increasing" or
103	"decreasing," and more. Small datasets from these other six experiments yield limited
104	information because of the need to model changing mean fitness in a population, including not
105	just the tracked lineages but also cells with an empty vector (see Estimating lineage fitness from
106	random peptide sequencing counts section). We therefore chose to restrict our analysis to
107	experiment 7. Experiment 7 consists of the numbers of reads of each random peptide sequence
108	in 5 replicate populations of <i>E. coli</i> at 4 time points. We assume that fitness is identical across
109	replicates, so we summed across all 5 replicates to obtain a total number of reads for each
110	polypeptide at each time point.

Following Neme et al. (2017), we took the 1061 peptides out of over one million that had ≥5 reads across all 5 replicates of experiment 7. Neme et al. (2017) used this cutoff because it is not possible to infer fitness with any reasonable resolution for individual peptides with fewer than five reads. The dramatic nature of this data reduction is unsurprising, firstly because each initial unique peptide was present in only one copy, and secondly because most peptides

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116	are likely deleterious	. We note therefore that	our analyzed subset	of peptides with at least five

- 117 reads are certainly non-lethal, and likely less deleterious than the average random peptide.
- 118 Nonetheless, we achieved enough resolution to distinguish between more and less harmful
- 119 peptides, with remarkably large effect sizes considering the restricted fitness range.
- 120 We further excluded the six peptides that, while meeting the criterion of ≥5 reads, had
- all of those reads at the same timepoint, leaving 1055 peptides for analysis.
- 122

123 Estimating lineage fitness from random peptide sequencing counts

124 The expected number of reads λ_{it} of peptide *i* at times *t*=1,2,3,4 was modeled as:

125
$$\lambda_{it} = N_t p_{i0} \prod_{k=1}^t \frac{\omega_i}{W_{k-1}}$$

where N_t is the observed total number of reads, p_{i0} is the initial frequency of peptide *i* at the beginning of the experiment (prior to the round of selection used to produce the first measured timepoint t = 1), $\frac{\omega_i}{W_t}$ is the fitness of bacteria with peptide *i* at time *t* (i.e. their propensity to contribute to the next time point), and W_k is population mean fitness at time *k*, including bacteria containing empty vectors for which we have no direct count data.

131 The likelihoods of observed peptide counts were estimated from this expectation and 132 two different error models. A Poisson distribution, which captures sampling error alone, was 133 used to generate our initial estimates of p_{io} , ω_i , and W_k (collectively yielding λ_{it}) because it is

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analytically tractable. Under a Poisson error function, the likelihood of observing n_{it} reads of

135 peptide *i* at time *t* is

136
$$f_{Poiss}(n_{it}|\lambda_{it}) = \frac{\lambda_{it}^{n_{it}}e^{-\lambda_{it}}}{n_{it}!}.$$

To also capture variance inflation κ due to PCR amplification, we used a negative binomial
distribution in the Polya form:

139
$$f_{NBP}(n_{it} | \lambda_{i,t}, \kappa) = \left(\frac{\Gamma\left(n_{it} + \frac{\lambda_{i,t}}{\kappa - 1}\right)}{n_{it}! \Gamma\left(\frac{\lambda_{i,t}}{\kappa - 1}\right)}\right) \left(\frac{1}{\kappa}\right)^{\frac{\lambda_{i,t}}{\kappa - 1}} \left(1 - \frac{1}{\kappa}\right)^{n_{it}}$$

140 where $\Gamma(\cdot)$ is the gamma function. We used the initial estimates of p_{io} , ω_i , and W_k to 141 numerically fit the negative binomial model. For the specifics of fitting the Poisson and negative 142 binomial models, see Supporting Information. Weights were calculated, for use in downstream 143 linear models, from this likelihood inference procedure, as the inverse of Fisher information 144 (see Supporting Information).

145 An existing software package for estimating lineage fitness from sequencing counts is 146 Fit-Seq (Li, et al. 2018), which captures the amplification of PCR error through a more 147 sophisticated distribution for the number of reads that is derived in the supplementary information of Levy et al. (2015). However, Fit-Seq assumes that mean fitness is a simple 148 average of all measured lineages' fitness, requiring all individuals to be tagged and measured. 149 150 But Neme et al.'s (2017) experiment included lineages carrying an empty plasmid, i.e. with the 151 selectable marker but no random peptide. Worse, the proportion of cells with an empty vector can be presumed to increase over time. In the absence of a reliable way to directly quantify 152

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153 cells with empty vectors, we instead consider mean population fitness over time to be a set of154 independent parameters to be fitted.

155 Clustering non-independent sequences

156 Upon visual inspection, we found that some peptide sequences were extremely similar, with 157 only one or two amino acid differences; these data points will not contain independent 158 information about the relationship between sequence and fitness. To account for non-159 independence, we clustered peptides by their Hamming distance, and either took only the 160 peptide whose fitness had the highest weight within its cluster, or took weighted means within 161 clusters, or included cluster in our regression models as a random effect term. Single-link 162 clustering with Hamming distance cutoffs of 6 to 29 amino acids all produced an identical set of 163 646 clusters for our 1055 peptides. The largest cluster had 228 random peptides, and the 164 second largest had only 13. The vast majorities of clusters contained only 1 sequence (Dataset 165 S1). A few peptides had mutations in their non-random regions; these mutations were counted 166 in our Hamming distance measurements.

Such similar sequences are highly unlikely to arise by chance if the peptides were truly random; $20^{50} \approx 10^{65}$ peptides are possible, far more than the ~2 × 10⁶ observed. Because we analyze only peptides with at least 5 reads, replicated sequencing error is an unlikely cause. We see the same nearly-identical sequences appearing in every experimental replicate, suggesting either that mutations occurred during Neme et al.'s (2017) initial growth phase, or that the "random" peptides synthesized for the experiment are not entirely random. We note that

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173 construction of the "random" peptide library involved ligations of a smaller set of "seed"

174 sequences, introducing non-randomness at this stage.

175 Predictors of fitness

- 176 All peptides are exactly 65 amino acids long with 50 amino acids of random sequence, so there
- 177 was no need to control for length.

178 GC content

179 Many amino acid sequences mapped to several possible nucleotide sequences, as part of the

180 same problem of mutation or non-random construction discussed above. To calculate one GC

181 content for each random peptide, we calculated a simple average of GC content across all the

182 nucleotide sequences in the dataset that map to the peptide with the largest weight in the

183 cluster.

184 To calculate GC content for the over two million peptides with at least one sequencing 185 read, we took a simple average of the GC content from the random portion of the peptides.

186 Disorder

Protein disorder was measured using IUPred2 (Dosztányi, et al. 2005; Meszaros, et al. 2018) for amino acid sequences, and using disorder propensity (Theillet, et al. 2013) for individual amino acids. IUPred2 returns an ISD score between zero and one for each amino acid in a sequence, with higher scores indicating greater intrinsic disorder. To calculate an ISD score for each random peptide, we took the average of the scores for the whole sequence (i.e. including nonrandom parts). We used a square root transform because it produced a more linear relationship

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193	with fitness than no transform. All measurements referring to ISD or IUPred used IUPred2
194	except Δ ISD, which used the original IUPred program – differences between the two are
195	minimal (Meszaros, et al. 2018).
196	Disorder propensity gives each amino acid a score based on the frequency it is found in
197	disordered proteins relative to ordered proteins (Theillet, et al. 2013). The disorder propensity

- score for a peptide was determined by averaging the disorder propensity scores for the amino
- acids in the random region. When we use the disorder propensity metric, we explicitly refer to
- 200 it as "disorder propensity" and not as "ISD."

201 Aggregation propensity

- Tango (Fernandez-Escamilla, et al. 2004; Linding, et al. 2004; Rousseau, et al. 2006) returns an
- aggregation score for each amino acid in a sequence. At least five sequential amino acids with a
- score greater than or equal to five indicates an aggregation-prone region. We scored peptide
- aggregation propensity as the number of amino acids within regions scored as aggregation-
- 206 prone, including contributions from non-random regions.

207 Solubility

- 208 CamSol (Sormanni, et al. 2015) returns a solubility score for each amino acid in a sequence, as
- 209 well as a simple average of all scores for a sequence, which CamSol calls a "solubility profile."
- 210 We used the solubility profile of the full sequences, including non-random regions.

211 Amino acid frequencies

- 212 We counted frequencies among the 50 amino acids in the random portion of each peptide.
- 213 The values for all the above predictors for each peptide are listed in Dataset S1.

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214 Statistics

- All statistical tests were carried out in R version 3.6.3 (R Core Team 2019), with figures
- 216 generated using "ggplot2" (Wickham 2016). Weighted linear mixed models were implemented
- using the "Imer" function from the "Ime4" package (Bates, et al. 2015), with cluster as a
- 218 random effect. See Supporting Information for details, including justification of a log-transform
- 219 for fitness. When R² values were needed, we instead averaged peptides within the same cluster
- into a combined datapoint, allowing us to avoid the use of random effect term. We calculated
- 221 R² and adjusted R² values using the base R "lm" function. Adjusted R² is a modification of R² to
- 222 penalize additional predictors, and is calculated using the formula:

223
$$R_{adj}^2 = 1 - (1 - R^2) \frac{n-1}{n-p-1}$$

- where *n* are the number of data points and *p* are the number of predictors. Raw P-values are
 reported unless otherwise noted, i.e. without correction for multiple comparisons.
- 226 Data and code availability
- 227 All code and supplemental tables are available on GitHub at
- 228 <u>https://github.com/MaselLab/RandomPeptides</u>. The original Neme et al. (2017) data can be
- found at Dryad <u>http://dx.doi.org/10.5061/dryad.6f356</u>, and the original sequences are available
- at the European Nucleotide Archive (ENA) under the project number PRJEB19640.

231 Results

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232 Estimating the fitness effects of random peptides

Assessing predictors of the fitness effects of random peptides requires those fitness effects to be measured accurately and precisely. Neme et al. (2017) tracked lineage frequencies over four days, and categorized a peptide as increasing or decreasing in frequency by comparing the DNA sequencing counts of day 4 to day 1 using DESeq2 (Love, et al. 2014).

237 We reanalyze the same data, instead using a custom maximum likelihood framework 238 (see Materials & Methods) to quantitatively estimate "fitness" and its associated confidence interval / weight. "Fitness" here refers to allele frequency changes over an entire cycle of 239 240 population growth and dilution, rather than per generation. Our method classifies peptides 241 quantitatively rather than qualitatively. It accounts for the fact that mean population fitness 242 increases over the four days (see Materials and Methods). Our use of all available data within 243 an appropriate maximum likelihood framework should make our method more sensitive and 244 specific for identifying benign vs harmful peptides (see Supplementary Text).

Note that some peptides are pseudoreplicates (see Materials & Methods). There were 646 total clusters, of which there was statistical support for increases in frequency for the highest-weighted peptide in 138 clusters, and for decreases in 488 clusters. Some of our statistics use cluster as a random effect within a linear mixed model. When fixed-effect models are used, such as to generate interpretable R² values, we collapse each cluster into a single pseudo-datapoint with value given by the weighted mean and weight given by the sum of weights.

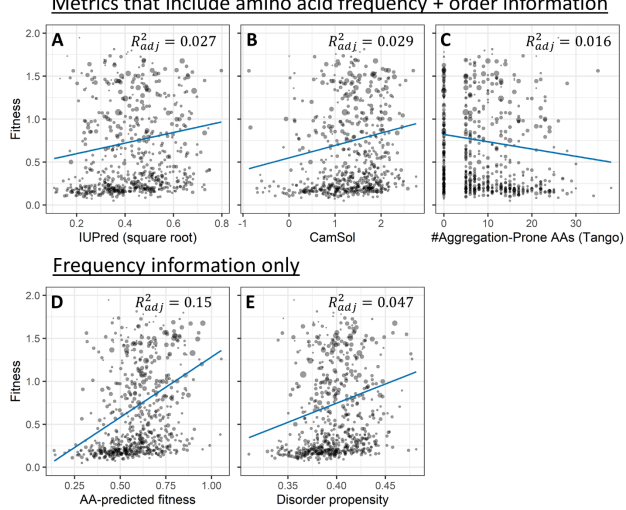
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252 Most predictive power stems from amino acid frequencies rather than amino acid

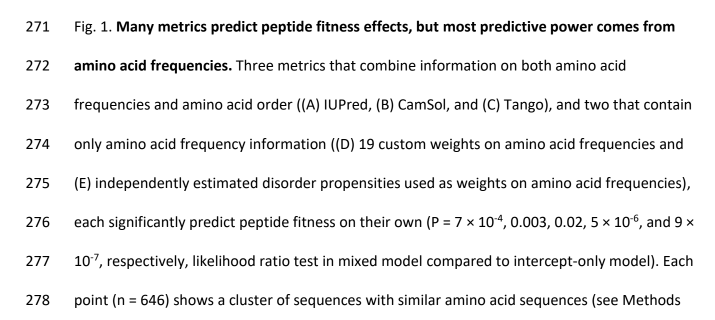
253 order

254	We estimated peptide disorder using several metrics that contain information both about
255	amino acid frequencies and about their order: IUPred as an estimate of intrinsic structural
256	disorder (Dosztányi, et al. 2005; Meszaros, et al. 2018), CamSol as an estimate of water
257	solubility (Sormanni, et al. 2015), and Tango as an estimate of general aggregation propensity
258	(Fernandez-Escamilla, et al. 2004; Linding, et al. 2004; Rousseau, et al. 2006). Fewer than 6% of
259	the random peptides have a predicted transmembrane helix (Dataset S1) from TMHMM (Krogh,
260	et al. 2001), so our choice of these predictors is guided by our assumption that the random
261	peptides are predominantly located in the cytosol. Having a predicted transmembrane helix did
262	not in itself predict random peptide fitness effects (P = 0.2, likelihood ratio test relative to
263	mixed model with only the intercept as a fixed effect). In contrast, each of our cytosol-
264	solubility-inspired metrics significantly predicted random peptide fitness (Fig. 1A – 1C), with
265	effects in the predicted direction (more disorder and more solubility are good, more
266	aggregation propensity is bad). Adjusted R^2 values for IUPred, CamSol, and Tango are 0.027,
267	0.029, 0.016, respectively. Another aggregation predictor, Waltz (Maurer-Stroh, et al. 2010),
268	that specializes in β aggregates, was in the right direction but did not quite meet statistical
269	significance (P = 0.06).

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Metrics that include amino acid frequency + order information

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279	for more details), and the area displayed for each point is proportional to summed weights
280	across that cluster. Blue lines are fixed-effect weighted linear regressions of cluster fitness on
281	the x-axis predictor, where clusters are collapsed to a single pseudo-datapoint by their
282	weighted average and weights are sums within each cluster. Metrics that include both
283	frequency and order information fail to outperform frequency-only based metrics, as shown by
284	regression slopes (blue lines) and adjusted R ² values (top right of each figure panel). Adjusted R ²
285	is calculated as $R_{adj}^2 = 1 - (1 - R^2) \frac{n-1}{n-p-1}$, where <i>n</i> is the number of data points and <i>p</i> is the
286	number of degrees of freedom in the predictor. Note that in part D the predictor (model-
287	predicted fitness) is a composite of 19 degrees of freedom that have all been trained on the
288	dataset, so care should be taken in comparing its blue regression line to that of the other
289	panels, each of which has a predictor with only one degree of freedom – this problem does not
290	apply to comparisons of adjusted R ² values. Seven clusters with fitness greater than 2 are not
291	shown here for ease of visualization; a complete y-axis is shown in supplemental fig. 1. Log-
292	transforming fitness would remove high fitness skew, but creates systematic heteroscedasticity,
293	and so was not done (supplemental fig. 2). The lack of systematic heteroscedasticity can be
294	seen here in the form of similar point size across fitness values.

295

296 Next we asked whether these sophisticated metrics offer additional predictive power 297 beyond mere amino acid frequencies, in the light of prior work on young genes in which little 298 additional predictive power was found (Wilson, et al. 2017). To do this, we fit a model of fitness 299 predicted by amino acid frequencies, measured from counts of each amino acid in each

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300	peptide's random region (Fig. 1D), and compared its performance to predictors that
301	incorporate ordering information (Figs. 1A-C). The amino acid frequency-only model was a
302	significant predictor of fitness (P = 4.5×10^{-6} , likelihood ratio test compared to an intercept-only
303	mixed model). It is also more biologically predictive than other metrics, with adjusted $R^2 = 0.15$
304	(adjusted to account for the number of predictors used) being far greater than the values of
305	0.027, 0.029, and 0.016 found in Figs 1A-1C. Another, non-adjusted, way to look at biological
306	effect size is the far steeper blue line in Fig. 1D than in Figs. 1A-1C. Statistically, when the
307	frequencies of each of the twenty amino acids are used as predictors (Fig. 1D), then IUPred,
308	CamSol, and Tango drop out of the model (P = 0.2, 0.2, and 0.3, respectively, likelihood ratio
309	test in mixed model, see Supplemental Table S1), suggesting that their predictive power in Figs.
310	1A-1C came largely from being metrics of amino acid frequencies. These results are surprising:
311	one might expect sophisticated metrics that incorporate both amino acid frequencies and order
312	information to offer more predictive power and explain a greater range of fitness than simple
313	amino acid frequencies, yet they fail to do so.
314	Our Fig. 1D model using the frequencies of the 20 amino acids involves 19 degrees of
315	freedom, while the other metrics we examine involve only one. This makes it inappropriate to
316	compare the slopes of the blue lines, although adjusted R ² values can still be compared, and the
317	fact that the other metrics drop out of a combined model is also informative. We also
318	investigated a one degree of freedom model of amino acid frequencies, in which relative
319	weights were specified in advance by a disorder propensity metric that assigns each amino acid
320	a score based on how frequently it is found in known disordered versus ordered proteins
321	(Theillet, et al. 2013). Average disorder scores over each peptide's random region significantly

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322	predicted random peptide fitness effects in a linear mixed model (Fig. 1E, $P = 9 \times 10^{-7}$, likelihood
323	ratio test compared to an intercept-only model). The effect size on predicted fitness from the
324	10% to the 90% quantiles of disorder propensity is 0.49 to 0.70, and the adjusted R^2 for the
325	disorder propensity model 0.047. For comparison to other predictors with a single degree of
326	freedom, the largest effect size model that incorporates both amino acid frequency and order
327	information was IUPred with an effect size from 0.51 to 0.69, and the best adjusted R^2 model
328	was CamSol with 0.029. This further suggests that predictive power resides with amino acid
329	frequencies, not order information.
330	To understand whether order information has additional predictive power beyond that
	To understand whether order information has additional predictive power beyond that
331	of amino acid frequencies, we next investigated a metric of ISD that is comprised of only order
331 332	
	of amino acid frequencies, we next investigated a metric of ISD that is comprised of only order
332	of amino acid frequencies, we next investigated a metric of ISD that is comprised of only order information. This can be calculated as the excess IUPred score of the real peptide in comparison

acid frequencies as predictors did not significantly improve the model (P = 0.2). This further

337 supports our conclusion that amino acid ordering plays only a minor role compared to amino

acid frequencies in the fitness effects of the random peptides examined here.

339

340 Small and disorder-promoting amino acids predict benign fitness effects

341 Next we quantify the statistical effect of each of the 20 amino acids on fitness. Naively, we

342 could take the associated slope coefficient in a multiple regression model, which represents the

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343	change in fitness when one amino acid is gained. But in a peptide of fixed length, one amino
344	acid cannot be gained without another amino acid being lost. We therefore instead calculate
345	the marginal fitness effect of each amino acid on fitness (see supplementary text and Table S2,
346	displayed in fig. 2, y-axis), representing the effect of gaining that amino acid and losing a
347	randomly selected alternative.
348	Amino acids with smaller volumes (Tsai, et al. 1999) and higher disorder propensities
349	(Theillet, et al. 2013) tend to have higher marginal fitness effects (fig. 2A and 2B; P = 0.01 for
350	both disorder propensity and volume, likelihood ratio test for dropping either term from a
351	weighted regression of marginal effect on both volume and disorder propensity). Volume and
352	disorder propensity together explain over half the weighted variation in marginal fitness effect
353	(weighted adjusted $R^2 = 0.52$). Other properties of amino acids, such as stickiness (Levy, et al.
354	2012), relative solvent accessibility (Tien, et al. 2013), amino acid cost in <i>E. coli</i> (Akashi and
355	Gojobori 2002), and isoelectric point (Liu, et al. 2004) did not provide significant explanatory
356	power on top of disorder propensity and volume (all P > 0.1, likelihood ratio test).

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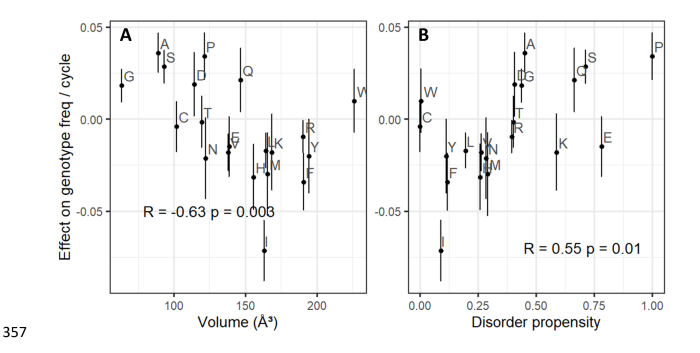


Fig. 2. Amino acids that are small and are associated with disorder promote higher fitness. The y-axis shows each amino acid's marginal effect on fitness, which is the change in fitness when one amino acid of the focal type replaces one randomly chosen amino acid of a different type in a random peptide (see Supporting Information). Error bars are +/- one standard error. Pvalues and correlation coefficients come from weighted Pearson's correlations, where weights for marginal effects are calculated as 1 / s.e. (marginal fitness effect)², and volume and disorder propensity are unweighted.

365

366 Tryptophan is an outlier for amino acid effects on fitness, with a slightly positive effect 367 on fitness despite both its large volume and its underrepresentation in disordered regions (fig. 368 2). Removing tryptophan from a weighted regression model of volume and disorder propensity 369 predicting marginal effect increases the weighted adjusted R² from 0.52 to 0.68. Tryptophan, 370 encoded only by UGG, is nearly 60% more common among peptides with at least 5 sequence

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371	reads than we expect from the 58% GC content of our dataset. Together with the confidence
372	interval for its marginal fitness effect including 1, this provides further evidence that tryptophan
373	is not harmful, making it a distinct outlier, for reasons that are not clear to us.
374	Isoleucine also stands out, as even more harmful than expected by its large size and
375	order propensity. Isoleucine's harmful effects may be exacerbated by its role in amyloid
376	formation. For example, familial amyloid cardiomyopathy is most commonly caused by a valine
377	to isoleucine mutation (Jacobson, et al. 1997; Dubrey, et al. 2015), suggesting that isoleucine
378	has potential to form dangerous amyloids where other hydrophobic amino acids do not.
379	Isoleucine, valine, and leucine are all hydrophobic amino acids with a branched carbon, but only
380	raised isoleucine levels are associated with a higher risk of Alzheimer's disease (Larsson and
381	Markus 2017), further suggesting that isoleucine may be especially prone to amyloid formation.
382	

383 Young animal sequences are enriched for amino acids that increase fitness in

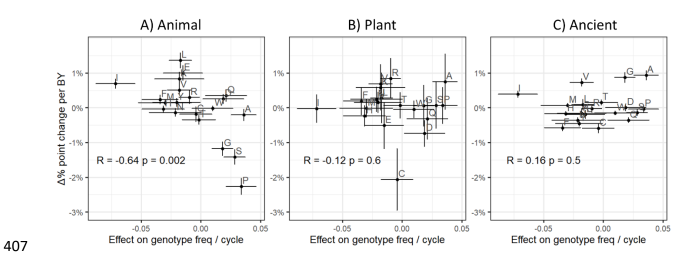
384 random peptides

As discussed in the Introduction, young domains have higher predicted ISD than their older counterparts. One hypothesis to explain this observation is that in order to be successfully born *de novo*, a protein sequence is especially constrained to first do no harm (Wilson, et al. 2017). However, the "phylostratigraphy" approach of assigning ages to genes is contentious. Detecting homologs is more difficult for fast-evolving sequences, which may be erroneously scored as young (Alba and Castresana 2007; Moyers and Zhang 2015, 2016). Disordered proteins tend to be fast evolving (Chen, et al. 2011), suggesting that highly disordered genes

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392	could be misclassified as young because of their fast evolutionary rate. If the amino acid
393	enrichments of higher fitness random peptides match the amino acid enrichments of young
394	genes, this would be evidence that the <i>de novo</i> gene birth process, rather than homology
395	detection bias alone, causes trends in protein properties as a function of apparent gene age.
396	To test this, we took the slopes of amino acid frequencies with protein domain age from
397	James et al. (2021), as quantified across over 400 eukaryotic species. As predicted, amino acids
398	that are good for random peptides are enriched among the youngest animal Pfams (fig. 3A).
399	This prediction was not, however, supported for trends among recent plant domains (fig. 3B)
400	nor among ancient (fig. 3C) domains older than 2.1 billion years. Plant and ancient trends
401	reflect a <i>de novo</i> gene birth process that enriches for the most abundant amino acids in their
402	respective lineages, such as cysteine, rather than for amino acids that promote ISD (James, et
403	al. 2021). It is interesting that we find that ISD still predicts harmlessness in <i>E. coli</i> , even though
404	we do not find evidence it shaped <i>de novo</i> gene birth in its distant ancestors. We also note that
405	ISD does shape recent <i>de novo</i> gene birth in viruses (Willis and Masel 2018).

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408	Fig. 3. Purportedly young animal Pfams are enriched for amino acids that predict high fitness
409	in random peptides. The y-axis represents how the frequency of each amino acid depends on
410	the age of the sequence in billion years (BY), estimated as a linear regression slope for non-
411	transmembrane Pfam domains (James, et al. 2021). Frequency is in number of percentage
412	points, e.g. a difference in glutamic acid content of 5% vs. 6% is a difference of one percentage
413	point. The x-axis shows each amino acid's marginal effect on fitness, which is the change in
414	fitness when one amino acid of the focal type replaces one randomly chosen amino acid of a
415	different type in a random peptide (see Supporting Information). Error bars are +/- one
416	standard error. Fitness effects predict A) animal, but not B) plant, or C) ancient (older than 2.1
417	billion years) Pfam phylostratigraphy slopes. Correlation coefficients and P-values come from
418	weighted Pearson correlations. Note that the P-value for animal phylostratigraphy slopes vs
419	marginal effects survives a conservative Bonferroni correction (P = 0.002 < 0.05/3 = 0.017).
420	

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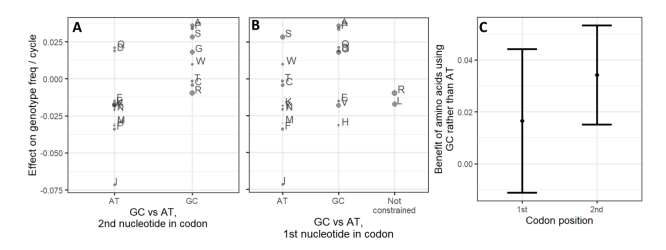
422 Fitness is better predicted by amino acid frequencies than by GC content

Long et al. (2018) proposed that selection acts directly on GC content, perhaps due to the three hydrogen bonds of G-C pairs. Amino acids encoded by Gs and Cs tend to promote higher ISD (Angyan, et al. 2012), making it difficult to distinguish between selection for high GC content and selection for disorder-promoting amino acids. To attempt to distinguish between the two, we compare amino acids that always have G or C to those that always have A or T, at both the first and second nucleotide positions in the codon. If selection were for GC nucleotides, we

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429	would expect GC to predict high marginal amino acid fitness effects at both positions. But if
430	results are dramatically different at the two positions, this would show that it is selection on
431	amino acid content that drives GC as a correlated trait. Results are statistically significant in the
432	predicted direction at the second position (fig. 4A, P = 0.001, weighted Welch's t-test), and in
433	the predicted direction but not statistically significant at the first (fig. 4B, P = 0.2). The effect
434	size of GC content on fitness could not be statistically distinguished between the first and
435	second position (fig. 4C), with wide and hence inconclusive error bars.
436	Linear models are compatible with partially independent contributions of both amino
437	acid frequencies and GC content to harm avoidance. GC content is a statistically significant
438	predictor of fitness by itself (P = 6×10^{-11} , likelihood ratio test for nested fixed-effect models
439	relative to intercept-only model). However, the weighted adjusted R ² of 0.06 for GC content is
440	much lower than the weighted adjusted R^2 of 0.15 (P = 10 ⁻¹⁸) for full amino acid frequency
441	information, suggesting it explains less of the variation than amino acid frequencies. Adding GC
442	content to the amino acid frequencies-only model offers a modest improvement (P = 0.004,
443	weighted adjusted R ² values improves from 0.15 to 0.16), while adding amino acid frequencies
444	to a GC content only model offers a notably larger improvement (P = 10 ⁻¹¹ , weighted adjusted
445	R ² improves from 0.06 to 0.16). These weighted adjusted R ² values suggest that while there
446	may be some direct selection on GC content, the effect of amino acid frequencies appears to be
447	well beyond what can be explained by GC content.

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449 Fig. 4. Amino acids that are constrained to use Gs and Cs tend to have higher marginal effects 450 on fitness than those constrained to use As and Ts. The difference is significant for constraints 451 at the second nucleotide position of a codon (A) (P = 0.001, weighted Welch's t-test), but not at 452 the first (B) (P = 0.2). Point area is proportional to weight, which is calculated as 1 / 2453 s.e.(marginal fitness effect)², as described in Supporting Information. The y-axis is the same as 454 the fig. 2 y-axis and fig. 3 x-axis. C) The mean advantage of amino acids constrained to use GC 455 rather than constrained to use AT is not distinguishable in size between the first and second codon positions. Y-axis gives the difference in the two weighted means of marginal fitness 456 457 effects from A) and B). Error bars represent 95% confidence intervals on the difference 458 between the means (calculated as difference +/- $t_{crit} \times se$), where $t_{crit} \approx 2.1$ is the critical value of the t-statistic with the appropriate degrees of freedom. Weighted Welch's t-test statistic and 459 460 the corresponding standard error of the difference in means were calculated using the 461 "wtd.t.test" function from the "weights" R package, version 1.0.1.

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463 The same amino acids predict benign fitness effects in random N-terminal tags

464 The degree to which benign effects are due to low expression of a random peptide, vs. benign 465 effects of the peptide once expressed, remains unclear. We therefore tested the ability of our 466 amino-acid-frequencies-only model, trained on the data of Neme et al. (2017), to predict residual fitness effects in a dataset that controls for peptide expression level. Goodman et al. 467 468 (2013) tagged the N-prime end of green fluorescent protein (GFP) with 137 different short 469 random sequences (11 amino acids long), allowing random peptide expression level to be 470 measured via fluorescence. Frumkin et al. (2017) measured the fitness effects of these random 471 peptide-tagged GFPs in *E. coli* using FitSeq (Li, et al. 2018). For 89 of them, Frumkin et al. (2017) were able to calculate a "fitness residual" based on the deviation from the fitness expected 472 473 from the level of GFP expression. Note that while this fitness residual controls for expression 474 level, it still contains the cost of inefficient expression in addition to the fitness effect of the 475 peptide itself. Frumkin et al. (2017) found that low fitness residuals were associated with 476 hydrophobic and expensive-to-synthesize amino acids. These findings are consistent with our 477 own estimates of direct peptide effects, as hydrophobic amino acids tend to be order-prone 478 (Linding, et al. 2004; Angyan, et al. 2012), and amino acid volume is highly correlated with synthesis cost in *E. coli* (Pearson's correlation coefficient = 0.85, P = 2×10^{-6} , cost for amino acid 479 480 synthesis in E. coli taken from (Akashi and Gojobori 2002)). Indeed, predicted fitness values for 481 Frumkin et al.'s (2017) N-terminal tags were significantly correlated with their actual fitness 482 residuals (fig. 5). The consistency between our results and the findings of Frumkin et al. (2017), 483 who control for peptide expression level, provides an external validation of our results and

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484 suggests that our findings are unlikely to be due to differences in peptide expression levels

485 alone.

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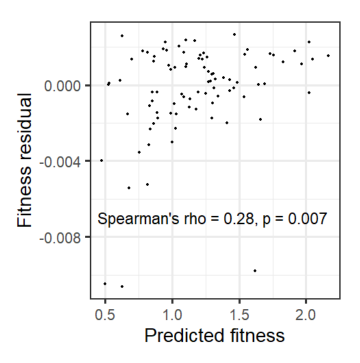


Fig. 5. Fitness predictions trained on the random peptides of Neme et al. (2017) also work for short random tags attached to the N-terminus of GFP. Predicted fitness comes from our amino acid frequencies-only mixed model. "Fitness residuals" of N-terminal tags are from Frumkin et al. (2017), and represent the difference between the fitness of the construct and the expected fitness from expression level. *n* = 89.

492 Discussion

- 493 We found that, while many metrics of peptide properties have some ability to predict the
- 494 fitness effects of random peptides expressed in *E. coli*, most predictive power stems from
- 495 amino acid frequencies. Simply knowing how many of which amino acids are present in these

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496	random peptides can account for 15% of the variance in fitness among lineages, and adding
497	more predictors to account for amino acid order fails to add more predictive power. This
498	indicates both the success of our statistical method for minimizing the noise in our fitness
499	estimates, and that mere amino acid frequencies without amino acid order can be informative
500	of peptide fitness effects. Amino acids that are small and promote disorder predict high fitness
501	in <i>E. coli</i> , and align with those that are enriched in young protein domains in animals.

502 Most studies of random peptides have focused on finding peptides that have specific 503 binding or function (e.g. Kaiser, et al. 1987; Keefe and Szostak 2001; Frulloni, et al. 2009). Some 504 were motivated as proof-of-concept that random peptides can exhibit properties of native 505 proteins, such as folding (Davidson and Sauer 1994; Chiarabelli, et al. 2006; LaBean, et al. 2011) 506 and being soluble (Prijambada, et al. 1996). Others focus on how to increase the percentage of 507 native-like random peptides, e.g. by showing that more hydrophilic random peptide libraries 508 have a higher percentage of stable and soluble peptides (Davidson, et al. 1995). Our work has a 509 different intent, identifying properties that make a peptide less likely to be harmful. Neme et 510 al.'s (2017) experiment was suitable for this purpose because it used a large library of peptides 511 with diverse properties, competed lineages growing under permissive conditions, and 512 measured relative growth rates (i.e. fitness). In contrast, a study design such as that of Knopp et 513 al. (2019), who selected random peptides that rescue viability in the presence of antibiotics, is 514 less suitable for our purposes because so few peptides, including harm-avoiding peptides, are 515 viable. Neme et al.'s (Neme, et al. 2017) study was also convenient because all peptides were 516 the same length – 65 amino acids with 50 amino acids of random sequence – allowing us to 517 neglect length in our analysis.

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518	Having a higher proportion of random peptides do no harm is expected to increase the
519	success rate of future screens for peptide with specific properties. Nucleotide sequences with
520	high %GC content tend to encode peptides with more benign fitness effects, suggesting that
521	higher %GC should be used in future random peptide libraries. However, very high GC content
522	will yield low complexity sequences, which our predictor has not been trained on. The marginal
523	fitness effects of each amino acid might be different in this very different context.
524	While the library used by Neme et al. (2017) was designed to have equal frequencies of
525	each nucleotide in the random region, and thus 50% GC content, the over two million random
526	peptides that had at least one sequencing read had a GC content of ~59% in their random
527	portion. The mean GC content of the peptide clusters we analyzed (see Materials and Methods)
528	was similar, at ~58%, with higher fitness peptides within this group having still higher %GC, as
529	discussed in the Results. The enrichment from 50% GC to ~59% GC might be because many
530	lower GC content sequences were so harmful that lineages that carried them went extinct prior
531	to detection via sequencing. Note that it might also reflect a bias toward GC in sequencing
532	methods (Benjamini and Speed 2012; Choudhari and Grigoriev 2017) – a bias that affects all

time points equally and so should not affect our fitness estimates.

Long et al. (2018) proposed that there is direct selection for high GC content, as evidenced in part by a preference for amino acids with G or C at the second position of codons, in excess of that predicted from mutation accumulation experiments. Our findings cannot exclude this hypothesis, but show stronger selection on amino acid frequencies, selection that is capable of driving increased GC content in coding regions as a correlated trait. In intergenic regions, elevated %GC is likely driven mostly by GC-biased gene conversion. However, elevated

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540	GC content could also be due, at least in part, to selection on peptides from non-coding regions
541	translated by error (Rajon and Masel 2011; Wilson and Masel 2011). Selection on translation
542	errors is for example strong enough to shape non-coding sequences beyond stop codons in
543	Saccharomyces cerevisiae (Kosinski and Masel 2020).
544	Fitness effects in Neme et al. (2017) might not be directly caused by peptide properties
545	alone but instead by the effect of both nucleotide and peptide properties on expression (Knopp
546	and Andersson 2018), with lower expression being less harmful. For example, auto-
547	downregulation at the mRNA level can cause significant difference in expression among
548	peptides, despite identical promoters. However, the properties we find to be predictive, such as
549	disorder and amino acid size, are not a priori related to auto-downregulation of mRNA in wild-
550	type <i>E. coli</i> , making the latter an unlikely explanation for our findings.
550 551	type <i>E. coli</i> , making the latter an unlikely explanation for our findings. While driven by amino acid frequencies, our findings are still consistent with the
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551 552 553 554	While driven by amino acid frequencies, our findings are still consistent with the hypothesis that peptides with low structural disorder tend to be harmful. Disorder-promoting amino acids may help a peptide remain soluble even if unfolded. Small amino acids also tend to be benign, perhaps because they are hydrophobic enough to promote some amount of folding
551 552 553 554 555	While driven by amino acid frequencies, our findings are still consistent with the hypothesis that peptides with low structural disorder tend to be harmful. Disorder-promoting amino acids may help a peptide remain soluble even if unfolded. Small amino acids also tend to be benign, perhaps because they are hydrophobic enough to promote some amount of folding but flexible enough to avoid too much hydrophobic residue exposure.
551 552 553 554 555 556	While driven by amino acid frequencies, our findings are still consistent with the hypothesis that peptides with low structural disorder tend to be harmful. Disorder-promoting amino acids may help a peptide remain soluble even if unfolded. Small amino acids also tend to be benign, perhaps because they are hydrophobic enough to promote some amount of folding but flexible enough to avoid too much hydrophobic residue exposure. Our findings suggest that the easiest way to avoid harm is through disorder and small

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560	transmembrane proteins, which need to be lipid soluble, presumably requiring different harm-
561	avoidance strategies than peptides that are located in the cytosol.

562 The correlation between the extent to which an amino acid is enriched in young animal 563 protein domains and its marginal fitness effect in random peptides in E. coli is intriguing, and 564 consistent with a body of literature that *de novo* gene birth favors protein disorder. What is 565 more, our ability to externally validate animal phylostratigraphy slopes against random 566 peptides in E. coli provides additional support that these slopes represent more than mere bias, 567 in contrast to suggests that all patterns are due to homology detection bias (Alba and Castresana 2007; Moyers and Zhang 2015, 2016). That is, if phylostratigraphy trends were due 568 to an artifact such as homology detection bias, such an artifact would be unlikely to bias our 569 570 random peptide analysis in the same direction. 571 Plants have different trends in amino acid frequencies as a function of sequence age

than animals do, with young genes seeming to prefer readily available amino acids, rather than amino acids that promote ISD (James, et al. 2021). This could be because: 1) plants are less susceptible to harm from random peptides, 2) other properties, such as amino acid availability, drive the emergence of *de novo* genes in plants, or 3) the plant data lack the resolution needed to identify a correlation with the properties studied here. We do not have the ability to differentiate between these three possibilities here.

578 Nevertheless, our finding of consistency between what is benign in *E. coli* and what is 579 benign in animals suggests the possibility of a deep concordance in what makes a peptide 580 harmful between two apparently disparate branches of life. The forces that drive protein birth

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581	therefore appear to share a key similarity between bacteria and Animalia. Monod once
582	suggested that what is true in <i>E. coli</i> must also be true in elephants; our work suggests that this
583	may apply to the properties that tend to make peptides less harmful. To modify Monod's
584	famous quote, what is harmful in <i>E. coli</i> is also harmful in elephants, but not necessarily in
585	eucalyptus.

586	A major idea in our understanding of proteins is that form – that is, the fold that is
587	determined by the exact sequence of amino acids – determines function and thus fitness.
588	However, for these random peptides in <i>E. coli</i> , the amino acid content but not the sequence in
589	which they occur is the main determinant of benign vs harmful effects. Random peptides likely
590	exist as a diverse ensemble of structural states, but the same is increasingly acknowledged to
591	be true of functional proteins. While the ordering of amino acids in functional proteins no
592	doubt plays a role, perhaps mere amino acid frequencies are also more important than once
593	thought in this context too, especially in structurally disordered protein regions.

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