

1 **Defining the functional significance of intergenic transcribed regions based on**
2 **heterogeneous features of phenotype genes and pseudogenes**

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28 **ABSTRACT**

29 With advances in transcript profiling, the presence of transcriptional activities in intergenic
30 regions has been well established in multiple model systems. However, whether intergenic
31 expression reflects transcriptional noise or the activity of novel genes remains unclear. We
32 identified intergenic transcribed regions (ITRs) in 15 diverse flowering plant species and found
33 that the amount of intergenic expression correlates with genome size, a pattern that could be
34 expected if intergenic expression is largely nonfunctional. To further assess the functionality of
35 ITRs, we first built machine learning classifiers using *Arabidopsis thaliana* as a model that can
36 accurately distinguish functional sequences (phenotype genes) and nonfunctional ones
37 (pseudogenes and unexpressed intergenic regions) by integrating 93 biochemical, evolutionary,
38 and sequence-structure features. Next, by applying the models genome-wide, we found that
39 4,427 ITRs (38%) and 796 annotated ncRNAs (44%) had features significantly similar to
40 benchmark protein-coding or RNA genes and thus were likely parts of functional genes.
41 However, ~60% of ITRs and ncRNAs were more similar to nonfunctional sequences and should
42 be considered transcriptional noise unless falsified with experiments. The predictive framework
43 established here provides not only a comprehensive look at how functional, genic sequences are
44 distinct from likely nonfunctional ones, but also a new way to differentiate novel genes from
45 genomic regions with noisy transcriptional activities.

46

47 INTRODUCTION

48 Advances in sequencing technology have helped to identify pervasive transcription in intergenic
49 regions with no annotated genes. These intergenic transcripts have been found in metazoa and
50 fungi, including *Homo sapiens* (human; ENCODE Project Consortium 2012), *Drosophila*
51 *melanogaster* (Brown et al. 2014), *Caenorhabditis elegans* (Boeck et al. 2016), and
52 *Saccharomyces cerevisiae* (Nagalakshmi et al. 2008). In plants, ~7,000 and ~15,000 intergenic
53 transcripts have also been reported in *Arabidopsis thaliana* (Yamada et al. 2003; Stolc et al.
54 2005; Moghe et al. 2013; Krishnakumar et al. 2015) and *Oryza sativa* (Nobuta et al. 2007),
55 respectively. The presence of intergenic transcripts indicates that there may be additional genes
56 in genomes that have escaped gene finding efforts thus far. Knowledge of the complete suite of
57 functional elements present in a genome is an important goal for large-scale functional genomics
58 efforts and the quest to connect genotype to phenotype. Thus the identification of functional
59 intergenic transcribed regions (ITRs) represents a fundamental task that is critical to our
60 understanding of the gene space in a genome.

61 Loss-of-function study represents the gold standard by which the functional significance
62 of genomic regions, including ITRs, can be confirmed (Niu and Jiang 2013). In *Mus musculus*
63 (mouse), at least 25 ITRs with loss-of-function mutant phenotypes have been identified
64 (Sauvageau et al. 2013; Lai et al. 2015), indicating that they are *bona fide* genes. In addition,
65 loss-of-function mutants have been used to confirm ITR functionality in mouse embryonic stem
66 cell proliferation (Ivanova et al. 2006; Guttman et al. 2009) and male reproductive development
67 (Heinen et al. 2009), as well as brain and eye development in *Danio rerio* (Ulitsky et al. 2011).
68 In human, 162 long intergenic non-coding RNAs harbor phenotype-associated SNPs, suggesting
69 that these expressed intergenic regions may be functional (Ning et al. 2013). In addition to
70 intergenic expression, most model organisms feature an abundance of annotated non-coding
71 RNA (ncRNA) sequences (Zhao et al. 2016), which are mostly identified through the presence of
72 expression occurring outside of annotated genes. Thus, the only difference between ITRs and
73 most ncRNA sequences is whether or not they have been annotated. Similar to the ITR examples
74 above, a small number of ncRNAs have been confirmed as functional through loss-of-function
75 experimental characterization, including but not limited to *Xist* in mouse (Penny et al. 1996;
76 Marahrens et al. 1997), *Malat1* in human (Bernard et al. 2010), *bereft* in *D. melanogaster*
77 (Hardiman et al. 2002), and *At4* in *A. thaliana* (Shin et al. 2006). However, despite the presence

78 of a few notable examples, the number of ITRs and ncRNAs with well-established functions is
79 dwarfed by those with no known function.

80 While some ITRs and ncRNAs are likely novel genes, intergenic transcription may also
81 be the byproduct of noisy expression that can occur due to nonspecific landing of RNA
82 Polymerase II (RNA Pol II) or spurious regulatory signals that drive expression in random
83 genomic regions (Struhl 2007). Thus, whether an intergenic transcript is considered functional
84 cannot depend solely on the fact that it is expressed. In addition to being biochemically active,
85 the genomic region must be under selection. This line of logic has revived the classical ideas on
86 differentiating “causal role” and “selected effect” functionality (Doolittle et al. 2014). A “causal
87 role” definition requires a definable activity to consider a genomic region as functional
88 (Cummins 1975; Amundson and Lauder 1994), which is adopted by the ENCODE Consortium
89 (2012) to classify ~80% of the human genome as having biochemical functions. This finding has
90 been used as evidence to disprove the presence of junk DNA that is not under natural selection
91 (see Eddy 2013). This has drawn considerable critique because biochemical activity itself is not
92 an indication of selection (Graur et al., 2013; Niu and Jiang, 2013). Instead, selected effect
93 functionality is advocated to be a more suitable definition for a genomic region with discernible
94 activity (Amundson and Lauder 1994; Graur et al., 2013; Doolittle et al. 2014). Under the
95 selected effect functionality definition, ITRs and most annotated ncRNA genes remain
96 functionally ambiguous.

97 Functional ITRs represent genic sequences that have not been identified with
98 conventional gene finding programs. Such programs incorporate sequence characteristics,
99 transcriptional evidence, and conservation information to define genic regions that are expected
100 to be functional. Thus, genes that lack the features typically associated with genic regions remain
101 unidentified. Due to the debate on the definitions of function post-ENCODE, Kellis et al. (2014)
102 suggested that evolutionary, biochemical, and genetic evidences provide complementary
103 information to define functional genomic regions. Integration of chromatin accessibility,
104 transcriptome, and conservation evidence was successful in identifying regions in the human
105 genome that are under selection (Gulko et al. 2014). Moreover, a comprehensive integration of
106 biochemical, evolutionary, and genetic evidence resulted in highly accurate identification of
107 human disease genes and pseudogenes (Tsai et al. 2017). However, it is not known if such
108 predictions are possible or if the features that define functional genomic regions in human are

109 applicable in other species. In plants, even though many biochemical signatures are known to be
110 associated with genic regions, these signatures have not been incorporated to assist in identifying
111 the functional genomic regions.

112 To investigate the functionality of intergenic transcription, we first identified ITRs in 15
113 flowering plant species with 17-fold genome size differences and evaluated the relationship
114 between the prevalence of intergenic expression and genome size. Next, we determined whether
115 93 evolutionary, biochemical, and sequence-structure features could distinguish functional
116 sequences (phenotype genes) and nonfunctional ones (pseudogenes) using *A. thaliana* as a
117 model. We then jointly considered all 93 features to establish functional gene prediction models
118 using machine learning methods. Finally, we applied the models to ITRs and annotated ncRNAs
119 to determine whether these functionally ambiguous sequences are more similar to known
120 functional or likely nonfunctional sequences.

121 **RESULTS & DISCUSSION**

122 **Relationship between genome size and intergenic expression indicates that intergenic** 123 **transcripts may generally be nonfunctional**

124 Transcription of an unannotated, intergenic region could be due to nonfunctional transcriptional
125 noise or the activity of a novel gene. If noisy transcription occurs due to random landing of RNA
126 Pol II or spurious regulatory signals, a naïve expectation is that, as genome size increases, the
127 amount of intergenic expression would increase accordingly. By contrast, we expect that the
128 extent of expression for genic sequences will not be significantly correlated with genome size
129 because larger plant genomes do not necessarily have more genes ($r^2=0.01$; $p=0.56$; see
130 Methods). Thus, to gauge if intergenic transcribed regions (ITRs) generally behave more like
131 what we expect of noisy or genic transcription, we assessed the correlation between genome size
132 and the amount of intergenic expression occurring within 15 flowering plant species.

133 We first identified genic and intergenic transcribed regions using leaf transcriptome data
134 from 15 flowering plants with 17-fold differences in genome size (Supplemental Table 1).
135 Identical numbers of RNA-sequencing (RNA-seq) reads (30 million) and the same mapping
136 procedures were used in all species to facilitate cross-species comparisons (see Methods).
137 Transcribed regions were considered as ITRs if they did not overlap with any gene annotation
138 and had no significant translated sequence similarity to plant protein sequences. As expected, the

139 amount of expression originating from annotated genic regions had no significant correlation
140 with genomes size ($r^2=0.03$; $p=0.53$; **Fig. 1A**). In contrast, the amount of intergenic expression
141 occurring was significantly and positively correlated ($r^2=0.30$; $p=0.04$; **Fig. 1B**). Because more
142 intergenic expression is occurring in species with more genome space, this is consistent with the
143 interpretation that a significant proportion of intergenic expression represents transcriptional
144 noise. However, the correlation between genome size and intergenic expression explained ~30%
145 of the variation (**Fig. 1B**), suggesting that other factors also affect ITR content, including the
146 possibility that some ITRs are truly functional, novel genes. To further evaluate the functionality
147 of intergenic transcripts, we next identified the biochemical and evolutionary features of
148 functional genic regions and tested whether intergenic transcripts in *A. thaliana* were more
149 similar to functional or nonfunctional sequences.

150 **Expression, conservation, and epigenetic features are significantly distinct between** 151 **benchmark functional and nonfunctional genomic sequences**

152 To determine whether intergenic transcripts resemble functional sequences, we first asked what
153 features allow benchmark functional and nonfunctional genomic regions to be distinguished. For
154 benchmark functional sequences, we used genes with visible loss-of-function phenotypes when
155 mutated (referred to as phenotype genes, $n=1,876$; see Methods). These phenotype genes were
156 considered functional based on the selected effect functionality criterion (Neander 1991) because
157 their mutations have significant growth and/or developmental impact and likely contribute to
158 reduced fitness. For benchmark nonfunctional genomic regions, we utilized pseudogene
159 sequences ($n=761$; see Methods). These pseudogenes exhibit sequence similarity to known
160 genes, but harbor disabling mutations, including frame shifts and/or in-frame stop codons, that
161 result in the production of presumably nonfunctional protein products. Considering that only 2%
162 of pseudogenes are maintained over 90 million years of divergence between human and mouse
163 (Svensson et al. 2006), it is expected that the majority of pseudogenes are no longer under
164 selection (Li et al. 1981).

165 We evaluated 93 gene or gene product features for their ability to distinguish between
166 phenotype genes and pseudogenes. These features were grouped into seven categories, including
167 chromatin accessibility, DNA methylation, histone 3 (H3) marks, sequence conservation,
168 sequence-structure, transcription factor (TF) binding, and transcription activity. Feature values

169 (Supplemental Table 2) were calculated for a randomly-selected 500 base pair (bp) window
170 inside a phenotype gene or pseudogene. We used Area Under the Curve - Receiver Operating
171 Characteristic (AUC-ROC) as a metric to measure how well a feature distinguishes between
172 phenotype genes and pseudogenes. AUC-ROC values range between 0.5 (random guessing) and
173 1 (perfect separation of functional and nonfunctional sequences), with AUC-ROC values of 0.7,
174 0.8, and 0.9 considered fair, good, and excellent performance, respectively. Among the seven
175 feature categories, transcription activity features were highly informative (median AUC-
176 ROC=0.88; **Fig. 2A**). Sequence conservation, DNA methylation, TF binding, and H3 mark
177 features were also fairly distinct between phenotype genes and pseudogenes (median AUC-ROC
178 ~ 0.7 for each category; **Fig. 2B-E**). By contrast, chromatin accessibility and sequence-structure
179 features were largely uninformative (median AUC-ROC=0.51 and 0.55, respectively; **Fig.**
180 **2F,G**). The poor performance of chromatin accessibility features is likely because the DNase I
181 hypersensitive site (DHS) datasets were sparse, as only 2-6% of phenotype gene and pseudogene
182 sequences overlapped a DHS peak. Further, median nucleosome occupancy of phenotype genes
183 (median normalized nucleosome occupancy = 1.22) is only slightly lower than that of
184 pseudogenes (median = 1.31; Mann Whitney U test, $p < 2e-4$). For sequence-structure features
185 based on dinucleotide structures (see Methods), we found that poor performance was likely due
186 to phenotype genes and pseudogenes sharing similar dinucleotide sequence compositions
187 ($r^2=0.99$, $p<3e-16$).

188 **Error rates for functional region predictions are high when only single features are** 189 **considered**

190 Within each feature category, there was a wide range of performance between features (**Fig. 2**,
191 Supplemental Table 3) and there were clear biological or technical explanations for features that
192 perform poorly. For the transcription activity category, 17 out of 24 features had an AUC-ROC
193 performance >0.8, including the best-performing feature, expression breadth (AUC-ROC=0.95;
194 **Fig. 2A**). However, five transcription activity-related features performed poorly (AUC-
195 ROC<0.65), including the presence of expression (transcript) evidence (AUC-ROC=0.58; **Fig.**
196 **2A**). This is because 80% of pseudogenes were considered expressed in ≥ 1 of 51 RNA-seq
197 datasets, demonstrating that presence of transcripts should not be used by itself as evidence of
198 functionality. For the sequence conservation category, maximum and average phastCons

199 conservation scores were highly distinct between phenotype genes and pseudogenes (AUC-
200 ROC=0.83 and 0.82, respectively; **Fig. 2B**). On the other hand, identity to best matching
201 nucleotide sequences found in *Brassicaceae* and algal species were not informative (AUC-
202 ROC=0.55 and 0.51, respectively; **Fig. 2B**). This was because 99.8% and 95% of phenotype
203 genes and pseudogenes, respectively, had a potentially homologous sequence within the
204 *Brassicaceae* family, and only 3% and 1%, respectively, in algal species. Thus, *Brassicaceae*
205 genomes were too similar and algal genomes too dissimilar to *A. thaliana* to provide meaningful
206 information. H3 mark features also displayed high variability. The most informative H3 mark
207 features were based on the number and coverage of activation-related marks (AUC-ROC=0.87
208 and 0.85, respectively; **Fig. 2E**), consistent with the notion that histone marks are often jointly
209 associated with active genomic sequences to provide a robust regulatory signal (Schreiber and
210 Bernstein 2002; Wang et al. 2008). By comparison, the coverage and intensity of H3 lysine 27
211 trimethylation (H3K27me3) and H3 threonine 3 phosphorylation (H3T3ph) were largely
212 indistinct between phenotype genes and pseudogenes (AUC-ROC range: 0.55-0.59; **Fig. 2E**).

213 Despite this high variability in performance, some features and feature categories had
214 high AUC-ROCs, suggesting that these features may individually provide sufficient information
215 for distinguishing between functional and nonfunctional genomic regions. To assess this
216 possibility, we next evaluated the error rates of function predictions based on single features. We
217 first considered expression breadth of a sequence, the best predicting feature of functionality.
218 Despite high AUC-ROC (0.95), the false positive rate (FPR; % of pseudogenes predicted as
219 phenotype genes) was 21% when only expression breadth was used, while the false negative rate
220 (FNR; % of phenotype genes predicted as pseudogenes) was 4%. Similarly, the best-performing
221 H3 mark- and sequence conservation-related features had FPRs of 26% and 32%, respectively,
222 and also incorrectly classified at least 10% of phenotype genes as pseudogenes. Thus, error rates
223 are high even when considering well-performing single features, indicating the need to jointly
224 consider multiple features for distinguishing phenotype genes and pseudogenes.

225 **Consideration of multiple features in combination produces accurate predictions of** 226 **functional genomic regions**

227 To consider multiple features in combination, we first conducted principle component (PC)
228 analysis to investigate how well phenotype genes and pseudogenes could be separated. Between

229 the first two PCs, which jointly explain 40% of the variance in the feature dataset, phenotype
230 genes (**Fig. 3A**) and pseudogenes (**Fig. 3B**) were distributed in largely distinct space. However,
231 there remains substantial overlap, indicating that standard parametric approaches are not well
232 suited to distinguishing between benchmark functional and nonfunctional sequences. Thus, we
233 instead considered all 93 features for phenotype gene and pseudogenes in combination using
234 random forest (referred to as the full model; see Methods). The phenotype gene and pseudogene
235 sequences and associated conservation, biochemical, and sequence-structure features were
236 separated into distinct training and testing sets and the full model was generated and validated
237 using independent data subsets (cross-validation). The full model provided more accurate
238 predictions (AUC-ROC=0.98; FNR=4%; FPR=10%; **Fig. 3C**) than any individual feature (**Fig.**
239 **2**; Supplemental Table 3). An alternative measure of performance based on the precision
240 (proportion of predicted functional sequences that are truly functional) and recall (proportion of
241 truly functional sequences predicted correctly) values among predictions generated by the full
242 model also indicated that the model was performing well (**Fig. 3D**). When compared to the best-
243 performing single feature (expression breadth), the full model had a similar FNR but half the
244 FPR (10% compared to 21%). Thus, the full model is highly capable of distinguishing between
245 phenotype genes and pseudogenes.

246 We next determined the relative contributions of different feature categories in predicting
247 phenotype genes and pseudogenes and whether models based on a subset of features would
248 perform similarly as the full model. Seven prediction models were established, each using only
249 the subset of features from a single category (**Fig. 2**). Although none of these category-specific
250 models had performance as high as the full model, the models based on transcription activity,
251 sequence conservation, and H3 mark features scored highly (AUC-ROC=0.97, 0.92, and 0.91,
252 respectively; **Fig. 3C**). Particularly, the transcription activity feature category model performed
253 almost as well as the full model (FNR=6%, FPR=12%). We emphasize that the breadth and level
254 of transcription are the causes of the strong performance of the transcription activity-only model,
255 not the presence of expression evidence.

256 To evaluate whether the strong performance of the full model is being driven solely by
257 transcription activity-related features, we also built a function prediction model did not consider
258 these features (full (-TX), **Fig. 3C,D**). We found that the model excluding transcription activity
259 features performed almost as well as the full model and similarly to the transcription activity-

260 feature-only model, but with an increased FPR (AUC-ROC=0.96; FNR=3%; FPR=20%). This
261 indicates that predictions of functional regions are not reliant solely on transcription data.
262 Instead, a diverse array of features can be considered to make highly accurate predictions of the
263 functionality of a genomic sequence. Meanwhile, our finding of the high performance of the
264 transcription activity-only model highlights the possibility of establishing an accurate model for
265 distinguishing functional genic and nonfunctional genomic sequences in plant species with only
266 a modest amount of transcriptome data.

267 **Functional likelihood allows the prediction of functional and nonfunctional genomic** 268 **regions**

269 To provide a measure of the potential functionality of any sequence in the *A. thaliana* genome,
270 including ITRs and ncRNAs, we utilized the confidence score from the full model as a
271 “functional likelihood” value (Tsai et al. 2017; see Methods). The functional likelihood (FL)
272 score ranges between 0 and 1, with high values indicating that a sequence is more similar to
273 phenotype genes (functional) and low values indicating a sequence more closely resembles
274 pseudogenes (nonfunctional). FL values for all genomic regions examined in this study are
275 available in Supplemental Table 4. As expected, phenotype genes had high FL values
276 (median=0.97; **Fig. 4A**) and pseudogenes had low values (median=0.01; **Fig. 4B**). To call
277 sequences as functional or not, we defined a threshold FL value of 0.35 (see Methods). Using
278 this threshold, 96% of phenotype genes (**Fig. 4A**) and 90% of pseudogenes (**Fig. 4B**) are
279 correctly classified as functional and nonfunctional, respectively, demonstrating that the full
280 model is highly capable of distinguishing functional and nonfunctional sequences.

281 We next applied our model to predict the functionality of annotated protein-coding genes,
282 transposable elements, and unexpressed intergenic regions. Most annotated protein-coding genes
283 not included in the phenotype gene dataset had high FL scores (median=0.86; **Fig. 4C**) and 80%
284 were predicted as functional. Of the 20% of protein-coding genes that were predicted as
285 nonfunctional, we expect that at least 4% represent false negatives based on the FNR of the full
286 model. The actual FNR among protein-coding genes may be higher, however, as phenotype
287 genes represent a highly active and well conserved subset of all genes. However, a subset of the
288 low-scoring protein-coding genes may also represent gene sequences undergoing functional
289 decay and *en route* to pseudogene status. To assess this possibility, we examined 1,940 *A.*

290 *thaliana* "decaying" genes that may be experiencing pseudogenization due to promoter
291 disablement (Yang et al. 2011) and found that, while these decaying genes represented only 7%
292 of all *A. thaliana* annotated protein-coding genes, they made up 45% of protein-coding genes
293 predicted as nonfunctional (Fisher's Exact Test (FET), $p < 1E-11$). In addition to protein-coding
294 genes, we evaluated the FLs of transposable elements (TEs) and randomly-selected, unexpressed
295 intergenic regions that are most likely nonfunctional. As expected, the FLs were low for both
296 TEs (median=0.03, **Fig. 4D**) and unexpressed intergenic regions (median=0.07; **Fig. 4E**), and
297 99% of TEs and all unexpressed intergenic sequences were predicted as nonfunctional, further
298 demonstrating the utility of the function prediction model. Overall, the FL measure provides a
299 useful metric to distinguish between phenotype genes and pseudogenes. In addition, the FLs of
300 annotated protein-coding genes, TEs, and unexpressed intergenic sequences agree with *a priori*
301 expectations regarding the functionality of these sequences.

302 **Exclusion of features from multiple tissues increases prediction performance for narrowly-** 303 **expressed sequences**

304 Although the full model performs exceedingly well, there remain false predictions. There are 76
305 phenotype genes (4%) predicted as nonfunctional (referred to as low-FL phenotype genes). We
306 assessed why these phenotype genes were not correctly identified by first asking what category
307 of features were particularly distinct between low-FL and the remaining phenotype genes. We
308 found that the major category that led to the misclassification of phenotype genes was
309 transcription activity, as only 7% of low-scoring phenotype genes were predicted as functional in
310 the transcription activity-only model, compared to 98% of high FL phenotype genes (**Fig. 5**). By
311 contrast, >65% of low-FL phenotype genes were predicted as functional when sequence
312 conservation, H3 mark, or DNA methylation features were used. This could suggest that the full
313 model is less effective in predicting functional sequences that are weakly or narrowly expressed.
314 While sequence conservation features are distinct between functional and nonfunctional
315 sequences when considered in combination, a significantly higher proportion of low-FL
316 phenotype genes were specific to the *Brassicaceae* family, with only 33% present in
317 dicotyledonous species outside of the *Brassicaceae*, compared to 78% of high-scoring phenotype
318 genes (FET, $p < 4e-12$), thus our model likely has reduced power in detecting lineage-specific
319 genes.

320 Given the association between transcription activity features and functional predictions,
321 we next investigated how functional predictions performed for conditionally-functional and
322 narrowly-expressed sequences. We found that genes with conditional phenotypes (see Methods)
323 had no significant differences in FLs (median=0.96) as those with phenotypes under standard
324 growth conditions (median=0.97; U test, $p=0.38$, Supplemental Fig. 1A). Thus, our model can
325 capture conditionally functional sequences. Next, we evaluated FL distributions among
326 sequences with different breadths of gene expression. For this comparison, we focused on non-
327 stress, single-tissue expression datasets (Supplemental Table 5), which was distinct from the
328 expression breadth feature in the prediction model that considered all datasets. While phenotype
329 genes were better predicted than pseudogenes among sequences with the same number of tissues
330 with expression evidence (U tests, all $p < 1.7E-06$; Supplemental Fig. 1B), 65% of the 62
331 phenotype genes expressed in ≤ 3 tissues were predicted as nonfunctional. Further, there was a
332 significant correlation between the number of tissues with expression evidence and FL values of
333 all sequences in our analysis ($r^2=0.77$; $p < 2E-16$). Thus, the function prediction model is biased
334 against narrowly-expressed phenotype genes.

335 We also found that 80 pseudogenes (10%) were defined as functional (high-FL
336 pseudogenes). Consistent with misclassifications among phenotype genes, a key difference
337 between high-FL pseudogenes and those that were correctly predicted as nonfunctional was that
338 high-FL pseudogenes were more highly and broadly expressed (**Fig. 5**). A significantly higher
339 proportion of high-FL pseudogenes came from existing genome annotation as 19% of annotated
340 pseudogenes were classified as functional, compared to 4% of pseudogenes identified through a
341 computational pipeline (Zou et al 2009) (FET, $p < 1.5E-10$). We found that high-FL pseudogenes
342 might be more recently pseudogenized and thus have not yet lost many genic signatures, as the
343 mean number of disabling mutations (premature stop or frameshift) per kb in high-scoring
344 pseudogenes (1.9) were significantly lower than that of low-scoring pseudogenes (4.0; U test, $p <$
345 0.02). Lastly, we cannot rule out the possibility that a small subset of high-scoring pseudogenes
346 represent truly functional sequences, rather than false positives (e.g. Poliseno et al. 2010; Karreth
347 et al. 2015). Overall, the misclassification of both narrowly-expressed phenotype genes and
348 broadly-expressed pseudogenes highlights the need for an updated prediction model that is less
349 influenced by expression breadth.

350 To tailor functional predictions to narrowly-expressed sequences, we generated a “tissue-
351 agnostic” model that attempts to minimize the contribution of biochemical activities occurring in
352 many tissues by excluding expression breadth and features that were available across multiple
353 tissues (see Methods). The tissue-agnostic model performed similarly to the full model (AUC-
354 ROC=0.97; FNR=4%; FPR=15%; Supplemental Fig. 2; Supplemental Table 4). Importantly, the
355 proportion of phenotype genes expressed in ≤ 3 tissues predicted as functional increased by 23%
356 (35% in the full model to 58% in the tissue-agnostic model, Supplemental Fig. 1C), indicating
357 that the tissue-agnostic model is more suitable for predicting the functionality of narrowly-
358 expressed sequences than the full model, although there was an increase in FPR (from 10% to
359 15%). We next sought to evaluate the FL of ITR and annotated ncRNA sequences utilizing both
360 the full model and the tissue-agnostic model, as these sequences were often narrowly-expressed
361 (Supplemental Fig. 3A).

362 **Intergenic transcribed regions and annotated ncRNAs are mostly predicted as** 363 **nonfunctional**

364 A subset of ITRs and ncRNAs likely represent novel genes or unannotated exon extensions of
365 known genes (Johnson et al. 2005; Moghe et al. 2013). Nevertheless, most ITRs and ncRNAs are
366 functionally ambiguous, as they are predominantly identified by the presence of expression
367 evidence and few have been characterized genetically. To evaluate the functionality of ITRs and
368 ncRNAs, we applied both the full and tissue-agnostic models to 895 ITRs, 136 TAIR ncRNAs,
369 and 252 Araport long ncRNAs (referred to as Araport ncRNAs; see Methods). The median FLs
370 based on the full model were low (0.09) for both ITRs (**Fig. 4F**) and Araport ncRNAs (**Fig. 4G**),
371 and only 15% and 9% of these sequences were predicted as functional, respectively. By contrast,
372 TAIR ncRNAs had a significantly higher median FL value (0.53; U tests, both $p < 5e-31$; **Fig.**
373 **4H**) and 68% were predicted as functional. The higher proportion of functional TAIR ncRNA
374 predictions compared to ITRs and Araport ncRNAs could be best explained by differences in
375 features from the transcription activity category (**Fig. 5**). We also note that a greater proportion
376 of ITRs and Araport ncRNAs are predicted as functional when considering only DNA
377 methylation or H3 mark features (**Fig. 5**). However, these two category-specific models also had
378 higher false positive rates (unexpressed intergenic sequences and pseudogenes, **Fig. 5**). Thus,

379 single feature models do not provide additional support for the functionality of most Araport
380 ncRNAs and ITRs.

381 We next applied the tissue-agnostic model that is less biased against narrowly-expressed
382 sequences (Supplemental Fig. 1C) to ITRs and TAIR/Araport ncRNAs that were generally
383 narrowly-expressed (Supplemental Fig. 3A). Compared to the full model, around twice as many
384 ITRs (30%) and Araport ncRNAs (19%) but a similar number of TAIR ncRNA (67%) were
385 predicted as functional. Considering the union of the full and tissue-agnostic model predictions,
386 268 ITRs (32%), 57 Araport ncRNAs (23%), and 105 TAIR ncRNAs (77%) were likely
387 functional. ITRs and annotated ncRNAs closer to annotated genes tended to be predicted as
388 functional (Supplemental Fig. 4A). Using the 95th percentile of intron lengths for all genes as a
389 threshold to call ITRs and annotated ncRNAs as proximal or distal to neighboring genes, 57% of
390 likely functional and 35% of likely nonfunctional ITRs and ncRNAs were proximal to
391 neighboring genes, respectively (FET, $p < 2E-09$). To assess if a subset these likely functional,
392 proximal ITRs/ncRNAs may be unannotated exons of known genes, we assessed whether they
393 tended to have similar features with their neighbors. Compared to feature similarities between
394 neighboring and random gene pairs (Supplemental Fig. 4B-D), likely functional ITRs/ncRNAs
395 were less similar to their neighbors, regardless of proximity (Supplemental Fig. 4C,D). Thus,
396 despite their proximity to annotated genes, it remains unclear if some ITRs or annotated ncRNAs
397 represent unannotated exon extensions of known genes or not. In addition, for proximal
398 functional ITRs/annotated ncRNAs, we cannot rule out the possibility that they represent false-
399 positive functional predictions due to the accessible and active chromatin states of nearby genes.
400 Given the challenge in ascertaining the origin of likely functional, proximal ITRs/ncRNAs, we
401 instead conservatively estimate that 116 distal, functional ITRs and annotated ncRNAs may
402 represent fragments of novel genes.

403 **Intergenic transcribed regions and annotated ncRNAs do not resemble benchmark RNA** 404 **genes**

405 Thus far, we predicted the majority of ITR and annotated ncRNA sequences as nonfunctional.
406 We demonstrated that the full model was able to predict conditional phenotype genes
407 (Supplemental Fig. 1A) and the tissue-agnostic model was more effective than the full model in
408 predicting narrowly expressed phenotype genes (Supplemental Fig. 1B,C). Thus, conditional or

409 tissue-specific functionality do not fully explain why the majority of ITRs and ncRNAs are
410 predicted as nonfunctional. However, the function prediction models so far were built by
411 contrasting protein-coding genes with pseudogenes and it remains possible that these protein-
412 coding gene-based models can not accurately predict RNA genes. To evaluate this possibility,
413 we generated a tissue-agnostic model using features calculated from a randomly-selected 100 bp
414 sequence within a phenotype protein-coding gene or pseudogene body (for features, see
415 Supplemental Table 6). The reason for using 100 bp sequence is that most RNA genes are too
416 short to be considered by earlier models, which were based on 500 bp sequences. In addition,
417 features from the tissue agnostic model are more suitable for RNA gene prediction as annotated
418 RNA genes tend to be more narrowly expressed than phenotype genes (U tests, all $p < 2e-05$;
419 Supplemental Fig. 3B). The 100 bp tissue-agnostic model performed similarly to the full 500 bp
420 model in distinguishing between phenotype protein-coding genes and pseudogenes, except with
421 higher FNR (AUC-ROC=0.97; FNR=13%; FPR=5%; Supplemental Fig. 5), but only predicted
422 three out of six RNA genes with documented mutant phenotypes (phenotype RNA genes) as
423 functional (Supplemental Fig. 5I). Further, other RNA Pol II-transcribed RNA genes exhibited
424 mixed predictions from the 100 bp tissue-agnostic model, as 15% of microRNA (miRNA)
425 primary transcripts (Supplemental Fig. 5J), 73% of small nucleolar RNAs (snRNAs;
426 Supplemental Fig. 5K), and 50% of small nuclear RNAs (snRNAs; Supplemental Fig. 5L) were
427 predicted as functional. Although the proportion of phenotype RNA genes predicted as
428 functional (50%) is significantly higher than the proportion of pseudogenes predicted as
429 functional (5%, FET, $p < 0.004$), this finding suggests that a model built with protein-coding
430 genes has a substantial FNR for detecting RNA genes.

431 To determine whether the suboptimal predictions by the phenotype protein-coding gene-
432 based models are because RNA genes belong to a class of their own, we next built a multi-class
433 function prediction model aimed at distinguishing four classes of sequences: benchmark RNA
434 genes ($n=46$), phenotype protein-coding genes (1,882), pseudogenes (3,916), and randomly-
435 selected, unexpressed intergenic regions (4,000). Benchmark RNA genes include six phenotype
436 RNA genes and 40 high-confidence miRNA primary transcript sequences (see Methods).
437 Unexpressed intergenic sequences were included to provide another set of likely nonfunctional
438 sequences distinct from pseudogenes. Expression breadth and tissue-specific features were
439 excluded from the four-class model and 100 bp sequences were used. In the four-class model,

440 87% of benchmark RNA genes, including all six phenotype RNA genes, were predicted as
441 functional sequences (65% RNA gene-like and 22% phenotype protein-coding gene-like; **Fig.**
442 **6A**). In addition, 95% of phenotype genes were predicted as functional (**Fig. 6B**), including 80%
443 of narrowly expressed genes, an increase of 22% over the 500 bp tissue-agnostic model
444 (Supplemental Fig. 1B). For the remaining two sequence classes, 70% of pseudogenes (**Fig. 6C**)
445 and 100% of unexpressed intergenic regions (**Fig. 6D**) were predicted as nonfunctional (either
446 pseudogenes or unexpressed intergenic sequences). Thus, the four-class model improves
447 prediction accuracy of RNA genes and narrowly expressed genes. However, the inclusion of
448 RNA genes in the model has significantly increased the ambiguity in pseudogene classification.

449 Since the four-class model was able to distinguish benchmark RNA genes from
450 nonfunctional sequence classes, we next evaluated whether ITRs and annotated ncRNAs
451 resemble functional sequences with the four-class model. Note that the 100 bp model used here
452 allowed us to evaluate an additional 10,938 ITRs and 1,406 annotated ncRNAs. We find that
453 34% of ITR, 38% of Araport ncRNA, and of 65% TAIR ncRNAs were predicted as functional
454 sequences (**Fig. 6E-G**). To provide an overall estimate of the proportion of likely-functional
455 ITRs and annotated ncRNAs, we considered the predictions from the four-class model (**Fig. 6**),
456 the full model (**Fig. 3,4**), and the tissue-agnostic models (Supplemental Fig. 2,5). Based on
457 support from at least one of the four models, we classified 4,437 ITRs (38%) and 796 annotated
458 ncRNAs (44%) as functional, as they resembled either phenotype protein-coding or RNA genes.
459 Our findings lend support that they are likely parts of novel or annotated genes. Meanwhile, we
460 find that a substantial number of ITRs (62%) and annotated ncRNAs (56%) are predicted as
461 nonfunctional. Moreover, at least a third of ITRs (**Fig. 6E**) and Araport ncRNAs (**Fig. 6F**) most
462 closely resemble unexpressed intergenic regions. Thus, we show that the majority of ITRs and
463 annotated ncRNA regions resemble nonfunctional genomic regions, and therefore could
464 represent regions of noisy transcription.

465 **CONCLUSION**

466 Discerning the location of functional regions within a genome represents a key goal in genomic
467 biology. Despite advances in computational gene finding, it remains challenging to determine
468 whether intergenic transcribed regions (ITRs) represent functional or noisy biochemical activity.
469 We established robust function prediction models based on the evolutionary, biochemical, and

470 structural characteristics of phenotype genes and pseudogenes. The prediction models accurately
471 define functional and nonfunctional regions and are applicable genome-wide. These results echo
472 recent findings that human phenotype genes could be distinguished from pseudogenes (Tsai et al.
473 2017). Given that function predictions were successful in both plant and metazoan model
474 systems, integrating the evolutionary and biochemical features of known genes will likely be
475 applicable to any species. The next step will be to test whether function prediction models can be
476 applied across species, which could ultimately allow the phenotype data and omics resources
477 available in model systems to effectively guide the identification of functional regions in non-
478 models.

479 Expression data was highly informative to functional predictions. We found that the
480 prediction model based on only 24 transcription activity-related features performs nearly as well
481 as the full model that integrates additional information including conservation, H3 mark,
482 methylation, and TF binding data. In human, use of transcription data from cell lines also
483 produced highly accurate predictions of functional genomic regions (AUC-ROC=0.96; Tsai et al.
484 2017). Despite the importance of transcription data, we emphasize that the presence of
485 expression evidence is an extremely poor predictor. Taken together, these results indicate that
486 function prediction models can be established in any species, model or not, with a modest
487 number of transcriptome datasets (e.g. 51 in this study and 19 in human). One caveat of the
488 current model is that narrowly-expressed phenotype genes are frequently predicted as
489 pseudogene and broadly-expressed pseudogenes tend to be called functional. To improve the
490 function prediction model, it will be important to explore additional features unrelated to
491 transcription. Because few phenotype genes are narrowly-expressed (5%) in the *A. thaliana*
492 training data, more phenotyping data for narrowly expressed genes will be crucial as well.

493 Upon application of the function prediction models genome-wide, we found that 4,427
494 ITRs and 796 annotated ncRNAs in *A. thaliana* are likely functional. Assuming each entry
495 equals a novel gene, this estimate represents a 19% increase in annotated gene space (excluding
496 annotated ncRNAs) for the model plant. However, considering the high false positive rates (e.g.
497 10% for the full and 31% for the four-class model), this is most likely an overestimate of the
498 number of novel genes contributed by functional ITRs and annotated ncRNAs. In addition, we
499 emphasize that the majority of ITRs and ncRNAs resemble pseudogenes and random
500 unexpressed intergenic regions. Similarly, most human ncRNAs are more similar to

501 nonfunctional sequences than they are to protein coding and RNA genes (Tsai et al. 2017).
502 Furthermore, the significant relationship between the amount of intergenic expression occurring
503 in a species and the size of a genome is consistent with the interpretation that intergenic
504 transcripts are generally nonfunctional. Thus, instead of assuming any expressed sequence must
505 be functionally significant, we advocate that the null hypothesis should be that it is not,
506 particularly considering that most ITRs and annotated ncRNAs have not been experimentally
507 characterized. The machine learning framework we have described provides an approach for
508 distinguishing between functional and noisy biochemical activity, and will help defining the gene
509 space in a genome.

510 **METHODS**

511 **Identification of transcribed regions in leaf tissue of 15 flowering plants**

512 RNA-sequencing (RNA-seq) datasets were retrieved from the Sequence Read Archive (SRA) at
513 the National Center for Biotechnology Information (NCBI; www.ncbi.nlm.nih.gov/sra/) for 15
514 flowering plant species (Supplemental Table 1). All datasets were generated from leaf tissue and
515 sequenced on Illumina HiSeq 2000 or 2500 platforms. Genome sequences and gene annotation
516 files were downloaded from Phytozome v.11 (www.phytozome.net; Goodstein et al. 2011) or
517 Oropetium Base v.01 (www.sviridis.org; VanBuren et al. 2015). Genome sequences were repeat
518 masked using RepeatMasker v4.0.5 (www.repeatmasker.org) if a repeat-masked version was not
519 available. Only one end from paired-end read datasets were utilized in downstream processing.
520 Reads were trimmed to be rid of low scoring ends and residual adaptor sequences using
521 Trimmomatic v0.33 (LEADING:3 TRAILING:3 SLIDINGWINDOW:4:20 MINLEN:20, Bolger
522 et al. 2014) and mapped to genome sequences using TopHat v2.0.13 (default parameters except
523 as noted below; Kim et al. 2013). Reads ≥ 20 nucleotides in length that mapped uniquely within a
524 genome were used in further analysis.

525 For each species, thirty million mapped reads were randomly selected from among all
526 datasets and assembled into transcript fragments using Cufflinks v2.2.1 (default parameters
527 except as noted below, Trapnell et al. 2010), while correcting for sequence-specific biases during
528 the sequencing process by providing an associated genome sequence with the -b flag. The
529 expected mean fragment length for assembled transcript fragments in Cufflinks was set to 150
530 from the default of 200 so that expression levels in short fragments would not be overestimated.

531 The 1st and 99th percentile of intron lengths for each species were used as the minimum and
532 maximum intron lengths, respectively, for both the TopHat2 and Cufflinks steps. Intergenic
533 transcribed regions (ITRs) were defined by transcript fragments that did not overlap with gene
534 annotation and did not have significant six-frame translated similarity to plant protein sequences
535 in Phytozome v.10 (BLASTX E-value < 1E-05). The correlation between assembled genome
536 size and gene counts was determined with data from the first 50 published plant genomes
537 (Michael and Jackson, 2013).

538 **Phenotype data sources**

539 Mutant phenotype data for *A. thaliana* protein-coding genes was collected from a published
540 dataset (Lloyd and Meinke 2012), the Chloroplast 2010 database (Ajjawi et al. 2010; Savage et
541 al. 2013), and the RIKEN phenome database (Kuromori et al. 2006) as described by Lloyd et al.
542 (2015). Phenotype genes used in our analyses were those whose disruption resulted in lethal or
543 visible defects under standard laboratory growth conditions. Genes with documented mutant
544 phenotypes under standard conditions were considered as a distinct and non-overlapping
545 category from other annotated protein-coding genes. We identified six RNA genes with
546 documented loss-of-function phenotypes through literature searches (Supplemental Table 7): *At4*
547 (AT5G03545; Shin et al. 2006), *MIR164A* and *MIR164D* (AT2G47585 and AT5G01747,
548 respectively; Guo et al. 2005), *MIR168A* (AT4G19395; Li et al. 2012b), and *MIR828A* and *TAS4*
549 (AT4G27765 and AT3G25795, respectively; Hsieh et al. 2009). Conditional phenotype genes
550 were those belonging to the Conditional phenotype group as described by Lloyd and Meinke
551 (2012). Loss-of-function mutants of these genes exhibited phenotype only under stress
552 conditions.

553 ***Arabidopsis thaliana* genome annotation**

554 *A. thaliana* protein-coding gene, miRNA gene, snoRNA gene, snRNA gene, ncRNA region,
555 pseudogene, and transposable element annotations were retrieved from The Arabidopsis
556 Information Resource v.10 (TAIR10; www.arabidopsis.org; Berardini et al. 2015). Additional
557 miRNA gene and lncRNA region annotations were retrieved from Araport v.11
558 (www.araport.org; Krishnakumar et al. 2015). A pseudogene-finding pipeline (Zou et al. 2009)
559 was used to identify additional pseudogene fragments and count the number of disabling

560 mutations (premature stop or frameshift mutations). Genes, pseudogenes, and transposons with
561 overlapping annotation were excluded from further analysis. Overlapping lncRNA annotations
562 were merged for further analysis. When pseudogenes from TAIR10 and the pseudogene-finding
563 pipeline overlapped, the longer pseudogene annotation was used.

564 *A. thaliana* ITRs analyzed include: (1) the Set 2 ITRs in Moghe et al. (2013), (2) the
565 novel transcribed regions from Araport v.11, and (3) additional ITRs from 206 RNA-seq datasets
566 (Supplemental Table 5). Reads were trimmed, mapped, and assembled into transcript fragments
567 as described above, except that overlapping transcript fragments from across datasets were
568 merged. ITRs analyzed did not overlap with any TAIR10, Araport11, or pseudogene annotation.
569 Overlapping ITRs from different annotated subsets were kept based on a priority system:
570 Araport11 > Set 2 ITRs from Moghe et al. (2013) > ITRs identified in this study. For each
571 sequence entry (gene, ncRNA, pseudogene, transposable element, or ITR), a 100 and 500 base
572 pair (bp) window was randomly chosen for calculating feature values and subsequent model
573 building steps. Feature descriptions are provided in the following sections. The feature values for
574 randomly selected 500 and 100 bp windows are provided in Supplemental Table 2 and 6,
575 respectively. Additionally, non-expressed intergenic sequences were randomly-sampled from
576 genome regions that did not overlap with annotated genes, pseudogenes, transposable elements,
577 or regions with genic or intergenic transcript fragments (100 bp, n=4,000; 500 bp, n=3,716). All
578 100 and 500 bp windows described above are referred to as sequence windows throughout the
579 Methods section.

580 **Sequence conservation and structure features**

581 There were 10 sequence conservation features examined. The first two were derived from
582 comparisons between *A. thaliana* accessions including nucleotide diversity and Tajima's D
583 among 81 accessions (Cao et al. 2011) using a genome matrix file from the 1,001 genomes
584 database (www.1001genomes.org). The python scripts are available through GitHub
585 (<https://github.com/ShiuLab/GenomeMatrixProcessing>). The remaining eight features were
586 derived from cross-species comparisons, three based on multiple sequence and five based on
587 pairwise alignments. Three multiple sequence alignment-based features were established using
588 aligned genomic regions between *A. thaliana* and six other plant species (*Glycine max*,
589 *Medicago truncatula*, *Populus trichocarpa*, *Vitis vinifera*, *Sorghum bicolor*, and *Oryza sativa*)

590 (referred to as conserved blocks). For each conserved block, the first feature was the proportion
591 of a sequence window that overlapped a conserved block (referred to as coverage), and the two
592 other features were the maximum and average phastCons scores within each sequence window.
593 The phastCons score was determined for each nucleotide within conserved blocks (Li et al.
594 2012a). Nucleotides in a sequence window that did not overlap with a conserved block were
595 assigned a phastCons score of 0. For each sequence window, five pairwise alignment-based
596 cross-species conservation features were the percent identities to the most significant BLASTN
597 match (if $E\text{-value} < 1E\text{-}05$) in each of five taxonomic groups. The five taxonomic groups included
598 the *Brassicaceae* family ($n_{\text{species}}=7$), other dicotyledonous plants (22), monocotyledonous plants
599 (7), other embryophytes (3), and green algae (5). If no sequence with significant similarity was
600 present, percent identity was scored as zero.

601 For sequence-structure features, we used 125 conformational and thermodynamic
602 dinucleotide properties collected from DiProDB database (Friedel et al. 2009). Because the
603 number of dinucleotide properties was high and dependent, we reduced the dimensionality by
604 utilizing principal component (PC) analysis as described previously (Tsai et al. 2015). Sequence-
605 structure values corresponding to the first five PCs were calculated for all dinucleotides in and
606 averaged across the length of a sequence window and used as features when building function
607 prediction models.

608 **Transcription activity features**

609 We generated four multi-dataset and 20 individual dataset transcription activity features. To
610 identify a set of RNA-seq datasets to calculate multi-dataset features, we focused on the 72 of
611 206 RNA-seq datasets each with ≥ 20 million reads (see above; Supplemental Table 5).
612 Transcribed regions were identified with TopHat2 and Cufflinks as described in the RNA-seq
613 analysis section except that the 72 *A. thaliana* RNA-seq datasets were used. Following transcript
614 assembly, we excluded 21 RNA-seq datasets because they had unusually high RPKM (Reads Per
615 Kilobase of transcript per Million mapped reads) values (median RPKM value
616 range= $272\sim 2,504,294$) compared to the rest ($2\sim 252$). The remaining 51 RNA-seq datasets were
617 used to generate four multi-dataset transcription activity features including: expression breadth,
618 95th percentile expression level, maximum transcript coverage, and presence of expression
619 evidence (for values see Supplemental Table 5). Expression breadth was the number of RNA-seq

620 datasets that have ≥ 1 transcribed region that overlapped with a sequence window. The 95th
621 percentile expression level was the 95th percentile of RPKM values across 51 RNA-seq datasets
622 where RPKM values were set to 0 if there was no transcribed region for a sequence window.
623 Maximum transcript coverage was the maximum proportion of a sequence window that
624 overlapped with a transcribed region across 51 RNA-seq datasets. Presence of expression
625 evidence was determined by overlap between a sequence window and any transcribed region in
626 the 51 RNA-seq datasets.

627 In addition to features based on multiple datasets, 20 individual dataset features were
628 derived from 10 datasets: seven tissue/organ-specific RNA-seq datasets including pollen
629 (SRR847501), seedling (SRR1020621), leaf (SRR953400), root (SRR578947), inflorescence
630 (SRR953399), flower, (SRR505745) and silique (SRR953401), and three datasets from non-
631 standard growth conditions, including dark-grown seedlings (SRR974751) and leaf tissue under
632 drought (SRR921316) and fungal infection (SRR391052). For each of these 10 RNA-seq
633 datasets, we defined two features for each sequence window: the maximum transcript coverage
634 (as described above) and the maximum RPKM value of overlapping transcribed regions (referred
635 to as Level in **Fig. 2**). If no transcribed regions overlapped a sequence window, the maximum
636 RPKM value was set as 0. For the analysis of narrowly- and broadly-expressed phenotype genes
637 and pseudogenes (Supplemental Fig. 1B,C), we used 28 out of 51 RNA-seq datasets generated
638 from a single tissue and in standard growth conditions to calculate the number of tissues with
639 evidence of expression (tissue expression breadth). In total, seven tissues were represented
640 among the 28 selected RNA-seq datasets (see above; Supplemental Table 5), and thus tissue
641 expression breadth ranges from 0 to 7 (note that only 1 through 7 are shown in Supplemental Fig.
642 1B,C due to low sample size of phenotype genes in the 0 bin). The tissue breadth value is distinct
643 from the expression breadth feature used in model building that was generated using all 51
644 datasets and considered multiple RNA-seq datasets from the same tissue separately (range: 0-
645 51).

646 **Histone 3 mark features**

647 Twenty histone 3 (H3) mark features were calculated based on eight H3 chromatin
648 immunoprecipitation sequencing (ChIP-seq) datasets from SRA. The H3 marks examined
649 include four associated with activation (H3K4me1: SRR2001269, H3K4me3: SRR1964977,

650 H3K9ac: SRR1964985, and H3K23ac: SRR1005405) and four associated with repression
651 (H3K9me1: SRR1005422, H3K9me2: SRR493052, H3K27me3: SRR3087685, and H3T3ph:
652 SRR2001289). Reads were trimmed as described in the RNA-seq section and mapped to the
653 TAIR10 genome with Bowtie v2.2.5 (default parameters; Langmead et al. 2009). Spatial
654 Clustering for Identification of ChIP-Enriched Regions (SICER) v.1.1 (Xu et al. 2014) was used
655 to identify ChIP-seq peaks with a false discover rate ≤ 0.05 with a non-overlapping window size
656 of 200, a gap parameter of 600, and an effective genome size of 0.92 according to Koehler et al.
657 (2011). For each H3 mark, two features were calculated for each sequence window: the
658 maximum intensity among overlapping peaks and peak coverage (proportion of overlap with the
659 peak that overlaps maximally with the sequence window). In addition, four multi-mark features
660 were generated. Two of the multi-mark features were the number of activating marks (0-4)
661 overlapping a sequence window and the proportion of a sequence window overlapping any peak
662 from any of the four activating marks (activating mark peak coverage). The remaining two multi-
663 mark features were the same as the two activating multi-mark features except focused on the four
664 repressive marks.

665 **DNA methylation features**

666 Twenty-one DNA methylation features were calculated from bisulfite-sequencing (BS-seq)
667 datasets from seven tissues (pollen: SRR516176, embryo: SRR1039895, endosperm:
668 SRR1039896, seedling: SRR520367, leaf: SRR1264996, root: SRR1188584, and inflorescence:
669 SRR2155684). BS-seq reads were trimmed as described above and processed with Bismark v.3
670 (default parameters; Krueger and Andrews 2011) to identify methylated and unmethylated
671 cytosines in CG, CHH, and CHG (H = A, C, or T). Methylated cytosines were defined as those
672 with ≥ 5 mapped reads and with $>50\%$ of mapped reads indicating that the position was
673 methylated. For each BS-seq dataset, the percentage of methylated cytosines in each sequence
674 window for CG, CHG, and CHH contexts were calculated if the sequence window had ≥ 5
675 cytosines with ≥ 5 reads mapping to the position. To determine whether the above parameters
676 were reasonable, we assessed the false positive rate of DNA methylation calls by evaluating the
677 proportion of cytosines in the chloroplast genome that are called as methylated, as the
678 chloroplast genome has few DNA methylation events (Ngernprasirtsiri et al. 1988; Zhang et al.
679 2006). Based on the above parameters, 0-1.5% of cytosines in CG, CHG, or CHH contexts in the

680 chloroplast genome were considered methylated in any of the seven BS-seq datasets. This
681 indicated that the false positive rates for DNA methylation calls were low and the parameters
682 were reasonable.

683 **Chromatin accessibility and transcription factor binding features**

684 Chromatin accessibility features consisted of ten DHS-related features and one micrococcal
685 nuclease sequencing (MNase-seq)-derived feature. DHS peaks from five tissues (seed coat,
686 seedling, root, unopened flowers, and opened flowers) were retrieved from the Gene Expression
687 Omnibus (GSE53322 and GSE53324; Sullivan et al. 2014). For each of the five tissues, the
688 maximum DHS peak intensity and DHS peak coverage were calculated for each sequence
689 window. Normalized nucleosome occupancy per bp based on MNase-seq was obtained from Liu
690 et al. (2015). The average nucleosome occupancy value was calculated across each sequence
691 window. Transcription factor (TF) binding site features were based on *in vitro* DNA affinity
692 purification sequencing data of 529 TFs (O'Malley et al. 2016). Two features were generated for
693 each sequence window: the total number of TF binding sites and the number of distinct TFs
694 bound.

695 **Single-feature prediction performance**

696 The ability for each single feature to distinguish between functional and nonfunctional regions
697 was evaluated by calculating AUC-ROC value with the Python scikit-learn package (Pedregosa
698 et al. 2011). AUC-ROC values range between 0.5 (equivalent to random guessing) and 1 (perfect
699 predictions) and values above 0.7, 0.8, and 0.9 are considered to be fair, good, and excellent,
700 respectively. Thresholds to predict sequences as functional or nonfunctional using a single
701 feature were defined by the feature value that produced the highest F-measure, the harmonic
702 mean of precision (proportion of sequences predicted as functional that are truly functional) and
703 recall (proportion of truly functional sequences predicted as functional). The F-measure allows
704 consideration of both false positives and false negatives at a given threshold. FPR were
705 calculated as the percentage of negative (nonfunctional) cases with values above or equal to the
706 threshold and thus falsely predicted as functional. FNR were calculated as the percentage of
707 positive (functional) cases with values below the threshold and thus falsely predicted as
708 nonfunctional.

709 **Binary classification with machine learning**

710 For binary classification (two-class) models that contrasted phenotype genes and pseudogenes,
711 the random forest (RF) implementation in the Waikato Environment for Knowledge Analysis
712 software (WEKA; Hall et al. 2009) was utilized. Three types of two-class models were
713 established, including the full model (500 bp sequence window, **Fig. 3C,D** and **Fig. 4**), tissue-
714 agnostic models (500 bp, Supplemental Fig. 2; 100 bp, Supplemental Fig. 5), and single feature
715 category models (**Fig. 3C,D**). For each model type, we first generated 100 balanced datasets by
716 randomly selecting equal numbers of phenotype genes (positive examples) and pseudogenes
717 (negative examples). For each of these 100 datasets, 10-fold stratified cross-validation was
718 utilized, where the model was trained using 90% of sequences and tested on the remaining 10%.
719 Thus, for each model type, a sequence window had 100 prediction scores, where each score was
720 the proportion of 500 random forest trees that predicted a sequence as a phenotype gene in a
721 balanced dataset. The median of 100 prediction scores was used as the functional likelihood (FL)
722 value (Supplemental Table 4). The FL threshold to predict a sequence as functional or
723 nonfunctional was defined based on maximum F-measure as described in the previous section.
724 We tested multiple -K parameters (2 to 25) in the WEKA-RF implementation, which alters the
725 number of randomly-selected features included in each RF tree (Supplemental Table 8), and
726 found that 15 randomly-selected features provided the highest performance based on AUC-ROC
727 (calculated and visualized using the ROCR package; Sing et al. 2005). Binary classification
728 models were also built using all features from 500 bp sequences (equivalent to the full model)
729 with the Sequential Minimal Optimization - Support Vector Machine (SMO-SVM)
730 implementation in WEKA (Hall et al. 2009). The results of SMO-SVM models were highly
731 similar to the full RF results: *PCC* between the FL values generated by RF and SMO-
732 SVM=0.97; AUC-ROC of SMO-SVM=0.97; FPR=12%; FNR=3%. By comparison, the full RF
733 model had AUC-ROC=0.98, FPR=10%, FNR=4%.

734 Tissue-agnostic models were generated by excluding the expression breadth feature and
735 95th percentile expression level and replacing all features from RNA-seq, BS-seq, and DHS
736 datasets that were available in multiple tissues. For multiple-tissue RNA-seq data, the maximum
737 expression level across 51 RNA-seq datasets (in RPKM) and maximum coverage (as described
738 in the transcription activity section) of a sequence window in any of 51 RNA-seq datasets were
739 used. For multi-tissue DNA methylation features, minimum proportions of methylated cytosines

740 in any tissue in CG, CHG, and CHH contexts were used. For DHS data, the maximum peak
741 intensity and peak coverage was used instead. In single feature category predictions, fewer total
742 features were used and therefore lower $-K$ values (i.e. the number of random features selected
743 when building random forests) were considered in parameter searches (Supplemental Table 8).

744 **Multi-class machine learning model**

745 For the four-class model, benchmark RNA gene, phenotype protein-coding gene, pseudogene,
746 and random unexpressed intergenic sequences were used as the four training classes. Benchmark
747 RNA genes consisted of six RNA genes with documented loss-of-function phenotypes and 40
748 high-confidence miRNA genes from miRBase (www.mirbase.org; Kozomara and Griffiths-Jones
749 2014). We generated 250 datasets with equal proportions (larger classes randomly sampled) of
750 training sequences. Two-fold stratified cross-validation was utilized due to the low number of
751 benchmark RNA genes. The features included those described for the tissue-agnostic model and
752 focused on 100 bp sequence windows. The RF implementation, *cforest*, in the *party* package of
753 R (Strobl et al. 2008) was used to build the classifiers. The four-class predictions provide
754 prediction scores for each sequence type: an RNA gene, phenotype protein-coding gene,
755 pseudogene, and unexpressed intergenic score (Supplemental Table 4). The prediction scores
756 indicate the proportion of random forest trees that classify a sequence as a particular class.
757 Median prediction scores from across 100 balanced runs were used as final prediction scores.
758 Scores from a single balanced dataset models sum to 1, but not the median from 100 balanced
759 runs. Thus, the median scores were scaled to sum to 1. For each sequence window, the maximum
760 prediction score among the four classes was used to classify a sequence as phenotype gene,
761 pseudogene, unexpressed intergenic, or RNA gene.

762 **FIGURE LEGENDS**

763 **Figure 1.** Relationship between genome size and number of nucleotides covered by RNA-seq
764 reads (expression) in 15 flowering plant species. (A) annotated genic regions. (B) intergenic
765 regions excluding any annotated features. Mb: megabase. Gb: gigabase. Dotted lines: linear
766 model fits. r^2 : square of Pearson's correlation coefficient.

767 **Figure 2.** Predictions of functional (phenotype gene) and non-functional (pseudogene) sequences
768 based on each individual feature. Prediction performance is measured using Area Under the

769 Curve - Receiver Operating Characteristic (AUC-ROC). Features include those in the categories
770 of (A) transcription activity, (B) sequence conservation, (C) DNA methylation, (D) transcription
771 factor (TF) binding, (E) histone 3 (H3) marks, (F) sequence structure, and (G) chromatin
772 accessibility. AUC-ROC ranges in value from 0.5 (equivalent to random guessing) to 1 (perfect
773 predictions). Dotted lines: median AUC-ROC of features in a category.

774 **Figure 3.** Predictions of functional and nonfunctional sequences based on multiple features.
775 Smoothed scatterplots of the first two principle components (PCs) of (A) phenotype gene and (B)
776 pseudogene features. The percentages on the axes in (A) indicate the feature value variation
777 explained by the associated PC. (C) AUC-ROC values of function prediction models built when
778 considering all features (Full), all except transcription activity (TX)-related features (Full (-TX)),
779 and all features from each category. The category abbreviations follow those in **Fig. 2.** (C)
780 Precision-recall curves of the models with matching colors from (B). The models were built
781 using feature values calculated from 500 bp sequence windows.

782 **Figure 4.** Functional likelihood distributions of various sequence classes based on the full
783 model. (A) Phenotype genes. (B) Pseudogenes. (C) Annotated protein-coding genes. (D)
784 Transposable elements. (E) Random unexpressed intergenic sequences. (F) Intergenic
785 transcribed regions (ITR). (G) Araport11 ncRNAs. (H) TAIR10 ncRNAs. The full model was
786 established using 500 bp sequence windows. Higher and lower functional likelihood values
787 indicate greater similarity to phenotype genes and pseudogenes, respectively. Vertical dashed
788 lines indicate the threshold for calling a sequence as functional or nonfunctional. The
789 percentages to the left and right of the dashed line indicate the percent of sequences predicted as
790 functional or nonfunctional, respectively.

791 **Figure 5.** Proportion of phenotype genes, pseudogenes, ITRs, and ncRNAs predicted as
792 functional in the full and single-category models. Percentages of sequence classes that are
793 predicted as functional in models based on all features and the single category models, each
794 using all features from a category (abbreviated according to **Figure 2.** The models are sorted
795 from left to right based on performance (AUC-ROC). The colors of and numbers within the
796 blocks indicate the proportion sequences predicted as functional by a given model. Phenotype
797 gene and pseudogene sequences are shown in three sub-groups: all sequences (All), and those

798 predicted as functional (high functional likelihood (FL)) and nonfunctional (low FL) in the full
799 model. ITR: intergenic transcribed regions.

800 **Figure 6.** Function predictions based on a four-class prediction model. (A) Stacked bar plots
801 indicate the prediction scores of benchmark RNA genes for each of the four classes: dark blue -
802 phenotype protein-coding gene (Ph), cyan - RNA gene (RNA), red - pseudogene (Ps), yellow -
803 random intergenic sequence (Ig). A benchmark RNA gene is classified as one of the four classes
804 according to the highest prediction score. The color bars below the chart indicate the predicted
805 class, with the same color scheme as the prediction score. Sequences classified as Ph or RNA
806 were considered functional, while those classified as Ps or Ig were considered nonfunctional.
807 Percentages below a classification region indicate the proportion of sequences classified as that
808 class. (B) Phenotype protein-coding gene prediction scores. (C) Pseudogene prediction scores.
809 (D) Random unexpressed intergenic region prediction scores. Note that no sequence was
810 predicted as functional. (E) Intergenic transcribed region (ITR), (F) Araport11 ncRNA regions.
811 (G) TAIR10 ncRNA regions.

812

813 SUPPLEMENTAL FIGURE LEGENDS

814 **Supplemental Figure 1.** Impacts of conditional phenotypes and expression breadth on the
815 function prediction model. (A) Functional likelihood distributions of phenotype genes with
816 mutant phenotypes under standard growth conditions (non-conditional) and non-standard growth
817 conditions such as stressful environments (conditional) based on the 500 bp full model. Feature
818 values were calculated from a random 500 bp region from within the sequence body. Higher and
819 lower functional likelihood values indicate a greater similarity to phenotype genes and
820 pseudogenes, respectively. (B,C) Distributions of functional likelihood scores for phenotype
821 genes (blue) and pseudogenes (red) for sequences with various breadths of expression for (B) the
822 500 bp full model and (C) the 500 bp tissue-agnostic model generated by excluding the
823 expression breadth and features available from multiple tissues. The tissue-agnostic model is
824 aimed toward minimizing the effects of biochemical activity occurring across multiple tissues
825 and predicts a greater proportion of narrowly-expressed phenotype genes as functional compared
826 to the full model.

827 **Supplemental Figure 2.** Distributions of functional likelihood scores based on the 500 bp tissue-
828 agnostic model. (A) Phenotype genes. (B) Pseudogenes. (C) Annotated protein-coding genes. (D)
829 Transposable elements. (E) Random unexpressed intergenic sequences. (F) Intergenic
830 transcribed regions (ITR). (G) Araport11 ncRNAs. (H) TAIR10 ncRNAs. Vertical dashed lines
831 display the threshold to define a sequence as functional or nonfunctional. The numbers to the left
832 and right of the dashed line show the percentage of sequences predicted as functional or
833 nonfunctional, respectively.

834 **Supplemental Figure 3.** Distributions of expression breadth of different sequence classes. (A)
835 Based on 500 bp feature regions. (B) Based on 100 bp feature regions.

836 **Supplemental Figure 4.** Distance of ITRs and annotated ncRNA regions to and feature
837 similarity with neighboring genes. (A) Distance from intergenic transcribed regions (ITRs) and
838 annotated ncRNAs to the closest neighboring gene. ITR and ncRNA sequences are separated by
839 whether they are predicted as functional (F) or nonfunctional (NF) by the 500 bp full model. (B)
840 Feature similarity based on Pearson's Correlation Coefficients (PCC) between random pairs of
841 ITRs, Araport11 ncRNAs, TAIR10 ncRNAs, or annotated genes. (C) Feature similarity between
842 proximal neighbors (within 95th percentile (456 bp) of intron lengths), and (D) Feature similarity
843 between distal neighbors (>456 bp). Pairs involving ITRs and annotated ncRNAs were divided
844 by whether the ITR or ncRNA sequence was predicted as functional (F) or nonfunctional (NF)
845 by the full model. Feature values were quantile normalized prior to calculating correlations.

846 **Supplemental Figure 5.** Distributions of functional likelihood scores based on the 100 bp tissue-
847 agnostic model. (A) Phenotype genes. (B) Pseudogenes. (C) Protein-coding gene. (D)
848 transposable elements. (E) Random unexpressed intergenic sequences. (F) Intergenic transcribed
849 regions (ITR). (G) Araport11 ncRNAs. (H) TAIR10 ncRNAs. (I) RNA genes with loss-of-
850 function mutant phenotypes. (J) MicroRNAs, (K) Small nucleolar RNAs, (L) Small nuclear
851 RNAs. The tissue-agnostic model was built with 100 bp features and while excluding the
852 expression breadth and tissue-specific features. Higher functional likelihood values indicate
853 greater similarity to phenotype genes while lower values indicate similarity to pseudogenes.
854 Vertical dashed lines display the threshold to define a sequence as functional or nonfunctional.

855 The numbers to the left and right of the dashed line show the percentage of sequences predicted
856 as functional or nonfunctional, respectively.

857 **SUPPLEMENTAL TABLES**

858 **Supplemental Table 1.** Leaf tissue RNA-sequencing datasets for 15 flowering plant species

859

860 **Supplemental Table 2.** Conservation, biochemical, and sequence-structure feature values
861 calculated from 500 bp sequences.

862

863 **Supplemental Table 3.** False positive and false negative rates for single feature classifications.

864

865 **Supplemental Table 4. Function** predictions for all models generated in this study.

866

867 **Supplemental Table 5.** RNA-sequencing datasets for identifying intergenic transcribed regions,
868 calculating transcription activity features, and assessing tissue-specific predictions.

869

870 **Supplemental Table 6.** Conservation, biochemical, and sequence-structure feature values
871 calculated from 100 bp sequences.

872

873 **Supplemental Table 7.** RNA genes with documented loss-of-function phenotypes.

874

875 **Supplemental Table 8.** K parameters tested for random forest runs.

876 **DATA ACCESS**

877 All data are available in the text of this article or in the supplemental material.

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884 **AUTHOR CONTRIBUTIONS**

885 J.P.L., Z.T.-Y.T., and S.-H.S. designed the research. J.P.L., Z.T.-Y.T., R.P.S., and N.L.P.
886 performed the research. J.P.L., Z.T.-Y.T., R.P.S., N.L.P., and S.-H.S. wrote the article.

887 **DISCLOSURE DECLARATION**

888 The authors have no conflicts of interest to disclose.

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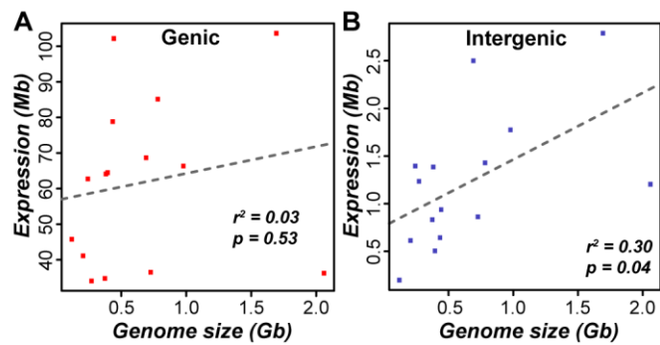
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- 1124

1125 **FIGURES**

Figure 1.



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Figure 2.

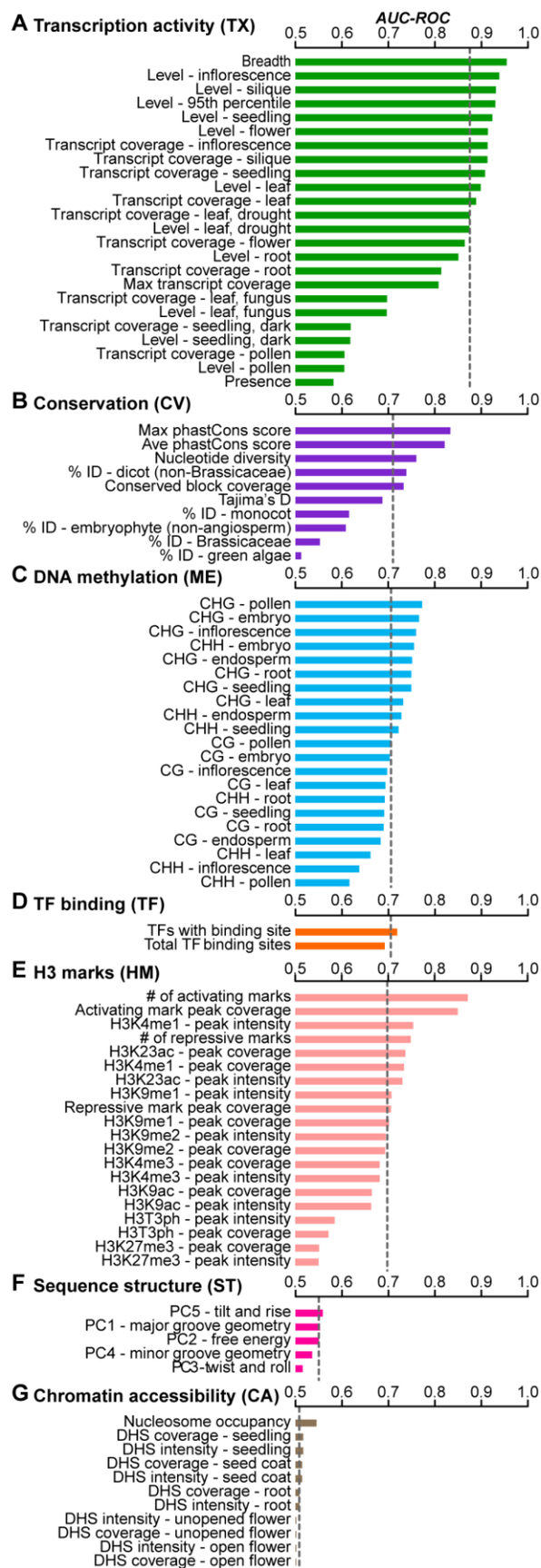
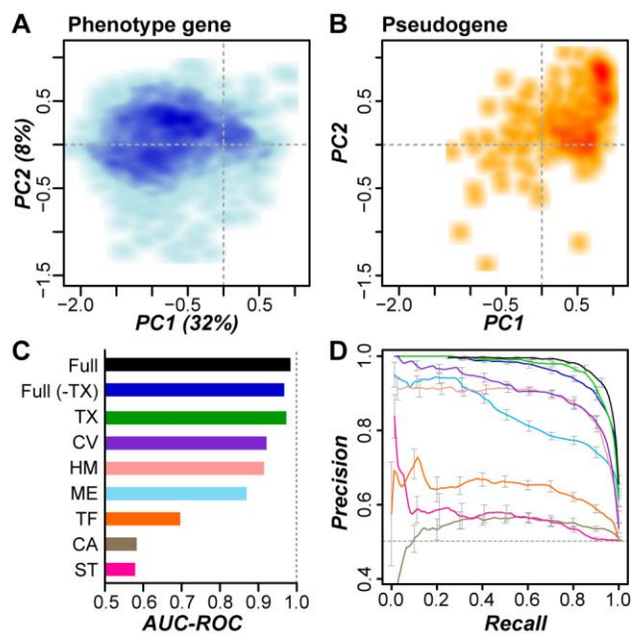


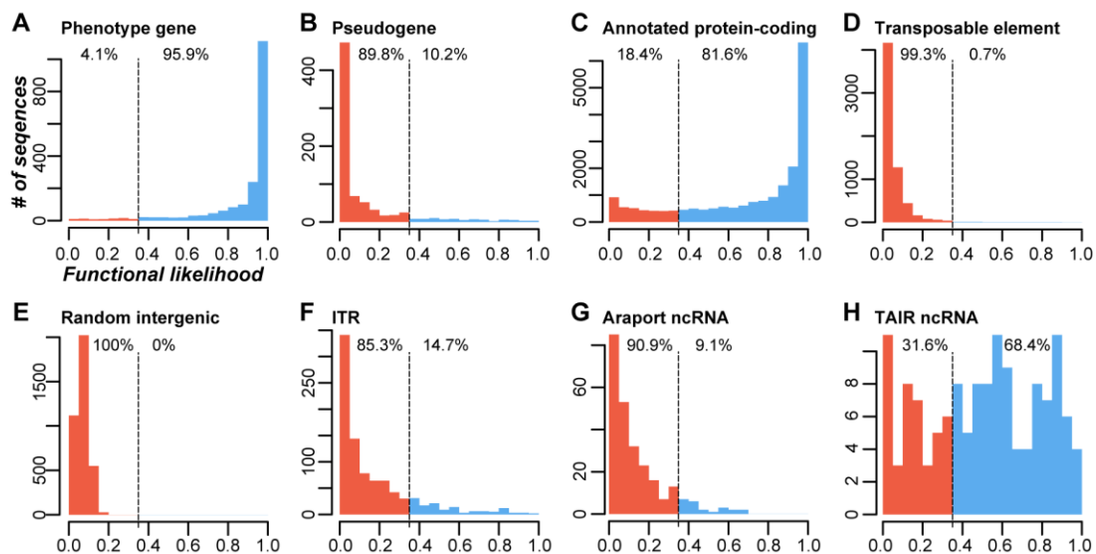
Figure 3.



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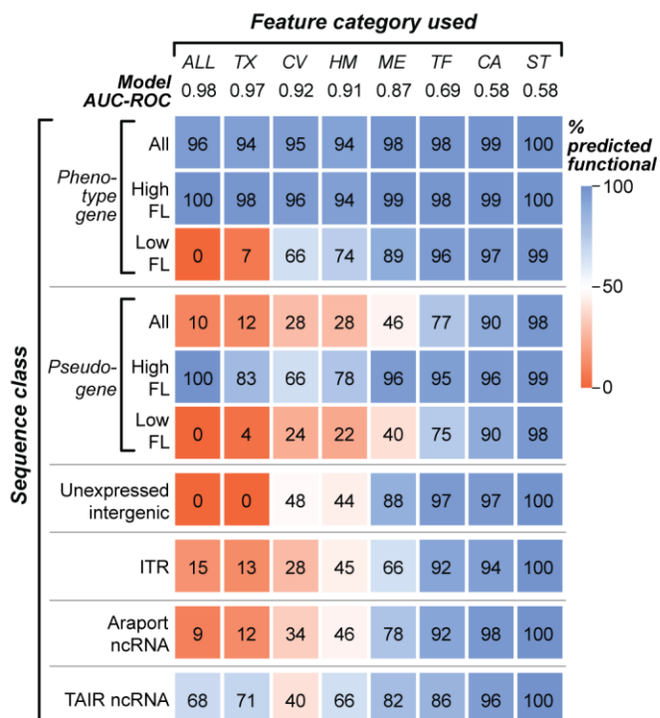
Figure 4.



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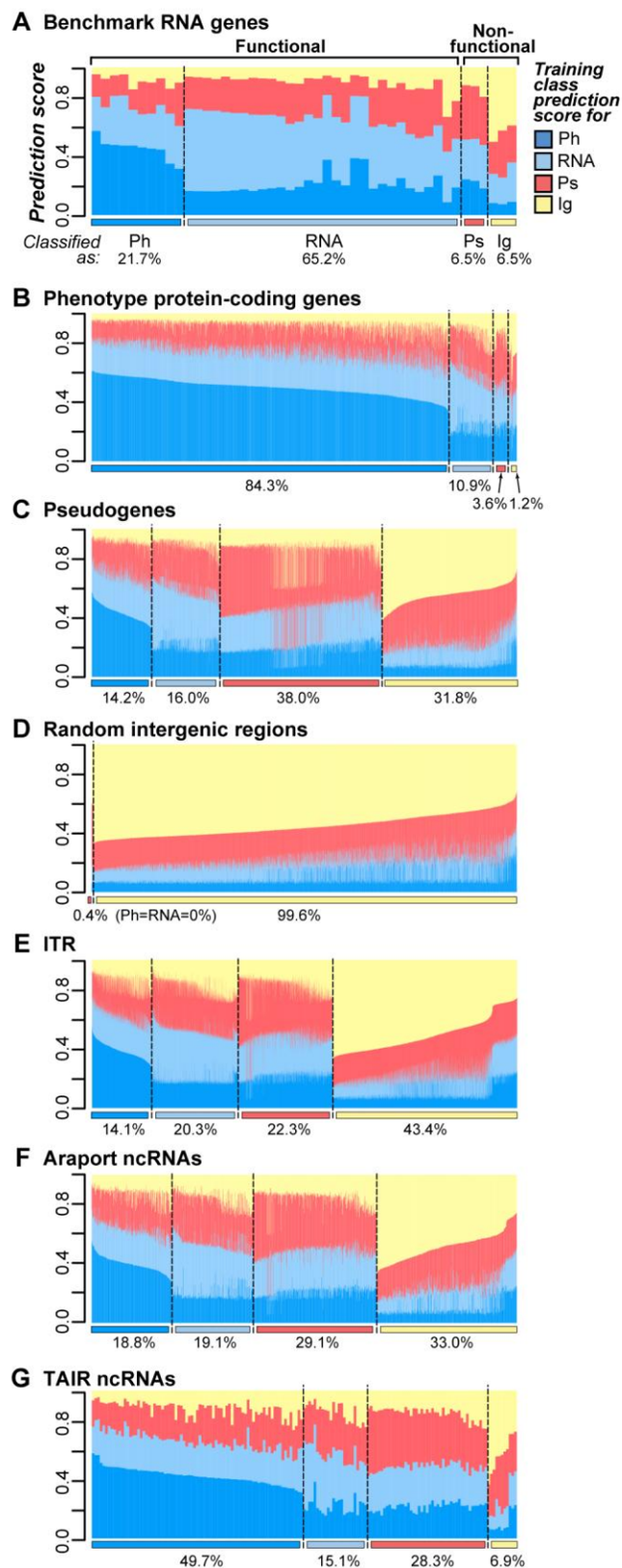
Figure 5.



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Figure 6.

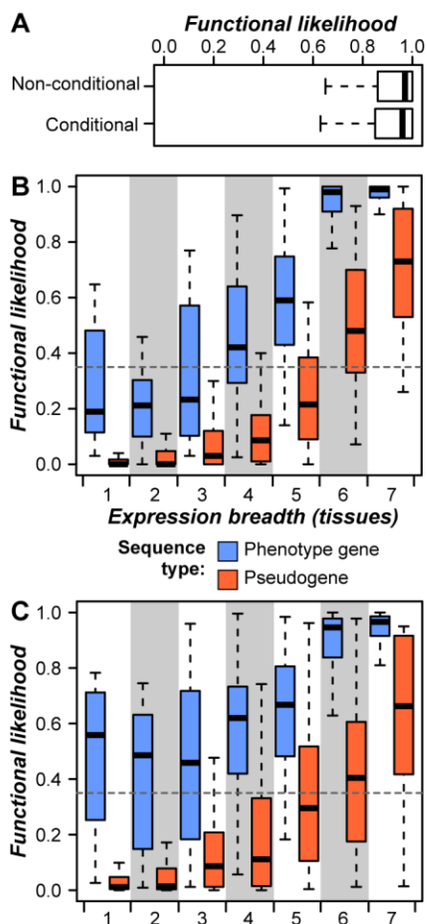


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1137 **SUPPLEMENTAL FIGURES**

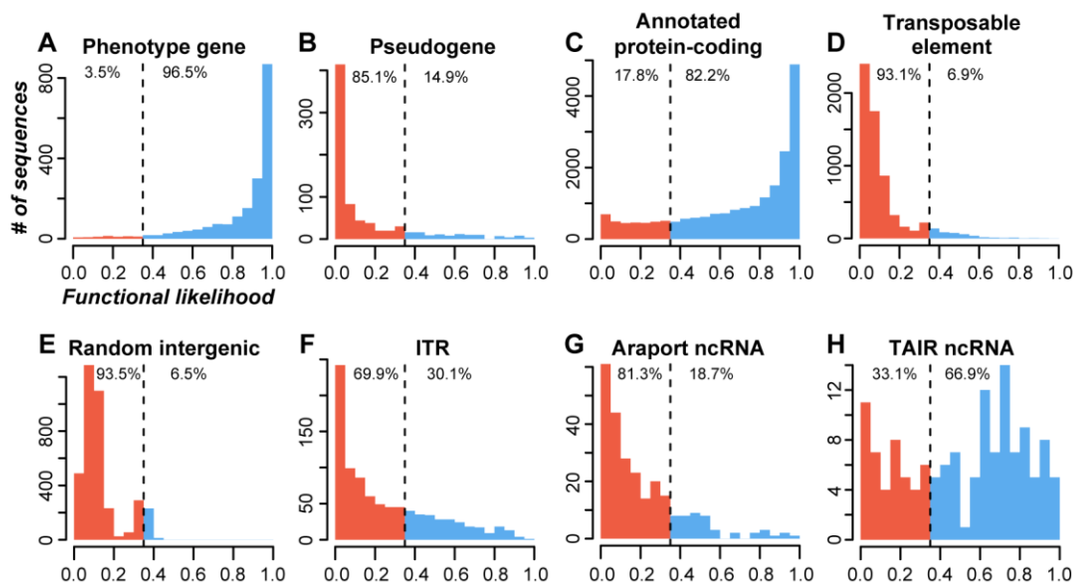
**Supplemental
Figure 1.**



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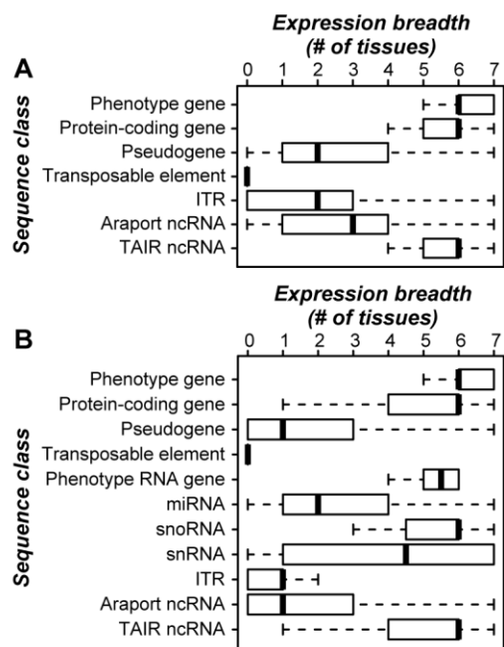
Supplemental Figure 2.



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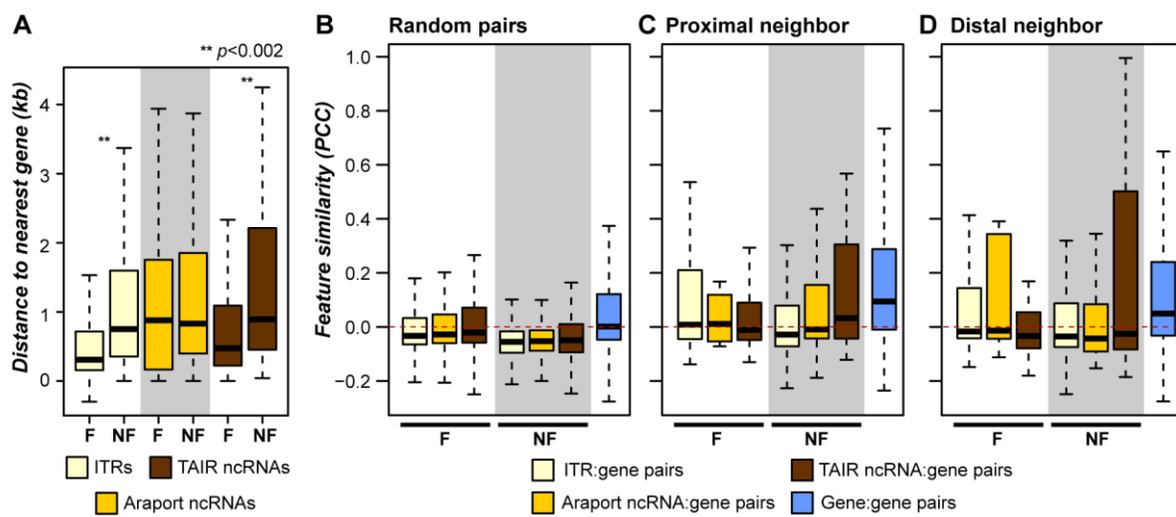
**Supplemental
Figure 3.**



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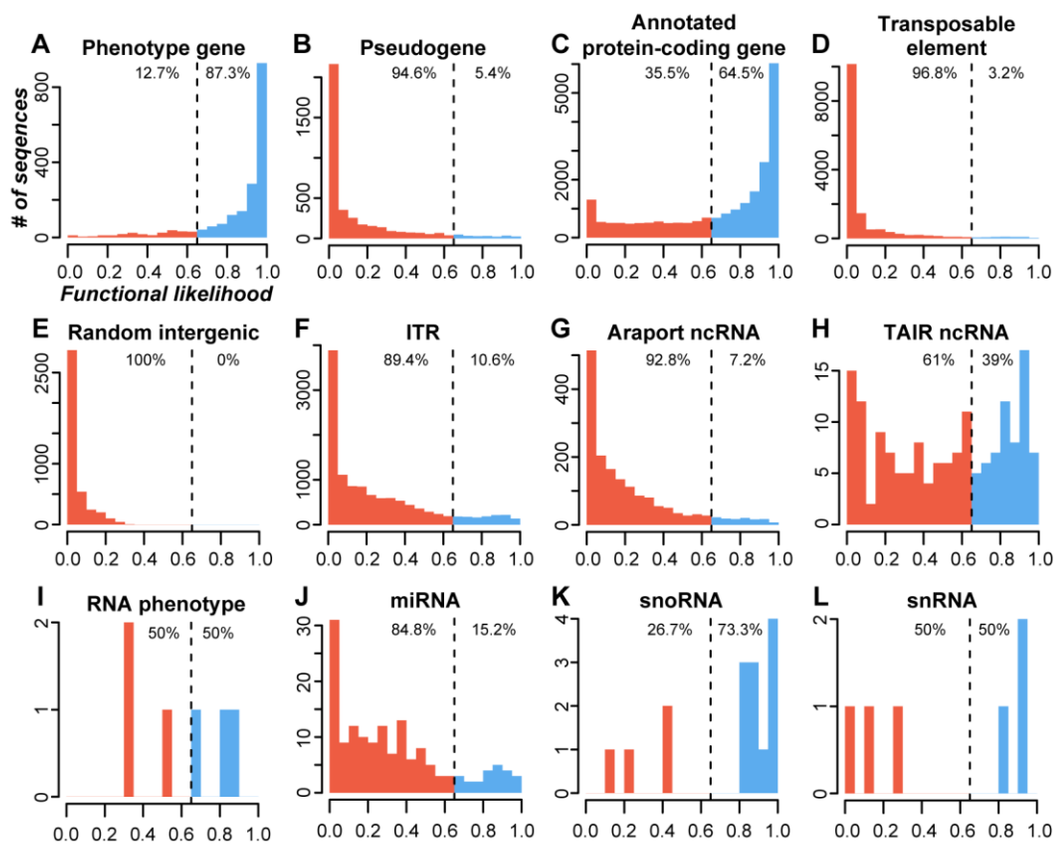
Supplemental
Figure 4.



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Supplemental Figure 5.



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