METHODOLOGY

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Tissue Enrichment Analysis: TEA

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Abstract

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Over the last ten years, there has been an explosive development in tools capable of measuring gene expression. These tools generate a large number of gene targets, but understanding these datasets, and forming hypotheses based on them remains challenging.

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We present a method for detecting tissue enrichment in *C. elegans* using the tissue ontology for this organism. We also present an efficient method for trimming the ontology that results in concise yet useful output.

Our tool, Tissue Enrichment Analysis (TEA), can be found at www.wormbase.org/tea

Keywords: Gene Ontology; Tissue Ontology; Wormbase

19Background

RNA-seq and other high-throughput methods in biology have the ability to identify thousands of genes that are altered between conditions. These genes are often correlated in their biological characteristics or functions, but identifying these functions remains challenging. In order to interpret these long lists of genes, biologists need to abstract genes into fewer terms that are biologically relevant in order to form thy protheses about what is happening in the data. One such abstraction method rehappening in the form of an directed acyclic graph[1–3] that provide detailed information the molecular, cellular or biochemical functions of the gene among others. For a given gene list, certain software programs can query whether a particular gene is enriched[4–6]. However, GO is often difficult to interpret due to the large number of terms associated with a given gene. There exist a number of GO analytic tools for use by the community but a shared complaint for many programs is the very large number of GO terms that are significantly associated with any given gene list.

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¹ A common tool for GO analysis, DAVID, clusters terms into broad categories that ¹
$^2{\rm are}$ amenable to exploration by researchers [7], whereas PANTHER, a different soft- 2
³ ware package [4, 8], attempts to solve this issue by employing a manually reduced ³
⁴ ontology, GOslim (pers. comm.).
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$_8$ Here we provide a new framework that analyses user-input list for $\mathrm{enrichment}_8$
$_{9}\mathrm{of}$ specific tissues. We believe that tissues are physiologically relevant units with $_{9}$
$_{10}{\rm broad},$ relatively well-understood functionalities amenable to hypothesis formation. $_{10}$
₁₁ As such, we believe that identification of tissues is likely to provide researchers _{1:}
$_{12}$ with enough information to be able to form hypotheses about the physiological $_{12}$
$_{13}{\rm responses}$ of an organism to a specified condition. Our analysis also cuts down on $_{13}{\rm responses}$
₁₄ result verbosity by filtering the ontology before testing using a small set of well- ₁₄
15 defined criteria to remove terms that don't contribute extra information. To our
16knowledge, such filtering has never been performed in an algorithmic fashion for
$_{17}$ an ontology before — indeed, tools such as DAVID do not employ term trimming $_{17}$
$_{18}a\ priori$ of testing, but rather fuzzy clustering $post$ testing to reduce the number $_{18}$
$_{19}$ of ontology terms. We believe our trimming methodology strikes a good balance $_{19}$
$_{20}$ between detailed tissue calling and conservative testing.
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We built our geftwere using a pre-established tiggue entels on for the many C. els.
We built our software using a pre-established tissue ontology for the worm, C. ele-
gans [9]. The C. elegans database, Wormbase[10], maintains a carefully curated list
of gene expression data from GFP-reporters. We use this gold-standard list to de-
velop a tissue enrichment analysis that reliably identifies even small tissues and show
that we can reliably discriminate between embryonic and larval tissues. Our tool is
available in Wormbase at the address http://mangolassi.caltech.edu/azurebrd/cgi-
²⁹ bin/testing/amigo/getWithPost.cgi and provides users with a text-based file of the
enrichment results as well as a simple and clear graph of the results that exhibit
the largest fold-change enrichment. Although we present results here for the worm,
we note that our software is species agnostic, and we are working to integrate tissue
ontologies from other databases to provide a broader service to the community.

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¹Methods ²Generating a Useful Dictionary ³Reducing term redundancy through a similarity metric ⁴As a first step to generate our tissue enrichment software, we wished to select tissue ⁵terms that were reasonably well-annotated, yet specific enough to provide insight ⁶ and not redundant with other terms. We also wanted to avoid testing tissues at ⁶ ⁷levels where redundancy becomes problematic. For example, several left and right 8 neurons have at least 25 annotating genes and we may want to include them for 8 $^9\mathrm{enrichment}$ testing. However, many left/right neuronal sisters have almost entirely 9 the same annotations, with at most one or two gene differences between them. We 10 reasoned that when two tissues have almost identical annotations, we cannot have statistical confidence in differentiating between them. As a result, testing these sister tissues provides no additional information compared with testing only the parent ¹⁴ node to these sisters. We refer to such sisters as 'redundant'. In order to identify ¹⁴ 15 redundancy, we defined a similarity metric 16 16 17 $s_i = \frac{|g_i|}{|\bigcup_{i=0}^k g_i|}$ $(1)^{18}$ 19

Where s_i is the similarity for a tissue i eith k sisters; g_i refers to the set of tissues₂₀ 2₁associated with tissue i and |g| refers to the cardinality of set g. For a given set₂₁ 2₂of sisters, we called them redundant if they exceeded a given similarity threshold.₂₂ 2₃We envisioned two possible criteria and built different dictionaries using each one.₂₃ 2₄Under a threshold criteron 'any' with parameter S between (0,1), a given set of₂₄ 2₅sisters j was considered redundant if the condition

$$s_{i,j} > S$$
 (2)₂₈

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was true for any sister i in set j. Under a threshold criterion 'avg' with parameter 30 S a given set of sisters i was considered redundant if the condition

S, a given set of sisters j was considered redundant if the condition

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was true for the set of sisters j (see figure 1).	1
2	2
³ Terminal branch terms and parent terms can be safely removed in an algorithmic	3
$_{4}fashion$	4
$_5$ Another problem arises from the fact that the tissue ontology is scarcely populated	5
$_{6}\mathrm{at}$ this point in time. Many nodes have 0-10 annotations, which we consider too few	6
$_{7}\mathrm{to}$ accurately test. To solve this issue, we implemented a straightforward trimming.	7
$_{8}$ algorithm. For a given terminal node, we test whether the node has more than a	8
9threshold number of annotations. If it does not, the node is removed. The next	9
$_{10}\mathrm{node}$ in the branch is tested and removed recursively until a node which satisfies	10
$_{11}$ the condition is found. At that point, no more nodes can be removed from that	11
$_{12}\mathrm{branch}.$ This is guaranteed by the structure of the ontology: Parent nodes inherit	12
$_{13} \mathrm{all}$ of the annotations of all of their descendants, so the number of annotating terms	13
$_{14} \mathrm{monotonically}$ increases with increasing term hierarchy (see figure 2). In this way,	14
$_{15} \mathrm{we}$ ensure that our term dictionary includes only those tissues that are considered	15
16 sufficiently well annotated for statistical purposes.	16
$_{17}$ $$ Finally, we also wanted to remove as many terms as possible from the dictionary	17
$_{18} \mathrm{with}$ the goals of reducing covariance between terms, decreasing multiple testing and	18
$_{19}\mathrm{removing}$ as many non-informative terms as possible. Decreasing covariance between	19
$_{20}\mathrm{terms}$ is important because we employ a frequent ist approach that assumes all terms	20
$_{21} \mathrm{are}$ independent. Large covariation coefficients between some terms means that if	21
$_{22}$ one of these tissues tests significant, the other terms are much more likely to pass	22
$_{23}{\rm significance}$ testing as well. This makes adequate correction for false positive rates	23
$_{24} {\rm considerably}$ more difficult. Moreover, from a data analysis perspective, we reasoned	24
$_{25}\mathrm{that},$ for any parent node, if all its daughters were selected for testing, there was no	25
$_{26}$ additional benefit to test the parent. In other words, if all the daughter nodes are	26
$_{27} \mathrm{tested},$ there is little additional information to be gained by including the parent	27
$_{28} \mathrm{node}.$ To address this issue we removed parent nodes from the analysis if all their	28
$_{29} {\rm daughter}$ nodes passed the annotation threshold (see figure 3).	29
Filtering greatly reduces the number of nodes used for analysis	30
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filtering are commutative, applying the ceiling filter is not commutative with either	33

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the trimming or the redundancy filter. If the ceiling filter is applied before any cother filter, only terminal nodes will remain, since all the parents have complete daughter sets. Since terminal nodes are the most poorly annotated, after applying daughter sets. Since terminal nodes will be left behind if any. On the other hand, the remaining filters very few nodes will be left behind if any. On the other hand, for applying the ceiling operator after trimming and redundancy filtering will result in forgreater numbers of nodes. We always applied the ceiling at the end. For validation forgreater numbers of nodes and the end of different dictionaries. The original ontology has for a factor of the set of the filtering of terms in granged from 21 to a maximum of 400 terms, which shows the number of terms in scarcely annotated ontology can be reduced by tenfold by application of a few simple filters.

These filters were used to compile a static dictionary that we employ for all anal-¹² ¹³yses. Because we have integrated our scripts to draw on the WormBase databases, ¹³ ¹⁴ our dictionary will remain up to date as tissue expression data improves. Our com-¹⁴ ¹⁵ pleted static trimmed dictionary is available for download at the following ftp URL: ¹⁵ ¹⁶XX. The final dictionary includes XX tissues for testing, and has XX annotating ¹⁶ ¹⁷ genes. All code was implemented in Python.

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19 Tissue enrichment testing via a hypergeometric model

 $_{20}$ Having built a static dictionary, we generated a Python script that implements $_{20}$ $_{21}$ significance testing algorithm based on the hypergeometric model. Briefly, the hy- $_{21}$ $_{22}$ pergeometric model assumes the existence of an urn with a pre-determined number $_{22}$ $_{23}$ of balls inside it. The balls can be painted one of several colors. The hypergeometric $_{23}$ $_{24}$ model provides an answer to the question: If an individual removes N balls, what $_{24}$ $_{25}$ is the probability of observing n_i balls of color i, if the balls are selected without $_{25}$ $_{26}$ replacement? Mathematically, this is expressed as:

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$$P(n_i|N, m_1, \dots, m_k, M) = \frac{\binom{m_i}{n_i} \binom{M - m_i}{N - n_i}}{\binom{N}{n_i}}$$
(4)29
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Here, n_i is the number of balls of type i drawn, N is the total number of draws, ³² m_i is tissue i and $M = \sum_i m_i$ is the total number of balls in the urn. In our specific ³³ case, M_i is equal to the total number of annotations in our dictionary. N is found ³³

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¹ by taking the user-input list and removing any genes that are not in our annotation ¹
$^2\mathrm{dictionary}.$ The remaining genes are then associated with their annotation $\mathrm{profiles}^2$
$^3-$ if a tissue is associated with s tissues, it generates s balls of s colors. Our program $^3-$
$^4\mathrm{counts}$ the number of times each tissue appears in the user list, and calculates the 4
$^5\mathrm{probability}$ of having with drawn as many or more balls for each tissue in the user 5
$^6\mathrm{list.}$ Due to the discrete nature of the hypergeometric distribution, this algorithm 6
$^7\mathrm{can}$ generate artifacts when the list is small. To avoid spurious results, a tissue is 7
$^8\mathrm{never}$ considered significant if there are no annotations for it in the user-provided 8
⁹ list.
Once the probability of drawing the labels has been quantified, we apply a stan- 10
$^{11}\mathrm{dard}\ \mathrm{FDR}\ \mathrm{correction}\ \mathrm{using}\ \mathrm{a}\ \mathrm{Benjamini\text{-}Hochberg}\ \mathrm{step\text{-}up}\ \mathrm{algorithm}[11].$ Genes that 11
$^{12}\mathrm{have}$ a q-value less than a given alpha are considered significant. Our default setting 12
13 is to set the alpha threshold at 0.1, but users will be able to modify this value either 13
14 in batch or in our web application. The program returns a text-based table showing 14
$^{15}{\rm the}$ tissues that tested significant, along with their associated q-value, the expected 15
$^{16}\mathrm{number}$ of hits for a list of that size, the observed number of hits and the enrichment 16
$^{17}\mathrm{fold}$ change (observed hits / expected hits). Finally, the program can also return a 17
$^{18}\mathrm{bar}$ chart of the enrichment fold change for the fifteen tissues with the largest en^{-18}
¹⁹ richment fold change. Our software relies heavily on the Pandas, Numpy, Seaborn ¹⁹
and SciPy modules to perform all statistical testing and data handling [12–14]. $$ 20
Our software is implemented in an easy to use GUI within WormBase. Users input ²¹
a gene-list (see figure 4) using any valid gene name for $C.$ elegans. These names are
processed into standard WBIDs and the result is displayed in the same window in 23
an easy to read format containing all the relevant information, and a graph of the 24
results is also displayed (see figure 5).
26
²⁷ Validation of the algorithm and parameter selection
In order to select an appropriate dictionary and validate our tool, we found a set of 28
29 30 gold standards based on microarray and RNA-seq literature which are believed
to be enriched in specific tissues [15–22]. Some of these studies went on to use GFP 30
to identify expression patterns and for this reason we generated a clean Since the 31
expression data is curated from GFP expression at this time and does not include 32
RNA-seq data, these gold standards are statistically independent from the dataset.

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¹We wanted to select a dictionary which included enough terms to be specific beyond ¹ ²the largest C. elegans tissues, yet would minimize the number of spurious results² ³and which had a good dynamic range in terms of enrichment fold-change. Selection³ ⁴of a dictionary based only on minimization of spurious results would result in a dic-⁴ ⁵tionary with a large number of annotations per tissue, and would therefore include ⁵ ⁶only the major tissues. On the other hand, selecting a dictionary that can detect⁶ ⁷smaller tissues will bias us towards tissues with lesser annotations. To our knowl-⁷ ⁸edge there is no good method for assessing false-positive or false-negative results⁸ ⁹for annotations. »»»; master As a first attempt to select a good dictionary, we generated all the possible combinations of dictionaries with minimal annotations of 10, 25, 50 and 100 genes and 11 similarity cutoffs of 0.9, 0.95 and 1, using 'average' or 'any' thresholding criteria for 12 ¹³the latter (see table 1). For these dictionaries, the number of tissues tested ranged ¹³ ¹⁴ from 97 to 676. The number of tissues was inversely correlated to the minimum ¹⁴ ¹⁵ annotation, as expected, and was largely insensitive to the redundancy threshold, ¹⁵ ¹⁶ at least in the range we explored (0.9-1). Next, we analyzed all 30 datasets using ¹⁶ each dictionary. Because of the large number of results, instead of analyzing each set 17 ¹⁸ of terms individually, we pooled all results for a given dictionary into histograms. ¹⁸ ¹⁹When we analyzed the distribution of significant q-values for the dictionaries, we ¹⁹ ²⁰ found that the similarity threshold mattered relatively little for any dictionary. We ²⁰ ²¹ also noticed that the 'any' thresholding method resulted in tighter histograms with ²¹ ²² a mode closer to 0 (data not shown). For this reason, we chose the 'any' method ²² ²³ for dictionary generation. The average q-value increased with decreasing annotation ²³ ²⁴cut-off (see figure 6), which reflects the decreasing statistical power associated with ²⁵ fewer annotations per term, but we remained agnostic as to how significant the ²⁵ trade-off between power and term specificity is. Based on these observations, we 26 ²⁷ ruled out the dictionary with the 100 annotation cut-off - it had the fewest terms ²⁸ and its q-values were not low enough to compensate the trade-off in specificity. To select between dictionaries generated between 50, 33 and 25 annotation cut- 30 offs, and also to ensure the terms that are selected as enriched by our algorithm are reasonable, we looked in detail at the enrichment analysis results. Most results were highly comparable and in line with what was expected. For some sets, all dictionaries seemed to perform well. For example, in our 'all neuron enriched sets' [16, 18]

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¹the results were an amalgamation of neuron related terms including mechanosensory ¹ ² neurons, thermosensitive neurons, interneurons, ganglions and male rays regardless ² ³ of the dictionary used. On the other hand, when we looked at a gene set enriched ³

⁴for germline precursor expression in the embryo [16], the dictionary with the 50⁴ ⁵cutoff was only able to identify 'oocyte WBbt:006797'; whereas the two smaller ⁵ ⁶dictionaries were able to single out cells germline precursor cells – at the 33-cutoff, ⁶ ⁷our tool identified 'Z2' and 'Z3' as being five-fold enriched; whereas at the 25 gene-⁷ ⁸cutoff the terms 'Psub4', 'Psub3' and 'Psub2' were identified in addition to 'Z2' and ⁸ ⁹'Z3'. We queried an embryonic stage intestine precursor geneset [16]. Notably, this⁹ ¹⁰gene set vielded no enrichment when using the 25 cutoff dictionary, nor when us-¹⁰ ¹¹ing the 50 cutoff dictionary. However, the 33 cutoff dictionary suggested, probably ¹¹ ¹²correctly, that the E lineage was heavily enriched in this set. Not all gueries worked ¹² ¹³equally well. For example, a number of intestinal enriched genes sets [16, 19] were ¹³ ¹⁴not enriched in intestine in any dictionary, but they were enriched for pharynx-¹⁴ ¹⁵ and hypodermis-related terms. We were somewhat surprised that intestinal gene ¹⁵ ¹⁶sets performed poorly, since the intestine is a relatively well-annotated tissue. We¹⁶ ¹⁷also assessed the internal agreement of our tool by using independent gene-sets ¹⁷ ¹⁸that we expected to be enriched in the same tissues. We had two independent pan-¹⁸ ¹⁹neuronal sets [16, 18]; two independent PVD enriched sets [16, 22]; two independent ¹⁹ ²⁰GABAergic gene sets [16, 17]; two independent pharvngeal gene sets [15, 16]; and ²⁰ ²¹two independent intestinal gene sets [16, 19]. Overall, the tool seems to have good²¹ ²²internal agreement. On most sets, the same terms were enriched, although order ²² ²³was somewhat variable. However, most high-scoring terms were preserved between ²³ ²⁴gene sets. The intestinal gene-sets and pharyngeal gene sets comparisons were ex-²⁴ ²⁵ceptions, since at least one gene set was missing each for intestine and pharynx²⁵ ²⁶in every dictionary, so we didn't consider them as informative for assessing inter-²⁶ ²⁷nal agreement. All comparisons can be found online in our Github repository (see²⁷ ²⁸Availability of data and materials). Overall, the dictionary generated by a 33 gene²⁸ ²⁹annotation cutoff with 0.95 redundancy threshold using the 'any' criterion. seemed ²⁹ ³⁰to perform well, with a good balance between specificity, verbosity and accuracy, ³⁰ ³¹so we selected this parameter set to generate our static dictionary. 32 32 33 33

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¹Results

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2 We applied our tool to the RNA-seq datasets developed by Engelmann et al. $\left[23\right]^2$
3 in order to attempt to gain further understanding of the biology underlying these 3
⁴ datasets. Engelmann et al. exposed young adult worms to 5 different pathogenic ⁴
⁵ bacteria or fungi for 24 hours, after which mRNA was extracted from the worms ⁵
$^6\mathrm{for}$ sequencing. We obtained the genes that Engelmann et al identified as up- or 6
7 down- regulated in their assay, and ran TEA using these lists. Initially we noticed 7
⁸ that genes that are down-regulated tend to be twice better annotated on aver-
⁹ age than genes that were up-regulated, suggesting that our understanding of the ⁹
$^{10}\mathrm{worm}$ immune system is scarce, in spite of important advances made over the last 10
¹¹ decade. Strikingly, 4 out of the five samples showed enrichment of neuronal tis- ¹¹
$^{12}\mathrm{sues}$ or neuronal precursor tissues (in the case of Harposporium sp) amongst the 12
$^{13}\mathrm{down\text{-}regulated}$ genes. A possible explanation for this might be that the infected 13
$^{14}\mathrm{worms}$ are sick and the neurons are beginning to shut down; an alternative hy- 14
$^{15}\mathrm{pothesis}$ would be that the worm is down-regulating specific neuronal pathways as 15
$^{16}\mathrm{a}$ behavioural response against the pathogen. Indeed, several studies[24, 25] have 16
$^{17}\mathrm{provided}$ evidence that $\mathit{C.~elegans}$ uses chemosensory neurons to identify pathogens 17
$^{18} \mathrm{Interestingly},$ one bacterium did not exhibit the same pattern of down-regulation of 18
19 neuronal-associated genes. $\it E.\ faecalis$ showed increased expression of genes associ- 19
$^{20}{\rm ated}$ with neuronal tissues, hinting that E. faecalis may have a different pathogenic 20
$^{21}\mathrm{profile}.$ Up-regulated tissues, when detected, included the hypodermis and excretory 21
$^{22}\mathrm{duct.}$ Our results highlight the involvement of various $\it C.~elegans$ neuronal tissues 22
in pathogen defense and/or illness.
24
²⁵ Discussion ²⁵
26 We have presented a tissue enrichment analysis tool that employs a standard hyper- 26
geometric model to test the $C.\ elegans$ tissue ontology. We have also presented the
28 first, to our knowledge, onto logy trimming algorithm. This algorithm, which is very 28
easy to execute, places strong limits on the number of terms selected for testing. ²⁹
Due to the nature of all ontologies as hierarchical, acyclical graphs with term in- 30
heritance, term annotations are correlated along any given branch. This correlation $^{\rm 31}$
32 reduces the benefits of including all terms for statistical analysis - for any given 32
33 term along a branch, if that term passes significance, there is a high probability 33

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¹that many other terms along that branch will also pass significant. If the branch ¹ ² is enriched by random chance, error propagation along a branch means that many ³more false positives will follow. Thus, a researcher might be misled by the number ³ ⁴of terms of correlated function and assign importance to this finding; the fact that ⁴ ⁵the branching structure of GO amplifies false positive signals is a powerful argu-⁵ ⁶ment for either reducing branch length or branch intracorrelation, or both. On the ⁶ ⁷other hand, if a term is actually enriched, we argue that there is little benefit to ⁷ ⁸presenting the user with additional terms along that branch. Instead, a user will⁸ ⁹benefit most from testing sparsely along the tree at a suitable specificity for hy-¹⁰pothesis formation. Related terms of the same level should only be tested when ¹⁰ ¹¹there is sufficient annotation to differentiate, with statistical confidence, whether ¹¹ ¹²one term is enriched above the other (see SI for a back-of-the-envelope calculation ¹² ¹³of when this can be the case). Our algorithm reduces branch length by identifying ¹³ ¹⁴ and removing nodes that are insufficiently annotated and parents that are likely to ¹⁴ ¹⁵include sparse information. It is important to note that our tool is not the first tissue enrichment model¹⁶ ¹⁷ for the worm that has been reported. Chikina et al [26] report a tissue enrichment ¹⁷ model based on an SVM classifier that has been trained on microarray studies. SVM 18 ¹⁹ classifiers are powerful tools capable of great sensitivity, but they require continuous retraining as tissue expression data widens. Our tool benefits from the fact that it 20 ²¹ will be integrated in WormBase and will therefore be updated continuously as new ²¹ $^{22}\mathrm{data}$ is integrated. 22 We have tried hard to benchmark our tool well. However, our analysis suffers from the drawback that is very hard to benchmark negative controls. Even for our set of positive controls, the statistical analysis sometimes throws out unexpected ²⁵ results. For example, the embryonic germline precursor gene set had the term 'AB' as the most enriched term in the dictionaries with cut off of 25 and 33. Is this an error, or does this hint at new biology? Although we were unable to determine false-positive and false-negative rates, we don't believe this should deter scientists from using our tool. Rather, we encourage researchers to use our tool carefully as a guide, integrating evidence from multiple sources to inform the most likely 31 hypotheses. As with any other tool based on statistical sampling, our analysis is most vulnerable to bias in the data collection stage. For example, we know that

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$^{1}\mathrm{tissue}$ expression reports are negatively biased against germline expression due to 1	L
2 the difficulty associated with extra chromosomal array expression in that tissue.	2
³ Support from the community will be crucial in correcting these flaws going forward;	3
⁴ indeed, without the community reports of tissue expression this tool would not be ⁴	1
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	6
⁷ Availability of data and materials	7
8Our web implementation is available at https://www.wormbase.org/tea. Our soft-s	3
9ware can also be downloaded using Python's pip installer via the command	9
pip install tissue_enrichment_tool	10
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noop, / damsoloos storatorio/ 1 isoto 2 in formion of the day of the	
All benchmark gene sets, benchmarking code and figures can also be found at the	13
14same address, under the 'tests' folder.	14
15	15
	16
The authors declare that they have no competing interests. 17	17
Author's contributions 18 DA and PWS conceived of the project; DA developed algorithm; RYL made intellectual contributions to the project; 1	18
PVI and IC developed the web CIII	19
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