Sma3s: A Three-Step Modular Annotator for Large Sequence Datasets

ANTONIO Muñoz-Mérida1, ENRIQUE Viguera2, M. GONZALO Claros3, OSWALDO Trelles1,4, and ANTONIO J. Pérez-Pulido5,*

Integrated Bioinformatics, National Institute for Bioinformatics, University of Málaga, Campus de Teatinos, Spain1; Cellular Biology, Genetics and Physiology Department, University of Málaga, Campus de Teatinos, Spain2; Molecular Biology and Biochemistry Department, University of Málaga, Campus de Teatinos, Spain3; Computer Architecture Department, University of Málaga, Campus de Teatinos, Spain4 and Centro Andaluz de Biología del Desarrollo (CABD, UPO-CSIC-JA), Facultad de Ciencias Experimentales (Área de Genética), Universidad Pablo de Olavide, Sevilla 41013, Spain5

*To whom correspondence should be addressed. Tel. +34 954-348-652. Fax. +34 954-349-376.
E-mail: ajperez@upo.es

Edited by Prof. Kenta Nakai
(Received 29 October 2013; accepted 6 January 2014)

Abstract

Automatic sequence annotation is an essential component of modern ‘omics’ studies, which aim to extract information from large collections of sequence data. Most existing tools use sequence homology to establish evolutionary relationships and assign putative functions to sequences. However, it can be difficult to define a similarity threshold that achieves sufficient coverage without sacrificing annotation quality. Defining the correct configuration is critical and can be challenging for non-specialist users. Thus, the development of robust automatic annotation techniques that generate high-quality annotations without needing expert knowledge would be very valuable for the research community. We present Sma3s, a tool for automatically annotating very large collections of biological sequences from any kind of gene library or genome. Sma3s is composed of three modules that progressively annotate query sequences using either: (i) very similar homologues, (ii) orthologous sequences or (iii) terms enriched in groups of homologous sequences. We trained the system using several random sets of known sequences, demonstrating average sensitivity and specificity values of ~85%. In conclusion, Sma3s is a versatile tool for high-throughput annotation of a wide variety of sequence datasets that outperforms the accuracy of other well-established annotation algorithms, and it can enrich existing database annotations and uncover previously hidden features. Importantly, Sma3s has already been used in the functional annotation of two published transcriptomes.

Key words: functional annotation; genome annotation; transcriptome annotation; bioinformatic tool

1. Introduction

Sequence annotation is the process of associating biological information to sequences of interest. Annotations can include the potential function, cellular localization, biological process or protein structure of a given sequence. Some sequences are annotated using direct experimental evidence, but most annotations are inferred from sequence similarities or conserved patterns associated with known characteristics. Large publically accessible databases of annotated sequences make it possible to automatically annotate large collections of unknown sequences. This is especially valuable for the interpretation of large sequence datasets generated by genome and expressed sequence tag (EST) sequencing projects as well as gene and protein expression experiments, such as DNA microarrays, and many other emerging research areas.

Sequence annotation is also important in transcriptomic experiments that aim to identify gene clusters with similar expression patterns that are linked to a particular biological process or experimental condition. Biological function can then be inferred from annotations shared within these clusters.
Several methods have been developed for the annotation of the large datasets generated by sequencing projects. These methods typically perform homology-based searches and infer annotations from sequences with high levels of homology to the query sequence. To detect sequence similarity, most methods use BLAST, a fast heuristic algorithm for identifying homology between sequences based on local sequence alignments.8

Blast2GO is one of the most widely used tools for genomic annotation.9 It performs three steps: a BLAST search, a step that maps similar sequences to existing annotation associations, and finally term annotation. As its name suggests, Blast2GO mainly uses terms from the Gene Ontology (GO) controlled vocabulary for annotation,10 although it also includes other annotation classes. However, Blast2GO is most commonly run online, which can make it more difficult to analyse large collections of sequences. Other methods can be installed locally, and so avoid problems associated with remote execution, such as sequence number limitations or resource thresholds. Blast2GO is also available in an offline form, but installation requires knowledge of relational databases that make it less suitable for routine use and difficult for non-specialist users. AutoFact11 can perform several types of BLAST search using locally downloaded domain and motif databases. However, it uses databases with descriptive non-standard annotation entries rather than controlled vocabularies such as GO. The resulting lack of homogeneity in terms of vocabulary or syntax can make it much more difficult to automatically assign annotations or evaluate annotation quality.

The above methods can be applied to a wide range of different sequence types and organisms. Many other tools are specialized for specific applications. Some tools only deal with single annotation classes, as is the case for Gotcha, which only uses GO terms.12 Others are specialized for the annotation of specific sequence types, such as the EST-specific ESTAnnotator13 and EST-PAC.14 In addition, some tools are designed especially for the annotation of specific organisms. For example, BLANNANOTATOR assigns non-standard annotations to bacterial sequences.15

Some platforms allow diverse data sources to be integrated for the purposes of sequence annotation and annotation analysis. For example, the Babelomics suite,16 of which Blast2GO is part, contains tools for assigning interactions, pathways or even regulatory annotations to analysed sequences. Another widely used platform is DAVID,17 which extracts functional annotations from a variety of public genome resources, and allows subsequent analysis by biological enrichment. One of the richest annotation sources is BioMart,18 which retrieves sequence annotations from Ensembl and other databases.

All of these annotation methods use BLAST to detect and evaluate the similarity between a query sequence and putative homologues.8 For each query sequence, BLAST attempts to create alignments with database sequences and uses these alignments to judge sequence similarity. Most annotation methods select the database sequence with the highest overall similarity score for each query sequence as the donor of annotation information. In addition, a minimum similarity threshold sets the limit below which sequences will be rejected even if they are the most similar sequence in the database. Defining the similarity threshold is not an easy task, as it must balance annotation quality and sequence coverage (SC). Moreover, it has been shown that the optimal similarity threshold can vary between different sequence types.19 Several other BLAST variables, such as how it scores alignment gaps, can also result in significantly different results. These factors can be difficult to configure, particularly for non-specialist users.

We have developed Sma3s, a fast, accurate and flexible annotation tool specifically designed for the annotation of large collections of sequences obtained from diverse gene libraries or coding sequence datasets. It is comprised of three modules that solve the annotation process with progressively less-stringent sequence similarity requirements, combined with methods to optimize specificity. Each module uses an initial exhaustive BLAST search as its starting point. The third Sma3s module enhances annotation quality using term enrichment to identify annotations shared by groups of similar sequences. We have defined optimal default Sma3s parameter values to minimize user intervention for most applications, thus aiding consistency and comparability. We show that Sma3s can rapidly produce high levels of prediction accuracy with minimal human supervision and modest computational resources.

2. Materials and methods

2.1. Implementation

Sma3s was written in Perl using modules of the BioPerl 1.6 project (http://www.bioperl.org) and is freely accessible from the Sma3s website (http://www.bioinfocabd.upo.es/sma3s/). Sma3s uses two programmes from the Blast package: blastp for amino acid sequences and blastx for nucleotide sequences, using an E-value threshold of $10^{-6}$, and blastclust for sequence clustering procedures. A Perl biological enrichment module, based on hypergeometric distribution, was used to calculate probability values (http://www.cse.huji.ac.il/course/2006/bioskill/Ex2/HyGe.pm).

Sma3s uses the UniProt database in plain-text format files (.file.dat), corresponding to the taxonomic division
of the organism under study, as its main data source. More details about programme requirements and operation are available on the website.

Running on a dual-processor Intel Pentium 4 CPU 3.00 GHz with 4 Gb of RAM, the programme takes an average of 0.7 s to annotate each sequence (mean length = 400 amino acids) against a division of the Swiss-Prot database (~45 000 sequences), using the three modules. In fact, the most time-consuming process (~90%) is the initial exhaustive BLAST (blastp or blastx), whose results are then reused in subsequent parameter tuning and annotation refinement steps.

2.2. Annotation types

UniProt\textsuperscript{20} is arguably the most complete publically available protein database. Human-based curation ensures high-quality annotations, which is particularly evident in the Swiss-Prot section, compared with the automatically annotated TrEMBL section (also part of UniProt). Sma3s uses Swiss-Prot as its main source for extracting annotations. The UniProt fields used by Sma3s are:

\textit{Gene ontology} GO\textsuperscript{10} provides a controlled vocabulary to describe genes and gene product attributes. It is organized into three biological ontologies: molecular function, biological process and cellular component. Standardized GO term annotations are included in the cross-reference (DR) field of UniProt.

\textit{Interpro} InterPro\textsuperscript{21} is an integrated documentation resource of protein families, domains and sites. It combines complementary sequence pattern information from several databases. UniProt provides InterPro identifiers also from the DR field.

\textit{Swiss-Prot keywords} The Swiss-Prot keywords constitute a well-defined and controlled vocabulary of terms used to annotate a UniProt protein entry. These keywords (the KW field in UniProt) describe functions, biological processes, structure, cellular localization and other protein characteristics.

\textit{Pathway annotation} This annotation provides a description of the metabolic pathway(s) in which a protein is involved. It is obtained from the comment (CC) field containing generic and specific metabolic pathway descriptors (e.g.: -!- PATHWAY: Nucleotide metabolism; purine metabolism). Sma3s gathers annotations of the most generic level (‘Nucleotide metabolism’ in the previous example). This type of annotation is particularly useful for identifying co-expressed genes that are active in the same metabolic pathway.

New annotation types can be incorporated into Sma3s with only minor changes to its algorithms.

2.3. Threshold calculation for the selection of multiple homologous sequences

Module 3 uses a modified version of Sander's formula,\textsuperscript{22} later updated by Rost\textsuperscript{23} to select alignments based on both identity and alignment length. The original formulas describe the relationship between sequence identity and alignment length observed in sequences sharing structural similarities as a curve. In the Rost equation, the identity threshold \((p)\) for an alignment of length \(L\) is defined as:

\[
p^\prime(n) = n + 480 \times L^{-0.32 \times (1 + e^{-1/100})},
\]

where \(n\) describes the distance in percentage from the original curve. This value allows curve adjustment to increase stringency and to ensure at least 40\% identity \((n = 20)\) at any alignment length in accordance with the ~40\% threshold for functional conservation described by Wilson \textit{et al.}\textsuperscript{24} and will be the default value used for Module 3 annotation.

2.4. Statistical significance analysis of annotations

Each term assigned by Sma3s' prediction algorithm is assigned a \(P\)-value to indicate annotation quality. Annotations from Module 3 take \(P\)-values from a biological enrichment calculation, while those from Modules 1 and 2 are derived from the highest rated BLAST hit.

The biological enrichment algorithm (see below) is based on hypergeometric distribution as follows: from \(M\) sequences in the database, \(K\) contains annotation \(X\); with \(m\) sequences in the annotated cluster, the probability \(P(X = k)\) that \(k\) sequences will have annotation \(X\) is:

\[
P(X = k) = \frac{\binom{K}{k} \binom{M - K}{m - k}}{\binom{M}{m}}
\]

Assuming that \(l\) sequences were found with annotation \(X\), the probability \(P\) that \(l\) can be adjusted to the null hypothesis is:

\[
P = \frac{\min(k, m)}{l} P(X = k),
\]

The cut-off \(P\)-value was determined experimentally to be \(\leq 0.1\) (see the Results section).

2.5. Accuracy estimation

Sma3s assesses three different ratios to evaluate method accuracy taking into account test sets of previously annotated sequences. These ratios are based on TP (true positives: predicted terms corresponding to the real annotation), FP (false positives: predicted
terms not corresponding to the real annotation) and FN (false negatives: non-predicted real terms).

It is worth pointing out that \( (TP + FN) \) corresponds to the number of real annotations and \( (TP + FP) \) represents the number of predicted annotations. In this way, we can calculate both sensitivity and specificity indices. We can also calculate the Relative Error Quotient (REQ), which evaluates the overall prediction quality by considering both the sensitivity and specificity measurements as described in the GOtcha method. \(^{12}\) REQ is calculated as:

\[
REQ = \frac{FN \times W + FP}{TP \times (1 + W)},
\]

where \( W \) is a weighting factor that moderates the influence of FN and FP. In this work, \( W \) has been set to 1 to make results comparable with the literature. Low REQ values indicate a low annotation error rate, whereas a high REQ indicates a higher proportion of errors. This measure of accuracy has the advantage that it encompasses TP, FP and FN, thus combining sensitivity and selectivity in a single value.

To evaluate prediction coverage percentages, two more values were calculated: the SC (the number of sequences with predicted annotations with respect to sequences with known annotations) and term coverage (TC, the number of predicted annotations with respect to known annotations).

### 2.6. Test datasets

#### 2.6.1. Random sequence datasets

All the databases used in this work belong to the UniProt Knowledgebase Release 2011_08, which were obtained from the EBI ftp-site: ftp://ftp.ebi.ac.uk/pub/databases/uniprot/current_release/knowledgebase/taxonomic_divisions/.

We downloaded four taxonomic divisions (bacteria, plants, invertebrates and mammals) and built five independent random datasets of 1 000 sequences (5 000 in total) for each taxonomic division. These datasets were processed with Sma3s. Query sequences were removed from the database before each run to avoid self-annotation.

#### 2.6.2. Sequence sets from DNA arrays

We extracted two independent DNA arrays from the GEO database: \(^{25}\) Affymetrix Murine 11K SubB Array (AC:GPL76) and the Affymetrix Arabidopsis ATH1 Genome Array (AC:GPL198). The nucleotide sequences from these arrays were downloaded, and sequences shorter than 150 were removed (to ensure a minimum amino acid sequence length of 50). Reference annotations were taken from both Affymetrix-assigned GO terms and GO terms extracted from the Swiss-Prot entries referred to in the Affymetrix annotation.

### 2.7. Blast2GO and Top-BLAST tests

We opened Blast2GO as a Java application (http://www.blast2go.com/b2ghome), selecting the 2Gb memory version, as would be typical for non-expert users performing a search with default parameters. The test did not include graphical data mining or enrichment analysis Blast2GO options. For Top-BLAST annotation, the best hit from a default BLAST search was used to annotate every sequence in the dataset. Self-prediction in Blast2GO (annotation based on annotations from the query sequence) was unavoidable, since dataset sequences cannot be removed from the database.

### 3. Results

#### 3.1. Sma3s algorithm overview

With Sma3s, we set out to develop a tool that can intelligently assign annotations to query sequences in a way that maximizes accuracy without sacrificing coverage, and works in a way that is robust enough to generate high-quality results without having to adjust the parameters.

To achieve these aims, we designed Sma3s as a modular system that processes query sequences in a series of discrete steps. The method starts with a BLAST similarity search, followed by a three-step annotation process which sequentially tests different degrees of similarity between query and database sequences (Fig. 1). The first module annotates query sequences using database homologues with very high levels of similarity (Fig. 1, M1). The second module performs a reciprocal BLAST on sequences unannotated in the first step to generate annotations from orthologous database sequences (Fig. 1, M2). The third module uses a novel strategy (Fig. 1, M3) that looks for more distantly related sequences and analyses these data for statistically significant sequence annotations.

Module 1: The first annotator uses a Top-BLAST strategy to check whether the query sequence (or one very similar to it) already exists in the UniProt database, and then directly assigns database annotations to the query sequence. We established a minimum sequence identity of 90% and a sequence overlap of at least 90% of the database sequence, although these parameters are customizable. In all cases, Sma3s uses annotations from Swiss-Prot in preference to TrEMBL when identical amino acid sequences are found in both databases (Table 1, see the next section for details).

Module 2: This module annotates using information from putative orthologues with lower similarity. The second module of Sma3s is based on reciprocal BLAST searches. It detects lower similarity sequences with at least 75% of per-residue identity, covering at least 75% of database sequence length. The best hit is then
used as an input seed in a blastp or tblastn search against the initial set of query sequences. If this second BLAST identifies the starting query sequence as the best hit, then an orthology relationship is established, and consequently, the annotation recovered. As in the previous step, Sma3s uses Swiss-Prot annotations in preference to TrEMBL (Table 1, see the next section for details).

Module 3: Module 3 recovers annotation information from multiple homologous sequences using a novel multi-step approach. The first step is to determine which homologues have sufficient similarity to the query sequence. Work by Sander\textsuperscript{22} and Rost\textsuperscript{23} found that minimum alignment length was a better criterion than overall percentage sequence identity for estimating sequence and structural homology, especially when comparing pairs of structurally matched proteins with lower levels of similarity. Intuitively, for short sequence alignments, a high percentage identity is needed to establish statistically significant relationships. Conversely, long pairwise alignments require lower identity to be qualified as significant. In the context of protein structure, a minimum threshold of \(~20\%\) was found to be a good predictor of protein structure, provided that the alignment had a length of at least 150 residues. Function is less well conserved than structure, but further studies have reported that biological function is typically conserved when two sequences exhibit 40\% sequence identity.\textsuperscript{24} Thus, based on the hypothesis that function conservation is also related to sequence alignment length, we have used a modified form of Rost’s equation (see Materials and Methods for details) that only selects sequences whose alignments have at least 40\% identity at any alignment length.

Although the sequences selected by this method can have significantly lower similarities to those identified by Modules 1 and 2, the combined analyses of multiple sequences can be used to increase annotation reliability. This is based on the hypothesis that annotations shared by several homologous sequences are more likely to reflect functions shared by the query sequence. Module 3 uses biological enrichment to only select those terms which appear more frequently in the identified homologues than would be expected by chance in the source database (see ‘Statistical significance assessment of annotations’ in the Materials and Methods section for details).

The presence of several redundant sequences in the search results could bias the term enrichment

**Table 1.** Sma3s results with different source databases

<table>
<thead>
<tr>
<th>Source Database</th>
<th>Sn</th>
<th>Sp</th>
<th>REQ</th>
<th>SC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Swiss-Prot</td>
<td>0.83</td>
<td>0.87</td>
<td>0.19</td>
<td>0.87</td>
</tr>
<tr>
<td>TrEMBL</td>
<td>0.61</td>
<td>0.86</td>
<td>0.45</td>
<td>0.94</td>
</tr>
<tr>
<td>UniProt</td>
<td>0.68</td>
<td>0.87</td>
<td>0.32</td>
<td>0.94</td>
</tr>
</tbody>
</table>

Sequence coverage (SC), specificity (Sp), sensitivity (Sn) and REQ values are shown together with the corresponding standard deviation.
algorithm. To reduce this possibility, very similar sequences are first combined by ‘clustering’. Redundant sequences are identified using blastclust (part of the Blast package), using 95% identity and alignment length as search criteria. Thus, when evaluating the frequency of annotation terms, each cluster counts only once, regardless of the number of sequences it contains.

In summary, Module 3 processes each query as follows (also see Fig. 1, M3): (i) the BLAST report is scanned for statistically significant alignments according to Rost’s equation; (ii) identification and grouping of clusters containing very similar members of the same family; (iii) a preliminary annotation set is formed by combining individual non-redundant annotations from each cluster and finally, (iv) biological term enrichment is used to select significantly overrepresented annotations in each group of homologous sequences. Owing to the modular implementation of the algorithm, each module can be run independently, allowing the user to easily customize annotation criteria.

3.2. Sma3s annotates random Swiss-Prot sequence collections with high accuracy

We trained Sma3s using random datasets extracted from the four Swiss-Prot taxonomic divisions. Five independent random sets of 1000 sequences (5000 in total) were extracted for each division. These sequences were then annotated using the corresponding division from Swiss-Prot, and each query sequence removed from the database at each step to avoid self-annotation.

We evaluated the annotation of each random set of sequences for SC, specificity, sensitivity and REQ values (an accuracy measurement which combines both specificity and sensitivity, with 0 being the best possible value). The results obtained varied significantly between taxonomic divisions, but not between different datasets from the same division (Fig. 2).

To identify the optimal default cut-off value, we tested Sma3s with different P-values using our random datasets as query sequences and the corresponding taxonomic divisions of the Swiss-Prot database as annotation sources. As can be seen in Fig. 2, more restrictive P-values generated consistent results with low REQ values and higher specificity (Sp) but at the cost of reduced sensitivity (Sn). The highest accuracy, as defined by the lowest REQ values, was obtained when specificity and sensitivity had equal or similar values. With the exception of the invertebrate dataset, a P-value of 0.1 resulted in average sensitivity and specificity values >0.8. These values compare favourably with those of other tools, such as Blast2GO \(^9\) or Gotcha,\(^{12}\) whose accuracy typically ranges between 0.65 and 0.7. On this basis, we established a default P-value of 0.1 for Sma3s.

When the same test datasets were annotated using the uncurated TrEMBL database or the complete UniProt database instead of Swiss-Prot, the average accuracy was reduced despite the high coverage obtained (Table 1). This can be explained by the fact that the unreviewed TrEMBL sequence collection is much larger than the manually annotated Swiss-Prot set. Thus, a large proportion of annotations assigned from the UniProt database are derived from TrEMBL, even though UniProt also contains Swiss-Prot sequences.

Accuracy was also calculated separately for each of the four annotation types recovered by Sma3s: GO terms, SW-Keywords, InterPro and Pathways (Fig. 3). Best results were obtained for Pathway and InterPro annotations, which had REQ values \(\leq 0.2\) for all except the invertebrate dataset. Keyword and GO annotations had poorer REQ values, but still achieved average sensitivity and specificity values of 0.8 (and as high as 0.97 for bacteria). The exception, again, was the invertebrate dataset for which Sma3s generated higher REQ values, associated with significantly lower sensitivity. The relatively poor results for the invertebrate dataset are probably due to the relatively small number of sequences present in the corresponding invertebrate database. Sequence length is also important. In fact, this dataset is the group with the highest proportion of very short sequences (10-40 amino acids), which are usually annotated with difficulty (Supplementary Fig. S1a).

3.3. The M3 module can enrich the annotation of previously analysed sequences

As described above, each dataset sequence is annotated by one of the three different modules in the Sma3s algorithm, which sequentially run from M1 to M3. To determine the effectiveness of each module, we calculated the percentage of random dataset sequences annotated by each module. On average, the M1 module annotated 53.76% of sequences (Table 2) with M2 and M3 annotating the remainder (19.70 and 26.54%, respectively). As modules M2 and M3 only annotate query sequences that were not annotated by the previous module(s), the \(\sim 26\%\) of sequences annotated by M3 suggests that it significantly increases Sma3s annotation coverage.

M1 and M2 modules provide highly accurate results since annotations are obtained from verified orthologues with high degrees of identity between query and donor sequences. However, as both modules employ the traditional Top-BLAST approach, each query sequence is annotated from only one putative orthologue. In contrast, the M3 module typically extracts annotations from several homologous sequences identified in the BLAST report. To better compare the relative benefits of the different approaches used by Sma3s, we compared the annotation of our random datasets
using the three modules as described above (M123) against annotation using Module 3 alone (M3). In the latter case, all sequences are annotated by Module 3, not just the sequences unannotated by Modules 1 and 2. Figure 4 shows that the mean specificity and sensitivity of the M3 module were slightly lower than the full Sma3s analysis, with the REQ value increasing from 0.19 to 0.32. However, M3 obtained more annotations (Fig. 4, column TC) due to its ability to extract annotations from multiple homologues. M3 accuracy was also compared with combined M1 and M2 modules (M12), which represent more conventional annotation algorithms. M12 obtained higher specificity (0.97) since it annotates only using very similar sequences (Fig. 4). However, sensitivity is rather less (0.47) with only 64% of dataset sequences annotated. This results in an REQ value of 0.71 for M12 versus the 0.32 for M3. Several examples illustrate this additional advantage of Sma3s M3 as a standalone annotation tool.

The best BLAST hit of the pig Qil1 protein (QIL1_PIG) in the mammalian dataset is the primate homologue (QIL1_MACFA). Because this protein entry does not contain any of the four annotation types used by Sma3s, pig Qil1 was not annotated by M1. However, the M3 module was able to annotate pig Qil1 localization using the GO term 'Mitochondrion', obtained from a more distant Qil1 mouse homologue (QIL1_MOUSE).

The Sma3s M3 module could be used even in cases where incomplete annotations are present in Swiss-Prot. For example, when M1 or M2 Sma3s modules are used with invertebrate datasets, the U3-lyctoxin-Ls1a protein from the wolf spider (TX308_LYCSI) only generates the GO term 'extracellular region'. In contrast,
the M3 module annotates the sequence with two additional GO terms: ‘pathogenesis’ and ‘calcium channel inhibitor activity’. Since this protein belongs to the spider toxin CSTX superfamily, which blocks mammalian neuronal calcium channels, these terms should be present in the annotation of this protein.
Another example is the SfsA protein (SFSA_PSEF5) from *Pseudomonas fluorescens*. SfsA was annotated in bacterial datasets with the Swiss-Prot keyword ‘Complete proteome’ and no GO terms. However, M3 predicted the following Keywords and GO terms: DNA binding and carbohydrate metabolic process. These terms seem valid because the bacterial SfsA family includes sugar fermentation stimulation proteins that contain a helix-turn-helix motif that probably binds DNA at its C-terminus.27

3.4. Accurate annotation of DNA array sequence sets

To test Sma3s with an experimental set of sequences, we chose collections belonging to two different Affymetrix DNA arrays (one mouse and one *Arabidopsis*). The real annotations for these datasets were extracted from the Affymetrix web server (March, 2012) as GO terms linked to the sequences, or as GO terms extracted from Swiss-Prot IDs linked to the sequences. Sma3s was used to annotate these arrays using default parameters, and accuracy measured using the reference annotations. We compared these results with the traditional Top-BLAST method, and the widely used Blast2GO tool with default annotation settings (Fig. 5).

### Table 2. Number of annotations assigned by each Sma3s module

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Module</th>
<th>Number of annotations</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random</td>
<td>M1</td>
<td>468.45 ± 164.25</td>
<td>53.76</td>
</tr>
<tr>
<td></td>
<td>M2</td>
<td>171.70 ± 45.12</td>
<td>19.70</td>
</tr>
<tr>
<td></td>
<td>M3</td>
<td>231.30 ± 80.72</td>
<td>26.54</td>
</tr>
<tr>
<td>DNA arrays</td>
<td>M1</td>
<td>386.5 ± 369.82</td>
<td>5.23</td>
</tr>
<tr>
<td></td>
<td>M2</td>
<td>462.5 ± 556.49</td>
<td>6.26</td>
</tr>
<tr>
<td></td>
<td>M3</td>
<td>6536.5 ± 4531.85</td>
<td>88.50</td>
</tr>
</tbody>
</table>

The average number of annotations identified by Sma3s is shown for the random and DNA arrays datasets, together with the corresponding standard deviation. Note that the percentages for each dataset group sum 100%, since Sma3s has applied its three modules consecutively (default configuration).

### Figure 4. Comparison of Sma3s performance using all modules (M123) or the M3 module alone. Corresponding sensitivity (Sn), specificity (Sp), sequence coverage (SC) and term coverage (TC) values appear above the bars. This figure appears in colour in the online version of *DNA Research.*

3.5. Sma3s is fast and has low-memory usage

In the post-genomic era, it is critical to have annotation tools that can be easily integrated into sequence analysis pipelines and applied quickly to large datasets. Therefore, it is important that annotation analyses run efficiently without excessive resource demands or sequence limitations. To this end, we estimated the speed of Sma3s analysis using the modest computer hardware (see Materials and Methods for details). We tested run times and memory usage using three datasets with progressively increasing sizes (Fig. 6). Measurements were taken separately for the BLAST search and sequence annotation steps. Sma3s processing times increased in

### Table 2. Number of annotations assigned by each Sma3s module

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Module</th>
<th>Number of annotations</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random</td>
<td>M1</td>
<td>468.45 ± 164.25</td>
<td>53.76</td>
</tr>
<tr>
<td></td>
<td>M2</td>
<td>171.70 ± 45.12</td>
<td>19.70</td>
</tr>
<tr>
<td></td>
<td>M3</td>
<td>231.30 ± 80.72</td>
<td>26.54</td>
</tr>
<tr>
<td>DNA arrays</td>
<td>M1</td>
<td>386.5 ± 369.82</td>
<td>5.23</td>
</tr>
<tr>
<td></td>
<td>M2</td>
<td>462.5 ± 556.49</td>
<td>6.26</td>
</tr>
<tr>
<td></td>
<td>M3</td>
<td>6536.5 ± 4531.85</td>
<td>88.50</td>
</tr>
</tbody>
</table>

The average number of annotations identified by Sma3s is shown for the random and DNA arrays datasets, together with the corresponding standard deviation. Note that the percentages for each dataset group sum 100%, since Sma3s has applied its three modules consecutively (default configuration).
proportion to the number of sequences analysed (Fig. 6a). Times were highly dependent on the initial BLAST step which increased in proportion to the sequence number. However, Sma3s annotation times were much lower and barely affected by increasing dataset sizes. Significantly, Sma3s run times were two orders of magnitude shorter than those of the Blast2GO method.

Memory usage was also measured using the same datasets and methods (Fig. 6b). Sma3s had an essentially constant memory usage in its two steps despite the sequence number increase, in contrast to the Blast2Go method. Our results show that Sma3s is able to efficiently annotate large sequence datasets using modest computational resources.

4. Discussion

Sequencing projects produce hundreds of thousands of new biological sequences that need to be quickly and accurately annotated since this information constitutes the basis for subsequent analysis, such as pathway design or genomics and transcriptomics studies. Therefore, fast and user friendly methods are needed to support sequence annotation at this stage.

Sma3s uses three consecutive modules: M1 (which searches for very similar sequences in the database), M2 (which searches for orthologues) and M3 (which extracts information from several homologous sequences). The novel approach used by the M3 module selects several homologues for each query sequence, using a modified Rost equation, which takes into account both alignment length and percentage identity. Annotations from all the non-redundant sequences are pooled, but only annotations that are significantly enriched versus the expected background frequency are assigned to the query sequence. This has the advantage that useful annotation information present in several homologues can be extracted, rather than only from the sequence with the highest homology.

Our tests have shown that Sma3s is a fast annotation method requiring minimal human supervision and computational resources. Mean Sma3s accuracy using different annotation types (GO, Swiss-Prot keywords and pathways, and InterPro) was higher than 0.8 (except for the invertebrate dataset), and was even higher (0.9) for specific datasets such as bacteria. In addition, REQ values were consistently very low. InterPro annotation was highly specific and sensitive (see Fig. 3), which can be explained by the fact that current InterPro annotation is strongly focused on sequence similarity, which has been shown to be effective for protein function prediction 26. On the other hand, it is difficult to judge Sma3s Swiss-Prot Pathway annotation given the limited coverage of this category in the UniProt database. Importantly, both InterPro and Pathway use short and controlled vocabularies with low term diversification. Some annotation methods can assign sentences derived from the database description line 15. Sma3s can also assign this descriptor (data not shown). However, the accuracy of these non-standard annotations...
is difficult to measure, whereas controlled terms such as GO or Keywords are more useful for experiments using massive datasets. Sma3s can be easily adapted to use different annotation classes, and each term predicted by Sma3s can be assigned a corresponding probability value. For example, Sma3s also includes annotations for the most probable gene names for each annotated sequence (data not shown), which can be useful for non-specialized users who only want to rapidly identify their sequences. Furthermore, Sma3s could be used with other annotation sources since the annotation assignment step can easily be disengaged from the main algorithm and used with other databases. For example, the annotation component could be extended to use data from the BioMart project, which links annotations from heterogeneous databases to UniProt identifiers. If Sma3s was converted to a web service, it could be directed to use BioMart public web services as an annotation source. This would also make Sma3s more versatile by dividing Blast and annotation steps into separate services.

4.1. The reference dataset and the BLAST results influence annotation accuracy

We demonstrated the accuracy of Sma3s with tests using different random datasets (Fig. 2). Moreover, Sma3s outperformed other widely used annotation methods when annotating DNA array datasets (Fig. 5). The only exception was the mouse dataset using Swiss-Prot as a reference. The annotation accuracy of this dataset is strongly dependent on reference annotations, which in this case came from Affymetrix. These annotations are assigned using NetAffx, which is based on homology transfer from heterogeneous databases, including Swiss-Prot, but also Unigene and LocusLink. Furthermore, these annotations depend on the external database entries associated with individual Affymetrix sequences, which have been shown to contain frequent inaccuracies. Thus, selection of a suitably high-quality reference annotation set seems to be an important factor for testing this or any other method. Nevertheless, Sma3s produced the best results in virtually all cases.

Further evidence for the idea that database annotation influences annotation accuracy is presented in Table 1, where the well-annotated Swiss-Prot division of UniProt provided the best accuracy (including specificity and sensitivity), while TrEMBL provided the lowest values. UniProt is the most widely used protein database since it contains rich information on each sequence. UniProt is comprised of Swiss-Prot, a curated section reviewed by skilled annotators and TrEMBL, an unreviewed section containing additional sequences that can be moved to Swiss-Prot once they have been manually reviewed. We have found that the best annotation accuracy is obtained when Swiss-Prot is used as the sole reference database (Table 1). Conversely, accuracy was reduced when using either TrEMBL or the complete UniProt database, mainly due to lower sensitivity. These lower values are expected partly because the test datasets were extracted from Swiss-Prot, and partly because TrEMBL is a much larger database. Thus, when UniProt or TrEMBL are used as reference sources, a higher proportion of Sma3s annotations are TrEMBL-derived, which consequently lowers the average annotation accuracy.

Similarly, it is important to highlight that results were also dependent on the quality of annotated reference sequences for each taxonomic group. For example, Sma3s annotation of random invertebrate sequences was of a much lower quality than those of bacterial sequences. The main difference between the two UniProt datasets is in the number of sequences they contain: 24,050 invertebrate sequences versus 326,570 bacterial sequences. In order for Module 3 to assign an annotation, it has to be statistically enriched within the homologous sequences. The probability of finding enriched annotations is more likely
with the higher numbers of candidate sequences available in the bacterial dataset. In fact, we have found that a combination of TrEMBL and SWISS-PROT databases increases sensitivity when annotating weakly conserved genomes (unpublished observation). However, this approach results in a higher rate of false positives.

Sequence length may be an important factor for successful annotation. For example, short sequences occur most frequently in invertebrates and to a lesser extent in plants (Supplementary Fig. S1a). These sequences are rarely annotated due to the low alignment scores they produce. The M1 module, which requires very high levels of similarity, is typically the most effective Sma3s component for annotation of these sequences (Supplementary Fig. S1b). This effect is likely to be enhanced by the higher diversity of the invertebrate dataset versus the bacterial one.

Another key factor for successful annotation is the initial BLAST search. In order for a sequence to be annotated by any of the three Sma3s modules, the initial BLAST must identify sequence alignments. For example, of the 30% of mouse array sequences which were not annotated by Sma3s (SC = 0.7 in Fig. 5), 86% lacked BLAST results, making annotation impossible. In the case of Arabidopsis array sequences, 40% lacked BLAST results thus limiting annotation coverage to a maximum of 60%. This dependency on BLAST results is shared with most annotation methods, including AutoFact 11, Blast2GO 9 and GOrtcha 12. In the case of Sma3s, the effect of the BLAST search is especially important given that Sma3s was able to annotate 90% of array sequences with BLAST alignments, largely due to the novel M3 module (Table 2).

4.2. The combined use of the three Sma3s modules generates accurate annotation, with enhanced coverage from the M3 module

We have shown that Sma3s can annotate different dataset types using its three module structure. Modules M1 and M2 are able to identify closely related database sequences and extract specific annotations, albeit with low overall coverage (TC in Figs 4 and 5). Thus, modules M1 and M2 provide high-quality annotations (with lower REQ values), increasing the overall sensitivity and specificity of Sma3s (Sn and Sp in Fig. 4).

Our findings suggest that the M3 module can increase SC in some cases, especially where close orthologues lack database annotation. The initial BLAST result can be reused by the M3 module to obtain new annotations for sequences previously annotated by M1 or M2. M3 annotation is broader, allowing new terms to be found (Fig. 4) but with lower specificity, thus increasing the chance of generating false positives. For most applications, using the three Sma3s modules sequentially balance the higher accuracy of Modules 1 and 2 with the identification of useful annotations from Module 3 for otherwise unannotated sequences.

Our results demonstrate that the Sma3s M3 module is able to generate new annotations for sequences that were unannotated in their original databases. Thus, the M3 module may be valuable as a database curation tool, adding new annotations to existing public databases. However, it is important to take into account that this module is likely to generate annotations with lower specificity (Sp in Figs 4 and 5). Ideally, users would be able to choose between high sensitivity and high specificity according to the goal of the experiment.

4.3. Sma3s has low computing requirements

The Sma3s algorithm comprises two different basic steps: similarity searching by BLAST and annotation by different strategies in each of the three modules. For high-throughput analyses, the BLAST itself accounts for most of the calculation time. Result execution times (Fig. 6) demonstrate that parallel BLAST searches could accelerate the process without affecting the rest of the algorithm. In fact, both Sma3s and Blast2GO can use separately generated BLAST reports, which may allow accelerated analysis if the user was to execute similarity searches in parallel. However, the speed advantage of Sma3s over Blast2GO is due to more than the speed of the BLAST search. Another reason for the speed of Sma3s is that it uses SwissProt, a relatively small high-quality database, which itself increases annotation accuracy (Figs 2, 3 and 5). These results together allow for a faster and more accurate annotation process when analysing large datasets. Thus, Sma3s can annotate a set of 20 000 sequences in ~3 h, without human intervention. Finally, memory usage is low (Fig. 6b), alleviating the need for high-performance computing facilities.

In conclusion, Sma3s is a fast and accurate method for annotating massive nucleotide or amino acid datasets in a way that is readily accessible for non-expert users. In fact, Sma3s has already been used in the functional annotation of both the olive29 and the maritime pine30 transcriptomes, and is now being used for the annotation of both a new bacterial genome and an EST database of almost 300 plant datasets containing a total of 8.6 million sequences (data not published yet).

Authors’ contributions

A.M.M. significantly helped in the design computational implementation, and testing of Sma3s algorithm. E.V. and M.G.C. contributed to computational design and rationale. O.T. managed and coordinated the initial stages, including the training. A.J.P.P. conceived the study, carried out the design, the tests and trainings, and wrote the computer program and the manuscript.
Acknowledgement: We thank John Pearson for his critical reading of the manuscript.

Supplementary data: Supplementary data are available at www.dnaresearch.oxfordjournals.org.

Funding

This work has been partially financed by the National Institute for Bioinformatics (www.inab.org), a platform of Genoma España and the EC project ‘Advancing Clinico-Genomic Trials on Cancer’ (contract no. 026996).

References

23. Rost, B. 1999, Twilight zone of protein sequence alignments, Protein Eng., 12, 85–94.