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Benchmarking mass spectrometry based proteomics algorithms using a simulated database

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Abstract

Protein sequencing algorithms process data from a variety of instruments that has been generated under diverse experimental conditions. Currently there is no way to predict the accuracy of an algorithm for a given data set. Most of the published algorithms and associated software has been evaluated on limited number of experimental data sets. However, these performance evaluations do not cover the complete search space the algorithmand the software might encounter in real-world. To this end, we present a database of simulated spectra that can be used to benchmark any spectra to peptide search engine. We demonstrate the usability of this database by bench marking two popular peptide sequencing engines. We show wide variation in the accuracy of peptide deductions and a complete quality profile of a given algorithm can be useful for practitioners and algorithm developers. All benchmarking data is available at<https://users.cs.fiu.edu/~fsaeed/Benchmark.html>

Keywords

Benchmarking; Peptide search algorithms; Proteomics; Mass-spectrometry

1 Introduction

For more than four decades, benchmarks have been used to assess the reproducibility and reliability of hardware (e.g. SPEC computer architecture benchmark) or software (e.g. BAliBASE for sequence alignments). Benchmarks provide a method of comparing the performance of given entity across various possible variables and gives a relative

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Author Contributions MA devised the method and database, wrote the software for generation of the database and wrote the manuscript. AA performed experiments and analyzed the results. FS proposed the initial idea, designed the experiments, supervised the research, and wrote and edited the manuscript.

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Data availability The proposed database is available at: <https://users.cs.fiu.edu/fsaeed/Benchmark.html>

Conflict of interest All authors declare that they have no conflict of interest.

Code availability Proposed databases were generated using the MaSS-Simulator software (Gul Awan and Saeed 2018).

performance by running a number of standard tests. These benchmarks ensure the reproducibility of the software for diverse conditions.

Mass spectrometry (MS) based proteomics (Aebersold and Mann 2003; Iglesias-Gato et al. 2016; Ebhardt et al. 2015; Tsai et al. 2015; PedroM and Bengt 2016; Saeed 2015) has revolutionized the study of system biology, and relies heavily on large number of software tools that automate the process of annotation and assessment of MS/MS spectra (Gul Awan and Saeed 2016; Kong et al. 2017; McIlwain et al. 2014). However, majority of the software tools that are published have been evaluated on a small set of experimental data which represents only a fraction of experimental conditions that would be encountered by the algorithm in real-world. The reliability of these algorithms and software packages then becomes questionable when they are encountered with novel data sets as demonstrated in Sect. 5, and 6. Further, it is up to the proteomics (or proteogeonomics/metaproteomics) practitioners to select a tool which would give the best accuracy for a given data set without using any quantifiable metric. The informed decision generally rests on what software tool the user is more comfortable with *instead* of what software would be best for this specific collected data set.

Similar problems have been encountered in other fields of science. One example most relevant to proteomics is the multiple-alignment problem in genomics analysis. Like peptide deduction, multiple-alignment is challenging because of many-solutions for a given data. In order to standardize the algorithm and software development, researchers came up with different benchmarks that could be used to assess the multiple alignment algorithms. Benchmarks such as BaliBase are a hallmark of alignment algorithm development (Arbelaez et al. May 2011; Zhenqin et al. 2018; Freytag et al. 2018) and have standardize the metrics that can be used for assessment.

In this paper, we present a standard reference database for proteomics as a bench-marking dataset for proteomics algorithms. Proposed database consists of spectra that have been simulated using MaSS-Simulator (Gul Awan M and Saeed F (2018)) while carefully adjusting its input parameters to cover a wide range of experimental conditions such as different dissociation strategies and peptide coverage. The benchmark data sets then can be used for assessing the accuracy and sensitivity of the existing and new algorithms, and thereby providing a scale against which all the algorithms can be evaluated.

2 Existing methods of algorithmic evaluation

Evaluating any algorithm requires ground truth data sets. In case of MS proteomics, these datasets consist of spectra that have been annotated with corresponding peptides. The methods of generating ground-truth datasets consists of annotating acquired MS/MS spectra using any existing peptide search algorithm and then evaluating these matches using statistical analysis tools specifically designed for MS proteomics data. These methods assign each peptide-to-spectra match (PSM) with statistical confidence value, each method may employ a different strategy or a variation of some existing method to determine statistical significance of these matches (Käll et al. 2008; Elias and Gygi 2007; Käll et al. 2007; Shteynberg et al. 2011; Keller et al. 2002). Among these, the most popular and widely

accepted metric has been percolator's False Discovery Rate (FDR). The Peptide Spectral Matches (PSMs) obtained from a search algorithm are processed by percolator which assigns each PSM with an FDR value. PSMs with FDR value of less than 1% are generally accepted as ground-truth data. Flow chart of FDR based generation of ground-truth has been shown in Fig. 1. Despite the ingenuity and wide acceptance of this method the ground truth data is far from perfect with wide variation in quality of spectra for a given FDR (Savitski et al. 2015).

To evaluate our argument, we used the proposed reference database to assess the reliability of a popular PSM assessment algorithm which assigns FDR to PSMs. Our results have shown that even with 1% FDR filtering, there are cases when up-to 35% of the PSMs are incorrect.

In this paper, we present a benchmarking database with large number of parameters that are used to simulate this data. Using this database, we demonstrate the shortcomings of existing ground-truth evaluation methods. We follow this discussion with two experimental demonstrations of how benchmarking database can be used to evaluate different types of peptide sequencing algorithms. For the sake of this study we use two algorithms, (1) Tide (Database search) (Diament and Noble 2011) and (2) Novor (Denovo sequencing) Ma (2015).

3 Proposed benchmarking database

The benchmarking reference database has been constructed by considering many possible combinations of six variables which govern the nature of MS/MS spectra that are generated during MS based proteomics experiments. A list of variables and the possible states that they can assume has been shown in Table 1. Real world MS spectra are stochastic and one can argue that six variables are not enough to simulate experimental conditions. Despite this argument, we have previously shown that correctly chosen small number of variables can be used to simulate MS/MS spectra that are very close to the real-world spectra with relatively small error percentage (Gul Awan and Saeed 2018). Since these are the variables that can easily be customized using MaSS-Simulator (Gul Awan and Saeed 2018), we chose them as the control variables for the proposed benchmarking database.

By assigning different values to each variable a unique experimental condition can be simulated using MaSS-Simulator. For this database we put together three hundred and twenty-four (324) possible experimental conditions by using variables and their states from Table 1. As a result, we have a very comprehensive benchmarking database covering a wide variety of dissociation strategies, noise content, relative intensities of signal peaks, peptide coverage, peptide length and peptide modifications. For each of the possible experimental conditions we simulate 1000 spectra in a single .ms2 file accompanied with a corresponding ground-truth file. In the table below, each variable can take several states. Using a combination of all possible states we have generated three hundred and twenty-four MS/MS spectra files with their corresponding peptide sequences available. Brief details of all the parameters have been provided in Table 2.

4 Assessment of FDR based method

A popular post-processing algorithm for evaluating PSMs from database search is Percolator from the Crux Toolkit (Käll et al. 2007). Percolator makes use of the Target-Decoy strategy and uses semi-supervised learning approach to assign FDR indicators to each PSM. Existing proteomics software evaluation is primarily done using percolator generated ground-truth spectra and is frequently considered as the gold-standard.

To evaluate how accurately Percolator assigns FDR values to PSMs we used the above discussed bench-marking database. The standard spectra were labelled with peptide sequences using Tide database search algorithm (Diament and Noble 2011) and then postprocessed using Percolator. Since we already had the ground-truth for simulated spectra available, we were able to evaluate the accuracy of FDR values assigned to each PSM. We filtered out PSMs which had been assigned an FDR value of 1% or less and then evaluated the accuracy by comparing the assigned peptide against the ground-truth peptides. It can be observed in Fig. 2 that for multiple experimental conditions Percolator is not able to accurately assign the FDR values. Plots in Fig. 2 give a comprehensive analysis of Percolator. Additional plots for remaining benchmark files can be found in Supplementary materials.

Our experiments and the results in the figure also suggest that spectra with low-coverage and peptides with shorter lengths are incorrectly assigned using Percolator.

These results support the hypothesis that low FDR PSM's cannot always be assumed to be a reliable ground-truth.

5 Evaluation of database search algorithm

To demonstrate the usability of the benchmarking reference database, we evaluated the performance of Tide database search software to observe its performance over various experimental conditions. Figure 3 shows plots which represent the behavior of Tide-Search engine across all the possible conditions covered by the proposed benchmarking database (additional plots available in supplementary materials). It can be observed that with decreasing peptide coverage the performance of Tide falls drastically and with a combination of low coverage and high noise the number of correct peptides even falls below 10%. In Fig. 3b it can be observed that introduction of two PTMs can also negatively affect the accuracy of Tide. But as shown in Fig. 3c, for longer peptide lengths Tide performs quite well in general but can still give a large number of incorrect matches when peptide coverage is low.

6 Evaluation of denovo sequencing algorithm

To demonstrate the usability of the benchmarking reference database, we evaluated the performance of Novor which is a denovo search software to observe its performance over various experimental conditions. Novor was used with its default settings, each .ms2 file was processed by Novor and the results were then evaluated by measuring recall value for each

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sequenced peptide and averaging the recall for each file in database (one file contains 1000 simulated spectra).

It can be observed in Fig. 4 that increasing Ion Coverage improves the recall significantly, across all lengths of peptides. Similarly, it can be seen that Companion Ions also have a significant effect on recall and low percentage of companion ions adversely effects Novor's performance. In general, performance of Novor is better for smaller peptides as compared to longer peptides. Results for additional datasets can be found in supplementary materials. It can be noted that the insights about the algorithm's performance, such as the effect of Companion Ions' population and peptide length on overall accuracy of the algorithm are not possible to understand when algorithms are published with handful of experimental datasets. This can be only possible when minor details of the spectra in the dataset are known and understood as was the case for the proposed benchmarking database.

7 Closing remarks

We have introduced a novel benchmarking strategy that can be used for evaluation of the algorithms on a wide variety of experimental conditions. This coverage of experimental conditions on our benchmarking data set will allow developers to report their accuracy of peptide deductions on a uniform scale and metrics. This benchmarking database strategy is the first step towards making peptide deduction algorithms more reliable and predictable in performance, and proteomics results more reproducible. Using this benchmarking database during algorithm development process can provide valuable insights into the design of the algorithms. Such benchmarking, if widely adopted, will help identify pitfalls and steer the algorithmic development process in correct direction. Profiling of these search algorithms will also make their performance more predictable for proteomics practitioners.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1.

Conventional work flow for generation ground-truth datasets for proteomics and evaluation of peptide sequencing algorithms

 \mathcal{C}°

 $-$ POS=3

Cos

A Peptide length fixed to state 1 and PTM to state 2.

 Be_{ak} Intensity

 $-POS=1$ $POS=2$

 $ln t_{\alpha}$ nsity

B Peptide length fixed to state 1 and PTM to state 1. Ion Coverage=3 Ion Coverage=2 Ion Coverage=1

C Peptide length was fixed to state 2 and PTM to state 1.

Plots showing percentage of Peptide correctly filtered using 1% FDR criteria

 $\rightarrow \text{POS}=1$ $\rightarrow \text{POS}=2$ $-$ POS=3

A Peptide length fixed to state 1 and PTM to state 2.

B Peptide length fixed to state 1 and PTM to state 1. Ion Coverage=1 Ion Coverage=2 Ion Coverage=3

C Peptide length was fixed to state 2 and PTM to state 1.

Plots showing percentage of Peptides correctly identified using Tide database search

A Peptide length fixed to state 1 and PTM to state 2.

B Peptide length fixed to state 1 and PTM to state 1. Ion Coverage=2 Ion Coverage=3 Ion Coverage=1

C Peptide length was fixed to state 2 and PTM to state 1.

Plots showing recall for Novor algorithm across the spectra from benchmarking database

Table 1

Each parameter can take several possible states Each parameter can take several possible states

Table 2

Description of different parameters used to develop the standard proteomics database Description of different parameters used to develop the standard proteomics database

