

Fast and accurate protein false discovery rates on human proteome study scale with Percolator 3.0

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Abstract

Percolator is a popular tool to assign reliable statistics, such as q-values and posterior error probabilities, to peptides and peptide spectrum matches (PSMs) using search results from mass spectrometry-based proteomics experiments. Percolator's processing speed has been sufficient for typical data sets with hundreds of thousands of PSMs. With our new scalable approach, we can now also analyze millions of PSMs in a matter of minutes on a commodity computer. Furthermore, with the increasing awareness for the need for reliable statistics on the protein level, we compared several easy-to-understand protein inference methods and implemented the best method in the Percolator package. We used Percolator 3.0 to analyze the data from a recent study of the draft human proteome containing 20 million spectra (PM:24870542).

Introduction

Percolator [6] has played a prominent part in the analysis pipelines of shotgun proteomics experiments for the last decade, by post-processing the results from database search engines such as SEQUEST [3], MASCOT [1], X!Tandem [2] and MS-GF+ [9]. Not only does Percolator often give a significant boost in the number of significant peptide spectrum matches (PSMs) or peptides, it also provides a consistent statistical framework in which to interpret the search results. As Percolator’s running time is usually much lower than that of the search engine, applying it as a post-processing step should be a no-brainer. As part of the continuous development and support of the Percolator package, we present two major additions aimed at supporting analysis of studies on the scale of the human proteome [8, 13].

As advances in technology are causing shotgun proteomic experiments to become progressively easy and affordable to carry out, the amount of data per study will keep rising steadily. While previous versions of Percolator are able to process the data from the vast majority of current studies in a decent time frame, certain limitations have come into sight for laboratories without access to an above average commodity computer. When processing millions of PSMs, the majority of Percolator’s processing time is spent on training support vector machines (SVMs). One could, however, surmise that the performance of the SVM would plateau pretty fast as a function of the number of input PSMs. Here, we propose to use Percolator’s semi-supervised learning algorithm to train SVMs on only a random subset of the PSMs and used the resulting score vectors to evaluate the rest of the PSMs.

Second, protein-level accuracy estimates have been on our feature wish list for quite some time now. One of the major obstacles was the question of how to deal with shared peptides and protein grouping. An implementation of Fido [12] has been part of the Percolator package for quite some time now and addressed these two issues, but is too computationally intensive on large-scale datasets with many shared peptides. On the other hand, large-scale studies have a deep coverage of the present peptides and therefore identify many peptides that uniquely identify a protein. This provides the option of ignoring shared peptides altogether and makes the task of protein inference much simpler and intuitive. Here, we compared several easy-to-understand protein inference strategies that only use these peptides that uniquely identify a protein and implemented the best candidate in the Percolator package.

Methods

We downloaded a set of spectra, comprising 2212 runs on 17 adult tissues, 7 fetal tissues, and 6 hematopoietic cell types with a total of 21 million spectra from [8]. The investigated peptides were analyzed on an LTQ Orbitrap Velos and Elite (Thermo Scientific) equipped with an Easy-nLC II nanoflow LC System (Waters). We will refer to this set as the *pandey* set.

Converting the RAW files to MS1 and MS2 files was done with Proteowizard [7]. Next, we assigned

high-resolution precursor masses and charges using information from the precursor scans with Hardklör [4] followed by Bullseye [5], through the Crux 2.0 package interface [10]. This was followed by a database search against the human Swissprot and Swissprot+Ensembl databases (accessed: 2015 Nov 12) using the Tide search engine, again through the Crux interface. The same search parameters as in [8] were used, with the exception of not using cyclization of N-terminal glutamine as a variable modification and using Tide’s default fragment tolerance. For the decoy proteins, we simply reversed the protein sequences. This process resulted in 73 million target and decoy PSMs.

To characterize the behavior of the scoring vectors based on subsets of the PSMs, we evaluated the performance for different sizes of the random subset. For each random subset size, we calculated the mean and standard deviation over 10 randomized runs of the number of PSMs and peptides with q value below 0.001.

Before applying the protein inference methods, we used the approach to handling shared peptides from *Nesvizhskii et al.* [11]. Here, proteins are grouped that are indistinguishable based on their theoretical proteolytically digested peptides, rather than their experimentally discovered peptides. We retain the peptides that are unique to such a protein group, rather than to a single protein. Especially for databases containing many proteoforms for a gene, this can decrease the number of shared peptides considerably.

We compared several straightforward protein inference methods: Fishers method for p-value combination, the picked target-decoy strategy, and the product of peptide-level posterior error probabilities (PEPs). We assessed the performance of the methods on the draft human proteome data, as well as on a standard protein mix. Using the standard protein mix we could characterize the accuracy and stability of the false discovery rate estimates, and the draft human proteome data could give us an indication of the performance on real data. We calculated false discovery rate (FDR) estimates from p-values for Fishers method, and using target-decoy models for all other methods.

Results

Searching all 73 million target+decoy PSMs resulted in 300 000 unique target peptides. Using subsets of even just 100,000 PSMs (0.1%) for SVM training did not reduce the number of identified peptides and generally even slightly increased this number. The standard deviation of the randomized runs for a fixed subset size did increase when taking increasingly smaller subsets, but this effect was limited. By using a subset of 500,000 PSMs to train the SVM, Percolators runtime was reduced from several hours to under 10 minutes.

We assessed the accuracy and stability of FDR estimates using three standard protein mixes. While Fishers method gave the most robust FDR estimates, it identified 5-10% fewer proteins than the picked target-decoy strategy and product of PEPs. These two strategies did give reasonably accurate FDR

estimates but were more sensitive to errors in the peptide identification.

For the draft human proteome set, at 1% FDR, the picked target-decoy strategy identified 12,300 proteins, multiplication of PEPs 11,600 and Fishers method only 8,800. From these results we concluded that the picked target-decoy strategy was the superior alternative and implemented this in the newest Percolator package.

Discussion

Acknowledgements

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