Systems biology

LuciPHOr2: site localization of generic post-translational modifications from tandem mass spectrometry data

Damian Fermin^{1,†}, Dmitry Avtonomov¹, Hyungwon Choi^{2,*} and Alexey I. Nesvizhskii^{1,3,*}

¹Department of Pathology, University of Michigan Medical School, Ann Arbor, MI, USA, ²Saw Swee Hock School of Public Health, National University of Singapore and National University Health System, Singapore, Singapore and ³Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, MI, USA

*To whom correspondence should be addressed. [†]Present address: Department of Pathology, Yale University, New Haven, CT, USA Associate Editor: Jonathan Wren

Received on September 9, 2014; revised on October 18, 2014; accepted on November 20, 2014

Abstract

We present LuciPHOr2, a site localization tool for generic post-translational modifications (PTMs) using tandem mass spectrometry data. As an extension of the original LuciPHOr (version 1) for phosphorylation site localization, the new software provides a site-level localization score for generic PTMs and associated false discovery rate called the false localization rate. We describe several novel features such as operating system independence and reduced computation time through multiple threading. We also discuss optimal parameters for different types of data and illustrate the new tool on a human skeletal muscle dataset for lysine-acetylation.

Availability and implementation: The software is freely available on the SourceForge website http://luciphor2.sourceforge.net.

Contact: hyung_won_choi@nuhs.edu.sg, nesvi@med.umich.edu

Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Post-translational modifications (PTMs) induce conformational changes in proteins that influence their enzymatic activity, subcellular localization and/or their binding affinity (Hoffman *et al.*, 2008; Silva *et al.*, 2013). High-throughput analysis of PTMs is typically achieved through mass spectrometry (MS) (Silva *et al.*, 2013), where tandem (MS/MS) mass spectra are searched against a sequence database allowing for PTMs. A major challenge in the interpretation of MS/MS data for PTMs is that database search engines can correctly match a peptide sequence with incorrectly localized PTM(s). These errors are not controlled by the type I error of peptide identification because each peptide has a different degree of uncertainty as far as site localization is concerned. For this reason, the conventional target-decoy framework often leads to underestimation of the error rates for site localization of PTMs (Fermin *et al.*, 2013). Hence, an

additional procedure to control the number of false site localizations is necessary when reporting PTM localization analysis results at the proteome scale.

2 Functional features

2.1 Improved implementation

To address this issue, we recently published a computational method called LuciPHOr for site localization analysis of phosphorylation events using MS/MS data (Fermin *et al.*, 2013). LuciPHOr was first implemented with a novel target-decoy framework specifically designed for phospho-site localization. Here, we have converted this tool into a general PTM localization program (LuciPHOr2) for estimating false localization rate (FLR) for any PTMs of a fixed mass. The new implementation provides enhancements over the previous

version: (i) it is written entirely in Java, making it operating system independent; (ii) the software now processes search results from any proteomics search engine.

2.2. Key parameters for site localization

In LuciPHOr2, the user is expected to specify key parameters for statistical modeling, which determine inclusion/exclusion of MS/MS peaks into the scoring algorithm. LuciPHOr2 constructs a probability model of peak intensity and mass accuracy across all spectra for correct and incorrect localizations respectively, and computes the likelihood ratio as the localization score for each candidate site. This requires an appropriate choice of low/high mass accuracy model (ALGORITHM parameter), fragment mass tolerance (MS2_TOL) and the criteria for selecting high confidence localizations to train the probability model (SELECTION_METHOD).

Two additional parameters are worth mentioning in this new version: the target modification and the mass shift of the PTM of interest. Since LuciPHOr2 was designed to score generic PTMs, the user must specify what amino acids should be treated as potential modification sites along with their respective mass shifts. The mass shift is required for the program to place decoy PTMs on arbitrary sites (non-specific residues) and accurately estimate the FLR.

To facilitate the optimal configuration, LuciPHOr2 generates a template input file that the user can edit for their specific needs. However, we note that the default parameters generated in this file are optimized for phosphorylation analysis.

3 Application to generic PTM analysis

We demonstrate the program through the analysis of a published dataset for lysine acetylation in 12 human skeletal muscle biopsies (Lundby *et al.*, 2012). The RAW files were converted into the opensource format mzML using MSConvert from the ProteoWizard package (Kessner *et al.*, 2008), and searched with X!Tandem (Fenyo and Beavis, 2003). The database search was performed as described by Lundby *et al.* with two exceptions: the precursor tolerance was increased to 20 ppm and up to five missed cleavages were allowed. The peptide search results were post-processed using PeptideProphet of the TPP (Deutsch *et al.*, 2010; Keller *et al.*, 2002). All peptide-to-spectrum matches (PSMs) with a lysine acetylation event were reanalyzed with LuciPHOr2 configured for lysine-acetylations (K+42.01056).

A total of 35482 PSMs were reported with acetylation by X!Tandem/TPP. To filter high-quality spectra, 30 602 PSMs within the 1% FDR threshold (PeptideProphet probability 0.809) were selected, and among these 20 754 PSMs had site ambiguity (68%). As expected for a non-labile modification, X!Tandem/TPP and LuciPHOr2 showed concordant site localization on the majority of ambiguous cases (19 557/20 754, 94%). For a non-negligible number of PSMs (673/19 557, 3.4%), however, LuciPHOr2 delta score was <3 between the top two localizations, indicating that those sites were localized based on merely a single fragment ion or less credible low intensity fragments, implying considerable ambiguity.

On the other hand, LuciPHOr2 and X!Tandem/TPP disagreed on 1197 site localizations, consisting of two major types. The first type were 987 PSMs with the ambiguity on two consecutive lysines at the end of the peptide sequences (e.g. VLTLELYKK), whereas the second type were 210 PSMs with the ambiguity in the middle of the sequences (e.g. DNKKPFIKEAAMAK). In nearly all of the latter cases, there was no major site discriminating fragment ion. This is

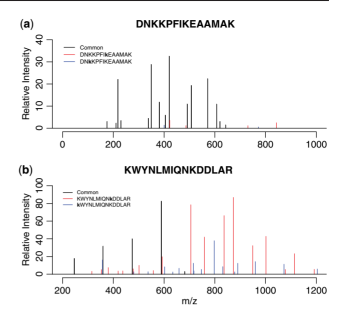


Fig. 1. MS/MS spectra with site determining ions for top two site localizations (matched peaks only). Black peaks are shard ions, while red and blue ones are the site determining ions for each site localization. The two site localizations are shown in the text in the upper left corner of each panel (lower case indicates PTM sites). (a) Peptide with a low score due to lack of clear site determining ions. (b) Peptide with a high score with evidence for both localizations, indicating the possibility of co-eluting positional isomers

illustrated in Figure 1a, where only a few minor peaks support both candidate sites (Supplemental Table S1).

Another advantage of LuciPHOr2 is that it produces a peak-bypeak score report for the two best site localizations, which can assist identification of positional isomers. The term 'positional isomers' refers to two co-eluting species of the same peptide with different site localizations (Courcelles *et al.*, 2012), often captured in the same MS/MS spectrum. To find these, the peak score report can be queried for the peptides with high likelihood scores (e.g. >30 for low mass accuracy data, >50 for high mass accuracy data) with a sufficient number of site determining ions for each localization. Figure 1b shows the spectrum for KWYNLMIQNKDDLAR, which has 18 and 16 site determining ions for the two lysines (Supplemental Table S1). This example was easily queried with the following criteria: the top two localizations score above 50 on the same spectrum and have more than five site determining ions.

4 Conclusions

In sum, the *post hoc* analysis of site localization of PTMs using the database search output is an essential step toward the control of type I errors in large-scale PTM analysis. The LuciPHOr2 package facilitates such analysis using the peptide identification from various database search engines, and is expected to contribute to more robust analysis of proteome-level PTM analysis in a wide range of applications.

Funding

This work was supported by Singapore MOE [grant R-608-000-088-112 to H.C.] and NIH [R01-GM-094231 to A.I.N.].

Conflict of Interest: none declared.

References

- Courcelles, M. et al. (2012) Occurrence and detection of phosphopeptide isomers in large-scale phosphoproteomics experiments. J. Proteome Res., 6, 3753–3765.
- Deutsch,E.W. et al. (2010) A guided tour of the trans-proteomic pipeline. Proteomics, 10, 1150–1159.
- Fenyo,D. and Beavis,R.C. (2003) A method for assessing the statistical significance of mass spectrometry-based protein identifications using general scoring schemes. *Anal. Chem.*, 75, 768–774.
- Fermin, D. et al. (2013) LuciPHOr: algorithm for phosphorylation site localization with false localization rate estimation using modified target-decoy approach. Mol. Cell. Proteomics, 12, 3409–3419.
- Hoffman, M.D. et al. (2008) Current approaches for global post-translational modification discovery and mass spectrometric analysis. Anal. Chim. Acta, 627, 50–61.
- Keller,A. et al. (2002) Empirical statistical model to estimate the accuracy of peptide identifications made by MS/MS and database search. Anal. Chem., 74, 5383–5392.
- Kessner, D. et al. (2008) ProteoWizard: open source software for rapid proteomics tools development. Bioinformatics, 24, 2534–2536.
- Lundby, A. et al. (2012) Proteomic analysis of lysine acetylation sites in rat tissues reveals organ specificity and subcellular patterns. Cell Rep., 2, 419–431.
- Silva, A.M. et al. (2013) Post-translational modifications and mass spectrometry detection. Free Radical Biol. Med., 65, 925–941.