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PINE: An Automation Tool to Extract and Visualize Protein-Centric Functional Networks

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Abstract

Recent surges in mass spectrometry-based proteomics studies demand a concurrent rise in speedy and optimized data processing tools and pipelines. Although several stand-alone bioinformatics tools exist that provide protein–protein interaction (PPI) data, we developed Protein Interaction Network Extractor (PINE) as a fully automated, user-friendly, graphical user interface application for visualization and exploration of global proteome and post-translational modification (PTM) based networks. PINE also supports overlaying differential expression, statistical significance thresholds, and PTM sites on functionally enriched visualization networks to gain insights into

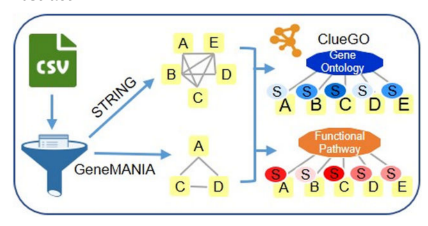
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proteome-wide regulatory mechanisms and PTM-mediated networks. To illustrate the relevance of the tool, we explore the total proteome and its PTM-associated relationships in two different nonalcoholic steatohepatitis (NASH) mouse models to demonstrate different context-specific case studies. The strength of this tool relies in its ability to (1) perform accurate protein identifier mapping to resolve ambiguity, (2) retrieve interaction data from multiple publicly available PPI databases, and (3) assimilate these complex networks into functionally enriched pathways, ontology categories, and terms. Ultimately, PINE can be used as an extremely powerful tool for novel hypothesis generation to understand underlying disease mechanisms.

Graphical Abstract



Keywords

proteomics; data visualization; enriched networks; automated bioinformatics tool; post-translational modifications

INTRODUCTION

The aim of mass spectrometry (MS) based proteomics studies is to identify and quantify a wide range of proteins and PTMs in biological samples of interest to uncover disease mechanisms and novel protein biomarkers. 1,2 For the last two decades, liquid chromatography-tandem mass spectrometry (LC-MS/MS) of complex mixtures of proteins in data-dependent acquisition (DDA) mode³ has been the standard in proteomics for sampling breadth and discovery. Recent improvements in mass spectrometer design and bioinformatics algorithms have resulted in the development of another peptide sampling method: data-independent acquisition (DIA)⁴ which aims to combine identification breadth of DDA with the sensitivity and reproducibility of targeted MS assays such as multiple reaction monitoring (MRM) or parallel reaction monitoring (PRM).^{5,6} These proteomic technological developments have led to a software evolution going from stand-alone software tools for MS data processing to large-scale complex computational pipelines transforming raw data into biological discoveries. With these advances, we can now routinely analyze large-scale quantitative differences between samples but more importantly we can also detect and quantify an array of protein PTMs that were previously challenging.8-10

The unambiguous identification and quantitation of PTM sites by classical bottom up methods still relies on proteotypic peptide identification, modified amino acid residue localization, peptide ionization and fragmentation, MS instrument accuracy, as well as its abundance in the biological sample of interest. ^{11,12} Despite these challenges, PTMs can be biologically insightful since they serve to diversify the function of proteins by acting as key regulators of protein function, structure, and PPIs in response to perturbations. ¹³ PTMs can occur on multiple sites of the same protein or the same site of a protein modified by different types of PTMs that could potentially serve as mechanistic switches. ¹⁴⁻¹⁶ An emerging challenge is therefore to decipher how PTMs, individually or in combination, modulate protein interactions to ultimately inform our understanding of key molecular processes and regulatory functions. ¹⁷

Several existing bioinformatics tools are available for visualizing and analyzing PPI networks. Databases targeted at collecting and storing PPIs, such as STRING, ¹⁸ GeneMA-NIA, ¹⁹ and Reactome²⁰ are publicly available and easily accessible. Similarly, KEGG (Kyoto Encyclopedia of Genes and Genomes), ²¹ Reactome, WikiPathways, ²² and Gene Ontology (GO)^{23,24} act as knowledge bases to provide functional and biological implications of proteins. Ingenuity Pathway Analysis (IPA)²⁵ builds on its comprehensive, manually curated knowledge base to provide biological insights from expression analysis experiments. PTMapper, ²⁶ Phospho-Path, ²⁷ and PTMOracle ²⁸ allow users to covisualize and coanalyze PTM data in the realm of PPI networks and helps improve our understanding of PTM-associated relationships. Analogously, piNET,²⁹ another PTM-centric visualization tool, maps peptides to proteins, performs upstream analysis of PTM sites, and conducts pathway analysis. Despite this, however, there is a sparsity of tools that (1) perform protein-centric PPIs and functional enrichment analysis while maintaining the protein-gene relationship from public databases which predominantly contains gene-centric information, (2) address ambiguity arising from combining data retrieved through various independent databases, and (3) simultaneously explore the regulatory role of PTMs and their involvement in PPIs and biological function.

With the goal of addressing these challenges, PINE has been developed as a fully automated, versatile, and easy to use Cytoscape-based tool³⁰ to facilitate mapping, annotation, and visualization of protein and PTM-level quantitative data generated by either targeted, discovery (DDA and DIA), or other proteomics approaches. It allows users to develop systematic searches for proteins of interest and explore complex network visualizations depicting PPIs and functional relationships. In order to improve proteome-wide coverage of PPIs, PINE supports extraction and integration of PPIs from different publicly available databases such as STRING and GeneMANIA with the ability to apply score thresholds based on the nature and quality of the supporting evidence, all while maintaining the protein—gene relationship and resolving ambiguity in the data. PINE utilizes ClueGO³¹ for functional enrichment analysis over other tools such as BiNGO³² and PIPE³³ because of its ability to (1) integrate GO terms and pathway analysis from multiple databases, (2) dynamically create annotation networks based on user query list, and (3) cluster genes and compare functional differences to provide accurate term statistics. PINE also supports overlaying differential expression, statistical significance thresholds, and PTM sites (if

applicable) on functionally enriched visualization networks to gain insights into proteomewide regulatory mechanisms as well as how PTMs might regulate PPIs.

To illustrate how PINE can be used to explore global proteomic signatures, crosstalk between PTMs, and their involvement in PPIs, we present case studies using two mouse models of nonalcoholic steatohepatitis, ³⁴ each chosen due to their extreme cellular methylation potential. Nonalcoholic fatty liver disease (NAFLD), the leading cause of chronic liver disease in Western countries, starts as a consequence of defects in diverse metabolic pathways that involve hepatic accumulation of triglycerides. In some cases, this progresses to nonalcoholic steatohepatitis (NASH), a condition characterized by the appearance of inflammation, cellular injury with or without fibrosis, together with steatosis. 35 Because there are different causes that can lead to steatosis, we hypothesized that different NASH subtypes may exist, reflecting the variety of mechanisms causing liver fat accumulation, and that identifying each subtype would help us characterize early proteomic changes in NAFLD development as well as key PTMs driving the progression toward NAFLD and subsequent NASH. We assessed this hypothesis by analyzing two mouse models of nonalcoholic steatohepatitis (NASH), (1) the glycine N-methyltransferase knockout (Gnmt -/-) mouse model, which has increased hepatic SAMe and is biologically hypermethylated, and (2) the methionine adenosyltransferase 1A knockout (Mat1a -/-) mouse model, which has hepatic SAMe deficiency and therefore is hypomethylated, to understand the different mechanisms causing liver fat accumulation in NASH.34,36-39 Both animal models spontaneously develop NASH and resemble many aspects of the major subtypes of human disease. ^{34,40} Interestingly, disease in both animal models can be treated by normalization of SAMe levels ,highlighting the importance of maintaining normal hepatic SAMe level and methylation capacity in NASH. ^{34,39,42} This is relevant because it may provide a better understanding of early molecular drivers of NASH and key PTMs that may drive disease progression which would be beneficial in developing early stage biomarkers and therapies for NASH.

METHODS

Sample Preparation for Proteomic Analysis.

Livers were obtained from 10-month old *Mat1a* –/– male mice in a C57Bl/6 background with hepatic lipid accumulation and their aged-matched wild-type (WT) male sibling littermates,³⁴ and three-month old *Gnmt* –/– mice in a C57Bl/6 background with hepatic lipid accumulation with age matched WT littermates.^{36,37} Frozen mouse livers were ground while frozen in a liquid N₂-cooled cryohomoginizer (Retsch). Samples were denatured in 8 M urea and 100 mM Tris–HCl, pH 8.0 and ultrasonicated (QSonica) at 4 °C for 10 min in 10 s repeating on/off intervals of 10 s and centrifuged at 16000x for 10 min at 4 °C. The soluble fraction was reduced with DTT (15 mM) for 1 h at 37 °C, alkylated with iodoacetamide (30 mM) for 30 min at room temperature in the dark, diluted to a final concentration of 2 M urea with 100 mM Tris–HCl, pH 8.0, and digested for 16 h on a shaker at 37 °C with a 1:40 ratio of Trypsin/Lys-C mix (Promega). Each sample was desalted using HLB plates (Oasis HLB 30 µm, 5 mg sorbent, Waters).⁴²

Titanium Dioxide Phospho Enrichment.

Titanium dioxide (TiO₂) phospho enrichment was performed as previously described. Briefly, 400 μ g of sample was digested as described above, individually desalted on Oasis HLB cartridges (Water, 10 mg), and eluted in 300 μ L of 80% ACN, 5% TFA, 1 mM glycolic acid. Each sample was then incubated in 50 μ L of titanium dioxide (TiO₂) slurry (30 mg/mL, Glygen Corp, Columbia, MD) at room temperature on a shaker overnight. Then the TiO₂ beads were washed twice with 200 μ L of 80% ACN, 5% TFA, once with 200 μ L of 80% ACN, 0.1% TFA, and eluted in 180 μ L of 30% ACN/1% NH4OH and neutralized with 200 μ L of 10% FA. Samples were then desalted on Oasis HLB μ -elution plates (Waters) and eluted in 80% ACN, 0.1% FA, dried in a speedvac, and then resuspended in 0.1% FA for LC–MS/MS analysis.

MS Acquisition.

MS data were acquired on Orbitrap Fusion Lumos (Thermo) mass spectrometer connected to an Ultimate 3000 nano LC system (Thermo Scientific) with a 60 min or 120 min gradient operating in DDA-MS or DIA-MS mode. ⁴² Briefly, for DDA analysis, precursor ions were fragmented using HCD and analyzed in the Orbitrap at 15000 K resolution. Monoisotopic precursor selection and dynamic exclusion was enabled and only MS1 signals exceeding 50000 counts triggered the MS2 scans. For DIA analysis, MS1 scans were acquired at 60,000 Hz from mass range 400–1000m/z. DIA MS2 scans were acquired in the Orbitrap at 15,000 Hz with fragmentation in the HCD cell. DIA was performed using 4 Da (150 scan events) windows over the precursor mass range of 400–1000 *m/z*, and the MS2 mass range was set from 100 to 1500 *m/z*.

Generation of Spectral Libraries and DIA Analysis.

A lysine PTM enriched peptide assay library⁴² was used for the DIA analysis and quantitation of individual specimens. Briefly, a hypermethylated DIA assay library was created by DDA acquisitions of 5 SCX fractions from *Gnmt* –/– mouse livers. This library was supplemented with a DDA acquisition of a methyl-Lys immunoprecipitation from a hepatoma cell line derived from *Gnmt* –/– mice.⁴¹ The library was further supplemented with DDA acquisitions of a "one-pot" acetyl-Lys and succinyl-Lys immunoaffinity enrichment of mitochondrial extract and whole liver lysate as previously described.⁴⁴

Briefly, raw intensity data for peptide fragments was extracted from DIA files using the open source openSWATH workflow⁴⁵ against the lysine PTM enriched peptide assay library. Target and decoy peptides were then extracted, scored, and analyzed using the mProphet algorithm⁴⁶ to determine scoring cut-offs consistent with 1% FDR. Peak group extraction data from each DIA file was combined using the "feature alignment" script, which performs data alignment and modeling analysis across an experimental data set.⁴⁷ Normalized transition-level data was then processed using the mapDIA software to perform pairwise comparisons between groups at the peptide level.⁴⁸ All data sets have been made publicly available at Panorama (PXD ID: PXD012621): https://panoramaweb.org/methylation_methods_1.url.

Phospho Enrichment Database Searches and Quantification.

DDA files were converted to mzXML and searched through the Trans Proteomic Pipeline (TPP) using three algorithms, (1) Comet, ⁴⁹ (2) X!tandem! Native scoring, ⁵⁰ and (3) X!tandem! K-scoring⁵¹ against a reviewed, mouse canonical protein sequence database, downloaded from the Uniprot⁵² database on January 24th, 2019, containing 17002 target proteins and 17002 randomized decoy proteins. Target-decoy modeling of peptide spectral matches was performed with PeptideProphet,⁵³ and peptides with a probability score of >95% from the entire experimental data set were imported into Skyline software⁵⁴ for quantification of precursor extracted ion intensities (XICs). Precursor XICs from each experimental file were extracted against the Skyline library, and peptide XICs with isotope dot product scores >0.8 were filtered for final statistical analysis of proteomic differences.⁵⁵ Raw peptide intensities were used to calculate pairwise comparisons between experimental groups using the linear mixed effects model built into the open sources MSSTATs (v3.2.2) software suite. ⁵⁶ Peptide abundance differences with an adjusted p-value < 0.05 were considered significantly different. The Skyline documents containing precursor XICs from each experimental file are available at Panorama (PXD ID: PXD017324): https:// panoramaweb.org/qt1CeZ.url

Software Design and Availability.

PINE is a tool designed to enable visualization of PPI networks and aids in forging biological interpretation through the creation of annotation networks. It allows users to integrate other types of protein data including fold changes, p-values, category, or PTM sites. To facilitate this, all the protein data must be formatted into a comma-separated (CSV) file. Protein data that is formatted into the required input format is parsed and visualized as PPIs networks on the visualization platform, Cytoscape, with nodes representing individual proteins and other protein data mapped onto the nodes. Concurrently, a list of annotation terms is generated from which users can select their terms of interest to create subnetworks depicting annotations linked to their related proteins. The features of PINE are described in more detail in the Results section. PINE is programmed in Python3, and the graphical user interface (GUI) for PINE was created with the Electron framework v6.0.7. Pyinstaller v3.4 was used to wrap PINE's Python code into an executable for use by the GUI. Prerequisites for using PINE include Cytoscape v3.7 or higher to be installed by users along with Cytoscape apps GeneMANIA and ClueGO. Programmatic access for the PINE workflow is enabled through Cytoscape's RESTful API methods⁵⁷ and the py2cytoscape⁵⁸ utility. The source code for PINE that invoke these respective APIs and the GUI along with additional usage documentation are included in the public GitHub repository, https://github.com/csmcvaneykjlab/pine, and an executable release is available for Windows 10.

RESULTS

Overview and Features of PINE.

PINE is a visualization tool that provides users with PPI and annotation networks for the integration of different types of protein data. To do so, the PINE framework consists of six primary steps: (1) preprocessing of input, (2) Uniprot mapping for protein to gene transformation, (3) retrieval of STRING PPIs, (4) simultaneous retrieval of GeneMANIA

PPIs, (5) merging from both public databases to construct single PPI network, and (6) performing enrichment analysis using Cytoscape plug-in, ClueGO to drive subnetwork creation (Figure 1A).

The preprocessing step parses the input containing proteins and their auxiliary data to accurately map protein data onto nodes in the final network. Considerations are made to ensure input data is valid (i.e., fold changes and *p*-values must be numeric), readily available (i.e., modified peptides must contain PTMs), and nonredundant. This step also allows the user to exclude ambiguous PTM sites with oxidized methionines, missed cleavages, and ragged ends.^{59,60} Next, the Uniprot mapping step performs an API call to Uniprot to map proteins to their corresponding primary genes. All subsequent steps use the primary gene to represent the input protein in order to remain consistent with various publicly available PPI databases. This step also conducts checks to ensure information is valid (i.e., protein identifiers are truly Uniprot IDs), readily available (i.e., proteins are successfully mapped to a primary gene), and nonredundant. At this level, data ambiguity is resolved by picking the first protein-to-gene match in cases such as (1) many proteoforms mapping to one primary gene and (2) one protein mapping to many primary genes. Additionally, the user also has an option to exclude all ambiguous identifiers from the analysis.

The next step retrieves PPIs from STRING and GeneMANIA and conducts checks to ensure information is valid (i.e., query genes are successfully mapped as a primary interactor), readily available (i.e., query genes are successfully mapped in STRING and/or GeneMANIA) and nonredundant interactions. The merging step consolidates PPI information from STRING and GeneMANIA to create a single, unified PPI network (as shown in Figure 1B). The ClueGO enrichment analysis step identifies enriched gene ontology terms and pathway annotation for all valid (i.e., one-to-one query gene mapping within ClueGO) and readily available (i.e., query genes successfully mapped in ClueGO) genes. Finally, PINE enables selection and visualization of top representative enriched terms into subnetworks using Cytoscape's RESTful API methods and the py2cytoscape utility. Cytoscape framework supports interactive network visualization through a Zoomable User Interface (ZUI) with customizable features such as zooming, panning, layout, and styling options to manage visual presentation.

Hereby, PINE provides a unique automation solution for robust and thorough data processing to avoid ambiguity, while simultaneously integrating information from various public databases for the creation of functionally enriched visualization networks.

PINE Interface for Visualization.

PINE provides a simple, easy-to-use, graphical user-interface having four main modules: (1) Initial Setup, (2) Parameter Settings, (3) Logging and Data Handling, and (4) Pathway Selection of enriched terms.

The first module is the Setup tab describing a one-time installation guideline required for running the tool. Subsequent uses of PINE automatically register the paths to the necessary packages. Next, the Settings tab (Figure 2A) enables users to customize their analysis with several mandatory and optional parameters. Mandatory parameters include input file

containing protein identifiers and related data columns in CSV format, type of run based on the proteomics experiment (for example, PTM-based run type can be selected in the presence of PTM sites), path to output directory and the species of interest. Additionally, in the case of PTM run types, users are required to provide the species database file in FASTA format, digestion enzyme used and modification sites of interest. Optional parameters include: fold change or *p*-value cutoffs, choice to add blue-colored outline to statistically significant protein or PTM nodes, choice to include or exclude all ambiguous data, choice of PPI databases to be used along with confidence score of interactions, option to include external (nonquery) interactors, functional enrichment analysis based on pathways (comprising of Reactome, KEGG, ClinVar, 61 CORUM and WikiPathways) or individual GO terms (biological process, cellular component or molecular function), ClueGO grouping level (i.e., representative or specific annotation terms), ClueGO enrichment *p*-value cutoff, and option to upload a background reference file for the enrichment analysis.

Once the analysis is submitted using user-defined settings, the interface is redirected to the Log tab which shows the progress of the run as well as records data handling at every step. Once the run is complete, the interface is redirected to the Pathway Selection tab (Figure 2B) which lists enriched annotation terms for the query data set along with term significance, corrected term significance, percent of genes in the cluster associated with the term and total number of term related genes. Top representative terms can be selected based on one or many of the following criterion: (1) terms with adjusted *p*-value 0.01, (2) terms with reasonable coverage based on % genes or number of genes, (3) terms most likely connected to the biological system of interest, and (4) terms comprising of candidate proteins of interest. Filtering and sorting options are also supported on the UI for easy and rapid selection of terms of interest for reanalysis and subnetwork creation which is ultimately rendered using Cytoscape's network visualization features.

Case Study 1: Identifying and Comparing Enriched Pathways and GO Terms.

Visualization of complex networks to uncover functionally enriched subnetworks can be challenging especially when large clusters of proteins and their PPIs need to be considered. To address this, we performed global quantitative proteomics using DIA across two mouse models to better understand the key commonalities and differences in the underlying functional pathways involved in causing liver fat accumulation in NASH (Figure 3). Using PINE, we first extracted and integrated PPI data from STRING and GeneMANIA for all statistically significant differentially expressed proteins (Bayesian False Discovery Rate 0.05) with 1.5 log2 fold change by comparing *Gnmt*—/— mice and *Mat1a*—/— mice relative to WT littermates. With ClueGO enrichment and Cytoscape operations automated within PINE, we created a subnetwork of top representative enriched terms to reduce visual complexity. Finally, we colored protein nodes based on the observed fold changes (with orange corresponding to upregulation and blue corresponding to downregulation) and enriched functional nodes were colored based on the percentage of upregulated or downregulated protein nodes to indicate activation or inhibition trend for that functional node (Figure 4).

The resulting subnetwork of enriched pathways contained six key functional processes for both NASH mouse models, respectively. The *Gnmt* –/– mice model showed activation of pathways such as fatty acid metabolism,⁶³ TCA cycle,^{63,64} glutathione/one carbon metabolism,^{34,65} peroxisomal protein import,^{34,66} glycolysis/gluconeogenesis,^{67,68} and inhibition of pathway such as metabolism of xenobiotics by cytochrome P450^{34,70} (Figure 4A). On the other hand, The *Mat1a* –/– mice model showed the reverse trends compared to *Gnmt* –/– mice model with inhibition of pathways such as fatty acid metabolism,⁶³ TCA cycle,^{63,64} peroxisomal protein import,^{34,66} pyruvate metabolism⁷¹ and activation of pathways such as glutathione metabolism^{34,65} and metabolism of xenobiotics by cytochrome P450^{34,69,70} (Figure 4B). These findings strongly correlate with known functional pathways altered in both NASH mouse models, although their proteomic expression profiles were regulated differently, elucidating key proteins could potentially be used as biomarkers to differentiate subtypes of NASH.^{34,42,72}

With PINE, we also visualized a subnetwork of enriched GO cellular components to identify similarities and differences in the subcellular localization between the two NASH mouse models (Figure 5). In PINE, we can represent differential expression data from multiple comparison groups (for example, *Gnmt* –/– mice and *Mat1a* –/– mice relative to their WT littermates) as bar charts wherein the height of the bar represents degree of fold change. Since mitochondria are well-known to play a critical role in the development and pathogenesis of NAFLD, ^{73,74} we focused our cellular subnetwork around the mitochondrial matrix to visualize proteomic signatures between the two mouse models. For example, carbamoyl-phosphate synthase (CPS1), a previously reported candidate biomarker for NASH, ⁷⁵ shown to be enriched in mitochondrial matrix ⁷⁶ was observed to be altered in both NASH mouse models but was significantly downregulated in *Mat1a* –/– mice relative to WT littermates (log2 fold change of –0.9) and nonsignificantly changed in *Gnmt* –/– mice relative to WT littermates (log2 fold change of –0.2), indicating that the two mice models might represent different pathophysiologies of NASH disease progression.

Case Study 2: Exploring Lysine and Arginine PTMs in the Context of Enriched Pathways and GO Terms.

In order to explore PTM crosstalk between methylation, succinylation and acetylation, we performed PTM analysis using DIA across the two NASH mouse models to visualize the relationships between quantified peptide/PTM/protein moieties in the context of NASH pathophysiology. Using PINE, we mapped all statistically significant differentially expressed PTM sites (Bayesian False Discovery Rate 0.05) with 1.5 log2 fold change between *Gnmt* –/– mice and *Mat1a* –/– mice relative to their WT littermates along with their functionally enriched pathways to provide a site specific PTM network view to shed light onto Lysine and Arginine controlled cascades that may be involved (Figure 6).

The resulting subnetwork of enriched pathways contained five key functional processes for both NASH mouse models including amino acid metabolism,^{34,77} lipid metabolism,^{34,77-79} PPAR signaling,⁷⁹ peroxisomal protein import,^{80,81} and glycolysis/gluconeogenesis,⁸² some of which were also enriched in the global proteome analysis (Figure 4A,B). PINE also provides the option to exclude ambiguous sites with oxidized methionines, miscleaved

versions, and ragged ends.^{59,60} In cases where PINE found ambiguity in PTM site identity, we indicated it using double asterisk (**) symbol next to the site. For example, BHMT has two dimethylated peptidoforms: one with oxidized methionine and one without. We also observed different PTMs localized to the same protein indicating possibility for modulating biochemical functions through positive and negative regulatory interactions.⁷⁷ For example, Betaine—homocysteine S-methyltransferase 1 (BHMT) is a well-known target for PTMs such as acetylation and trimethylation, which may be responsible for BHMT localization.⁸³ This enzyme plays an important role in regulation of hepatic lipid metabolism to protect the liver from triacylglycerol accumulation.^{34,64,84} A closer look at this protein revealed previously unreported lysine monomethylation and dimethylation events at site 179 that are regulated differently across the two NASH models, but their biological significance is yet to be determined. Understanding the crosstalk between these PTMs and regulation of their enzymatic mediators may represent a promising avenue for therapeutics in fatty liver disease.⁸⁵

Case Study 3: Exploring Protein Phosphorylation in the Context of Enriched Pathways and GO Terms.

Phosphoproteomics, or the large-scale determination of protein phosphorylation status, can be used to characterize and compare a variety of disease states but has been applied to NASH in only a handful of studies. For this use case, we performed global phosphoproteomics analysis using DDA from *Gnmt* –/– mice relative to WT littermates to look for differentially expressed phosphorylation sites to elucidate the role of kinase activity and regulation in NASH. Using PINE, we identified multiple enriched pathways including TGF-beta signaling, ^{86,87} insulin signaling, ^{88,89} glucagon signaling pathway, ^{88,89} ChREBP mediated metabolic gene expression, ^{88,89} bile acid and bile salt metabolism, ⁸⁹ signaling by Rho GTPases, ⁹⁰ EGFR1 signaling pathway, ⁹¹ AMPK signaling, ^{92,93} mTORC1-mediated signaling ^{42,94,95} and translation factors. ^{42,96} Since the complex interplay of AMPK and mTORC1 is rapidly gaining clinical relevance in metabolic disease, ⁹⁷⁻⁹⁹ we created a subnetwork focused around the crosstalk between mTORC1 and AMPK (Figure 7). AMPK activation has been known to enhance mitochondrial biogenesis and mitophagy and inhibit mTORC1, thus preventing excess-nutrient-induced hepatic lipid accumulation. ¹⁰⁰

Our phosphoproteome-based subnetwork confirmed these findings with several upregulated phosphosites from proteins involved in AMPK signaling as well as several downregulated phosphosites from proteins involved in mTORC1-mediated signaling and translation factors. ^{42,94-96} In order to take a complete look at all observed phosphosites associated with these pathways, we included both significant and nonsignificant differentially expressed sites indicated by blue node outline and no node outline, respectively. This network also revealed two significantly differentially expressed phosphosites (Threonine 214 and Threonine 507) of Eukaryotic translation initiation factor 4 gamma proteins (EIF4G1 and EIF4G2) that have been previously reported to be phosphorylated. ^{101,102} We noted that both phosphosites were down-regulated (adjusted *p*-value 0.05 and log2 fold change of –1.0) and have been shown to be involved in mTORC1 regulated translation control, mitosis and protein synthesis. Thus, AMPK could potentially act as a metabolic checkpoint by activating catabolic processes and inhibiting anabolic processes, in part, by negatively

regulating mTORC1 signaling and enhancing autophagy flux. ^{100,103,104} Although more clinical research is required to elucidate the role of AMPK and mTORC1 in NASH, novel therapies for metabolic disease could emerge from pharmacological manipulation of this interplay ^{97-99,103}

DISCUSSION

We have developed a fully automated, easy-to-use bioinformatics tool called PINE to aid researchers in extracting and visualizing differentially expressed proteins or PTMs and their interaction or functional networks from different types of mass-spectrometry data sets including DDA-, DIA-, and PTM-based experiments. PINE performs accurate protein identifier mapping to resolve ambiguity, retrieve PPIs from multiple publicly available interaction databases, and assimilate these complex networks into meaningful functionally enriched pathways visualized as snapshots of the proteome. Using two mouse models of NASH, we demonstrated that PINE can be used to gain biological insights that can be derived from linking observed protein modification changes to significantly altered protein pathways.

We acknowledge that there are certain areas for improvement within PINE, for example, we can increase support beyond the three commonly used organisms in proteomics profiling, i.e., Homo sapiens, Mus musculus, and Rattus norvegicus, as well, the number of public databases accessed for both interactions and annotations can be expanded along with improved filtering features based on confidence score or interaction sources. An interesting approach would be to compute edge weights based on how many database sources the interactions are present in or to apply the method carried out by Hierarchical HotNet¹⁰⁵ which simultaneously combines network interactions and vertex scores to construct a hierarchy of altered subnetworks. In the future extension of PINE, we also plan to improve network coverage by allowing synonym (or secondary) interactors and include upstream regulators to depict enzyme-substrate relationships. Identification of Single Amino Acid Polymorphisms (SAPs) could also be an important addition to this tool, playing a role in exploring how PTM sites containing SAPs affect underlying disease mechanisms. Additionally, due to the challenges involved in the unambiguous identification and localization of PTM, we recommend assessing the validity of these sites using tools such as Ascore, ¹⁰⁶ PTMProphet, ¹⁰⁷ LuciPHOr2, ¹⁰⁸ PIQUed, ¹⁰⁹ Peptide Collapse plugin for Perseus, ¹¹⁰ Inference of Peptidoforms (IPF), ¹¹¹ and Thesaurus ¹¹² prior to visualizing these data sets in PINE.

In summary, PINE provides a versatile, user-friendly, automated solution for network and pathway visualization specialized to handle bottom-up proteomics data. Using different case studies, we have demonstrated that PINE can seamlessly combine data from various public databases, while providing a variety of customization options, enabling profound exploration and visualization of global proteome and PTM relationships. Ultimately, PINE can serve as an extremely powerful tool for novel hypothesis generation toward understanding disease mechanisms and potential therapeutics.

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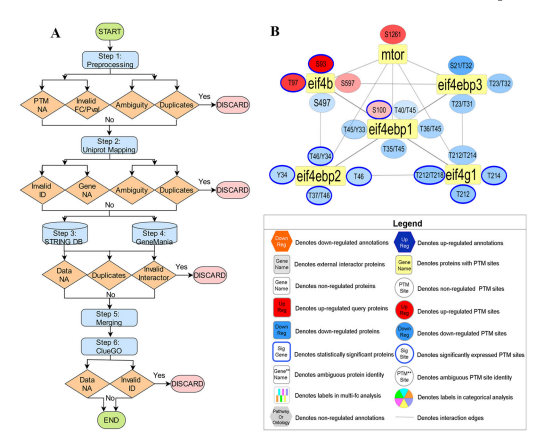


Figure 1.

Overview of PINE Framework. (A) PPIs derived from public databases (e.g., STRING, GeneMANIA) and functional enrichment-based network annotations from ClueGO are processed by the Cytoscape framework to create PPIs networks. By using PINE, protein or PTM associated differential expression (e.g., fold change) or statistical significance data (e.g., p-value) can also be overlaid onto the Cytoscape networks. (B) Example PPIs network depicting both statistically significant and nonsignificant PTM sites. The rectangular yellow nodes denote proteins (represented by their primary genes), connected to circular nodes representing PTM sites. The color refers to either down-regulated (blue) or up-regulated (orange) sites with statistically significant sites outlined blue. The legend describes the various node types and colors used in visualization of PPI networks

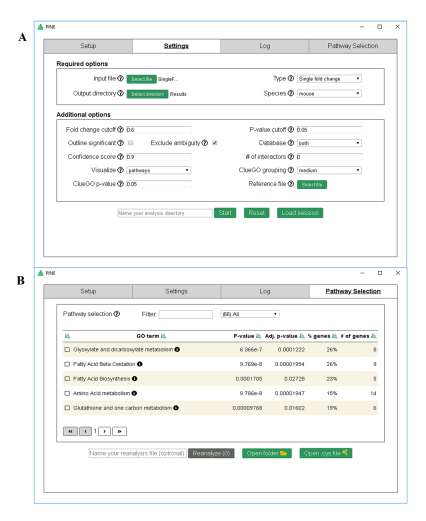


Figure 2.
PINE graphical user interface. (A) Screenshot illustrating the settings tab for easy query upload, providing users the choice to modify parameters for PPIs and annotation analysis.
(B) Screenshot illustrating the Pathway Selection tab that lists annotation terms, additionally providing the option to filter, sort, and subsequently select annotation terms of interest in order to generate ontology network.

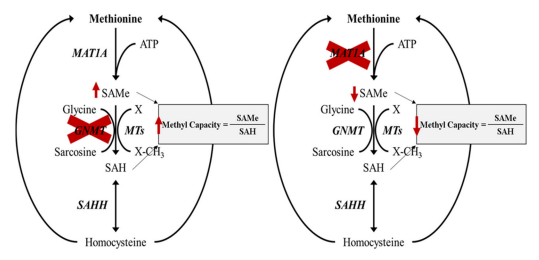
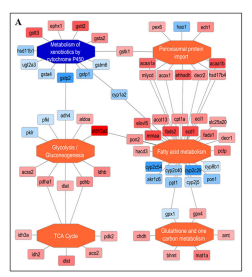


Figure 3.Hepatic methionine metabolism in different NASH models. Two mouse models of nonalcoholic steatohepatitis (NASH): *Gnmt* –/– mice (left) have increased hepatic SAMe level and methylation capacity, being biologically hypermethylated, while *Mat1a* –/– mice (right) are SAMe deficient with reduced methylation capacity and therefore hypomethylated.



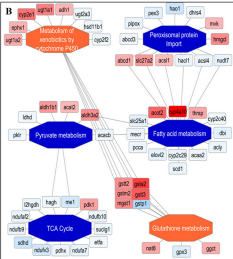


Figure 4.Enriched functional networks generated by PINE across different NASH models. (A) The *Gnmt* –/– mice model indicates five activated pathways shown as orange central nodes and the one inhibited pathway shown as blue central nodes along with fold-changes from the different protein nodes. (B) The *Mat1a* –/– mice model indicates two activated pathways shown as orange central nodes and the four inhibited pathway shown as blue central nodes along with fold-changes from the different protein node.

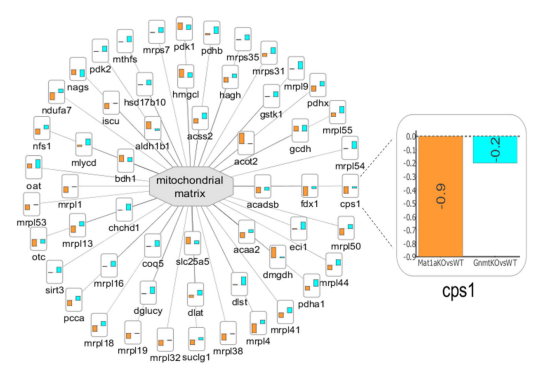


Figure 5.
Enriched cellular subnetwork for mitochondrial matrix generated by PINE across different NASH models. Altered cellular components are denoted by central gray nodes. Differential expression data from multiple comparison groups are represented as bar charts and the bar height indicates degree of fold change. (Example: Zoomed view into node CPS1 indicates degree of fold change for *Mat1a* –/– mice vs WT littermates shown using orange bar and *Gnmt* –/– mice vs WT littermates shown using blue bar.)

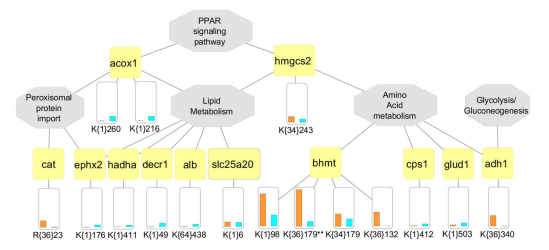


Figure 6.

Multi-PTM network illustrating crosstalk between methylation, acetylation and succinylation. Functionally enriched pathways are shown by central gray nodes along with gene connections shown by yellow nodes. The PTM site level quantitative differences from each mouse model are shown as a bar chart (*Gnmt* –/– mice indicated by blue bar and *Mat1a* –/– mice by orange bar). Sites are indicated with unimod accession in curly brackets (Example, K{34}179 and K{36}179 on gene BHMT would indicate lysine monomethylation and lysine dimethylation events on site 179). Ambiguous sites are indicated using double asterisk (**) symbol next to the site. (Example, K{1}110** on gene GLUD1.)

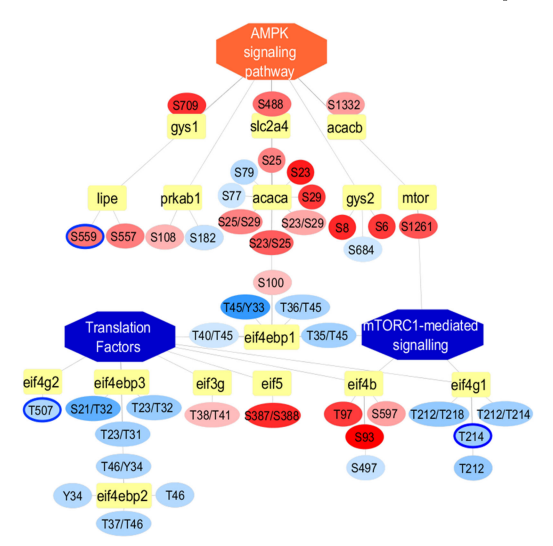


Figure 7.

Phospho-proteome network illustrating significant differentially expressed PTM sites using PINE. Inhibited pathways such as mTORC1-mediated signaling and translation factors are represented as central blue nodes whereas activated pathway such as AMPK signaling pathway are represented as central orange nodes along with gene connections represented as yellow nodes. The PTM site level quantitative differences between *Gnmt* –/– mice relative to WT littermates are represented as blue for downregulated nodes or orange for upregulated nodes connected to their respective gene nodes. We included both significant and nonsignificant differentially expressed sites indicated by blue node outline and no node outline, respectively.