Characterizing the Nitroproteome Using Bioinformatic Tools: A Mini-Review

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Abstract

Protein tyrosine nitration (PTN) is a post-translational modification that occurs due to the action of reactive nitrogen species (RNS). The specific tyrosine residues are mainly nitrated in the presence of nitrating agents. It is also related to neurodegenerative diseases, cellular signaling. So the characterization of nitroproteome with their functions is very much essential. In recent times bioinformatic tools become a powerful apparatus to characterize the biological objects. In this mini-review we have attempted to give an outline about the bioinformatic tools which can be used to characterize the nitration/denitration system.

Keywords: Protein tyrosine nitration; Reactive nitrogen species; Nitroproteome; Bioinformatic tools

Abbreviation: RNS: Reactive Nitrogen Species; NO$_2$-TYR: 3-Nitrotyrosine; PTN: Protein Tyrosine Nitration

Introduction

Nitrosative stress is a condition in which the cellular redox homeostasis is changed because of excessive production of different reactive nitrogen species (RNS) [1]. Under nitrosative stress the ratio of nitrosants to antioxidants is always >1. One of the major markers of nitrosative stress is the 3-Nitrotyrosine (NO$_2$-TYR), a stable post-translational modification of protein. 3-Nitrotyrosine forms due to the reaction of tyrosine and nitrating agents. A nitro (-NO$_2$) group is added in the ortho position of the phenolic hydroxyl group of tyrosine during the reaction of tyrosine and nitrating agent which results in the formation of 3-Nitrotyrosine (NO$_2$-TYR). Generally the natural abundance of tyrosine residues is about 3% in the proteins. But the nitration may introduce negative charge at neutral pH which may results in the change of the local physiological and chemical environment of the biomolecules. Due to this event the structure and function of the proteins are also altered. Thus, the cellular mechanism is also changed. 3-Nitrotyrosine has an impact on clinical biology [2]. It is related with the cell signaling and disease initiation and progression like neurodegenerative diseases, cancer and cardiovascular injury. Alzheimer’s disease, the most common neurodegenerative disease is associated with the formation of 3-Nitrotyrosine. Alzheimer’s disease is induced by accumulation of nitrated tau and mis folded αβ proteins in the brain [3]. It is also reported that proteins like alcholo dehydrogenase [4], aldolase [5], isocitrtae dehydrogenase [6] can be nitrated during nitrosative stress. Some enzymes are also present to counteract the niosative insult e.g. catalase [7], Cytochrome C [8] etc. but there are some reports that suggest these stress response enzymes can also be nitrated in the presence of nitrating agents. Interestingly, some reports suggest that PTN is also related to cellular signaling. The proteins of the mating signaling pathway can be nitrated in Saccharomyces cerevisiae [9]. So this event gives an idea about denitration. But the harmony between the balance of nitration and denitration is yet to be elucidated.

Thus, it is assumed that the nitration/de nitration pathway is just phosphorylation/dephosphorylation. So right now one of the major challenges for biologist is to characterize the nitration/denitration pathway. For this purpose one of the most important tool is bioinformatic. In this study we have tried to give an outline about the bioinformatics tools which can be used to understand the biochemistry of protein tyrosine nitration and de nitration.

Bioinformatic tools

Bioinformatic tools are used in the proteomics study not only to characterize the nitroproteins, their structure and function but this specific [10]; sensitive process is also related to the secondary structure of protein, Solvent accessibility [11], and protein-protein interactions [12]. The study of protein tyrosine nitration/denitration is started from the identification of nitroproteins. The mostly used software to identify the nitroproteins is Turbo SEQUEST [10]. One of the important software is Scaffold software which helps to compute protein
and peptide probability for nitration [13]. MS-BLAST is also applied to identify the homolog's of different nitro proteins which needs de novo correct tagging. To ensure the correct tagging SPIDER software is used where de novo sequencing and homology mutations are taken into account [14].

The secondary structure of protein is needed to be characterized to identify the nitration-prone tyrosine residues. PSIPRED (Software for Protein Identification from Sequence Tags with De Novo Sequencing Error), JNET, PROF Etc. [15-19]. Algorithms are used to characterize the secondary structure. Neutral networking is one of the best methods to characterize the secondary structure. JPred4 is the latest version of JPred to predict the secondary structure of proteins. JNet algorithm is used in JPred4 software. JNet algorithm is an error-free method which results in the higher accuracy of JPred 4 [20]. Useful software for characterizing the PTN site is GPS-YNO having high accuracy [21]. Solvent accessibility another important factor to characterize the protein structure is efficiently predicted by SANN, a nearest neighbor method [22].

Protein networking another important bioinformatics method used to determine the role and interaction of a protein in certain protein pool. PIN (protein interaction network) and PSN (Protein-Signaling Network) are two major network models for proteomics. The post translational modifications are actually studied by PSN whereas PIN is based on the protein-protein bindings [12]. The databases for protein interactions are DIP, BIND, MIPS, and MINT etc. One of the strongest software JGIP is used in DIP data model which consists of binary protein interactions. The information about the interacting proteins is also got from DIP data model [23]. DIP is mainly used for budding yeast because a huge amount of external information for budding yeast is only available now [24]. One of the largest collections of freely available information about pair wise molecular interactions and complexes is BIND (Bimolecular Interaction Network Database). The interaction between biological objects like DNA, RNA, gene, protein is characterized by BIND. The specification of BIND data is available as XML DTD and ASN.1 [25]. The indirect and genetic interaction is predicted by MINT (The Molecular Interaction database). Information regarding the modification of enzymes, their kinetics and their binding domains are stored by MINT [26]. The dataset of high-quality experimental protein-protein interaction in mammals is MIPS mammalian protein–protein interaction database (MPPi) [27]. The increasing data content is a problem for visualization. Cytoscape, stronger freely available, open-source java-based network visualization and analysis tool is used for visualization [28].

These softwares may be helpful to characterize the protein tyrosine nitration and denitration system. Protein networking is the choice of bioinformatic tool to establish the probable pathway because it is hypothesized that several enzymes are involved in denitration system. Prediction of secondary structure and Solvent accessibility are also very important to predict the structure and stability of the protein. The probable model for nitration/denitration is outlined in Figure 1.

**Conclusion**

PTN is a topic of ongoing research. PTN has the drastic effect on neurodegenerative diseases. Nitration of neuroproteins results in the tangle formation. The available medicine for these diseases is very few and having side effects. So it is very important to characterize the denitration system. Denitration is the mechanism which can reverse back the nitration, this reaction results in the prevention of tangle formation. So “denitrase” enzyme can be used as the therapeutic medicine for neurodegenerative diseases. But still a huge area of research is needed to understand the nitration/denitration properly.

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**References**


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