



Article

PredNTS: Improved and Robust Prediction of Nitrotyrosine Sites by Integrating Multiple Sequence Features

Andi Nur Nilamyani ¹, Firda Nurul Auliah ¹, Mohammad Ali Moni ², Watshara Shoombuatong ³, Md Mehedi Hasan ^{1,4},* and Hiroyuki Kurata ^{1,*}

- Department of Bioscience and Bioinformatics, Kyushu Institute of Technology, 680-4 Kawazu, Iizuka, Fukuoka 820-8502, Japan; nurnilamyani.514@gmail.com (A.N.N.); Firdana.525@gmail.com (F.N.A.)
- WHO Collaborating Centre on eHealth, UNSW Digital Health, School of Public Health and Community Medicine, Faculty of Medicine, UNSW Sydney, Sydney, NSW 2052, Australia; m.moni@unsw.edu.au
- Center of Data Mining and Biomedical Informatics, Faculty of Medical Technology, Mahidol University, Bangkok 10700, Thailand; watshara.sho@mahidol.ac.th
- ⁴ Japan Society for the Promotion of Science, 5-3-1 Kojimachi, Chiyoda-ku, Tokyo 102-0083, Japan
- * Correspondence: hasan.md-mehedi922@mail.kyutech.jp (M.M.H.); kurata@bio.kyutech.ac.jp (H.K.)

Abstract: Nitrotyrosine, which is generated by numerous reactive nitrogen species, is a type of protein post-translational modification. Identification of site-specific nitration modification on tyrosine is a prerequisite to understanding the molecular function of nitrated proteins. Thanks to the progress of machine learning, computational prediction can play a vital role before the biological experimentation. Herein, we developed a computational predictor PredNTS by integrating multiple sequence features including K-mer, composition of k-spaced amino acid pairs (CKSAAP), AAindex, and binary encoding schemes. The important features were selected by the recursive feature elimination approach using a random forest classifier. Finally, we linearly combined the successive random forest (RF) probability scores generated by the different, single encoding-employing RF models. The resultant PredNTS predictor achieved an area under a curve (AUC) of 0.910 using five-fold cross validation. It outperformed the existing predictors on a comprehensive and independent dataset. Furthermore, we investigated several machine learning algorithms to demonstrate the superiority of the employed RF algorithm. The PredNTS is a useful computational resource for the prediction of nitrotyrosine sites. The web-application with the curated datasets of the PredNTS is publicly available.

Keywords: nitrotyrosine; post-translational modification; feature encoding; RFE feature selection; machine learning



Citation: Nilamyani, A.N.; Auliah, F.N.; Moni, M.A.; Shoombuatong, W.; Hasan, M.M.; Kurata, H. PredNTS: Improved and Robust Prediction of Nitrotyrosine Sites by Integrating Multiple Sequence Features. *Int. J. Mol. Sci.* 2021, 22, 2704. https://doi.org/10.3390/ijms22052704

Academic Editor: Alexandre G. de Brevern

Received: 21 January 2021 Accepted: 3 March 2021 Published: 8 March 2021

Publisher's Note: MDPI stays neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https://creativecommons.org/licenses/by/4.0/).

1. Introduction

Nitrotyrosine, which is generated by numerous reactive nitrogen species, is a type of protein post-translational modification. It occurs when tyrosine is nitrated by reactive nitrogen species (RNS) such as peroxynitrite anions (ONOO $^-$), which are carried out in vivo through the rapid reaction of nitric oxide (NO $^-$) and superoxide (O $_2$ $^-$) [1–3]. Nitration of proteins changes their chemical properties. Excessive levels of peroxynitrite anion are observed during the inflammation process in a large number of diseases including diabetics, cancer, neurodegenerative disorders, asthma, and ageing [4,5].

Tyrosine nitration occurs within a protein interaction region such as enzyme—substrate or receptor—ligand complexes and brings several effects, like decreasing the electron intensity of the phenolic ring of tyrosine, and negatively affects their interactions. It modifies the enzymes and receptors, reducing their biological activities [2,6]. Besides, tyrosine nitration may interfere directly with the phosphorylation of tyrosine residues responsible for cellular signalling pathways [7,8]. Therefore, potential guidance for developing new therapeutic

strategies and drugs can be obtained through the identification of site-specific nitration modifications on tyrosine.

To reveal the mechanism and function of nitroprotein, identification of potential nitrotyrosine sites is essential. To date, large-scale proteomic studies have been performed to identify nitrated proteins based on the molecular signature of nitrotyrosine sites [3,5,8]. Although the number of experimentally verified nitrotyrosine sites is increasing, the mechanism of site-specific nitration modification on tyrosine remains largely unknown [9–11], probably owing to technological measurement constraints. The conventional experimental methods provide insights into a biological role of nitrotyrosine sites, but they are very time-consuming and expensive. Therefore, as an alternative strategy, an in silico approach can be proposed to predict nitrotyrosine sites, which functions for all proteome annotations because of their efficiency and convenience.

Until now, only a few predictors have been presented to identify nitrotyrosine sites [12–15]. Initially, Liu et al. [15] developed a predictor based on a group-based prediction system (GPS) using four statistical procedures (i.e., selection of motif length, K-means clustering, matrix mutation, and weight training), named GPS-YNO2. Xie et al. [12] developed the DeepNitro that implemented deep learning with four encoding schemes (i.e., positional amino acid distributions, sequence contextual dependencies, physicochemical properties, and position-specific scoring features). The NTyroSite [13] was constructed using sequence evolutionary information. The iNitro-Tyr was developed based on the pseudo amino acid composition [14]. From the results of previous studies, it is known that the previous predictors use training datasets and receive good performance by five-fold cross-validation (CV) tests. In this study, we have developed a computational predictor PredNTS by integrating multiple sequence features of K-mer, composition of k-spaced amino acid pairs (CKSAAP), AAindex, and binary. A workflow of the PredNTS is shown in Figure 1. We implemented the recursive feature elimination (RFE) to select the important features via a random forest (RF) classifier. Finally, the successive RF scores were combined with a linear regression model. A user-friendly web server was developed and is freely available at http://kurata14.bio.kyutech.ac.jp/PredNTS/ (accessed on 1 March 2021).

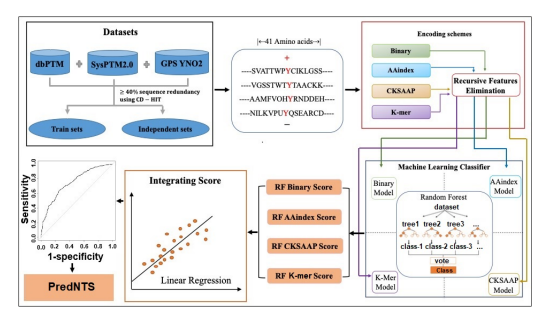


Figure 1. A framework for the PredNTS predictor and its server application. GPS, group-based prediction system; RF, random forest; RFE, recursive feature elimination; CKSAAP, composition of k-spaced amino acid pairs.

2. Results and Discussion

2.1. Sequence Preference Analysis

We visualized the curated positive and negative samples using a graphical sequence logo to check the significant preference of the amino acid residues surrounding nitrotyrosine proteins [16], as shown in Figure 2. Some significant differences of amino acid sequences were detected between the positive and negative samples. We found that charged residues such as K, R, and E frequently appeared in the enriched positions, while Y, S, F, and L were frequently observed in the depleted section. However, in the depleted section, no stacked residue was found at positions of -16, +3, and +13. The above analysis of amino acid residue preference between the positives and negatives suggested that a combination of frequency-based encodings with position specific encodings is effective in designing nitrotyrosine site prediction.



Figure 2. Sequence preference analysis of positive and negative samples of nitroproteins.

2.2. Single Encoding-Employing RF Model on the Training Dataset

We used the four encoding schemes (AAIndex, binary, CKSAAP, and K-mer) to generate numerical feature vectors. The window size was set to $41 \, (-/+20)$ for all the encoding schemes. The prediction performances were measured using five-fold CV through the RF classifier. The average performance of the four single encoding-employing RF models without any feature selection is summarized in Table 1. Without any feature selection, the K-mer encoding performed better than any other encodings, which achieved Ac of 0.796 and Matthew's correlation coefficient (MCC) of 0.593.

Table 1. Prediction performance of the single encoding-employing models without any feature selection on the training dataset by five-fold cross-validation (CV). CKSAAP, composition of k-spaced amino acid pairs. MCC, Matthew's correlation coefficient.

Encoding Scheme	Sn	Sp	Acc	MCC
Binary	0.519	0.808	0.658	0.331
	(0.19)	(0.01)	(0.04)	(0.14)
AAindex	0.473	0.803	0.638	0.293
	(0.24)	(0.00)	(0.02)	(0.16)
CKSAAP	0.731	0.808	0.709	0.519
	(0.16)	(0.00)	(0.11)	(0.20)
K-mer	0.785	0.807	0.796	0.593
	(0.11)	(0.01)	(0.06)	(0.13)

Values in parentheses represent the standard error (SE).

Note that the high-dimensional features may contain irrelevant or redundant attributes that affect accuracy reduction [17,18]. To discriminate the relative contribution and importance of each feature, the RFE method was considered. Different feature subsets were selected for each encoding, which controlled the high ranked features ranging from the top 50 to all with an interval of 50. The curated subset features were inputted to RF separately and their individual performances were estimated using five-fold CV (Figure S1). This approach selected the 400- for AAindex, 300- for binary, 200- for CKSAAP, and 500-dimesional

Int. J. Mol. Sci. 2021, 22, 2704 4 of 11

features for K-mer. Then, we measured the four statistical measures of Sp, Sn, Ac, and MCC via a five-fold CV test on the training dataset, as shown in Table 2. Use of the RFE improved the performance of our models. In the model with feature selection, the Acc was ~2% to 4% higher than the model without any feature selection. Figure 3A shows the ROC curves for the four single encoding-employing models with feature selection on the training datasets. The CKSAAP and K-mer encodings provided better prediction than the other two encoding schemes. The CKSAAP and K-mer encodings achieved AUCs of 0.900 and 0.895, respectively, while the binary and AAindex encodings provided AUCs of 0.773 and 0.771, respectively. The feature selection improved the performance for all the encoding schemes (binary, AAindex, CKSAAP, and K-mer).

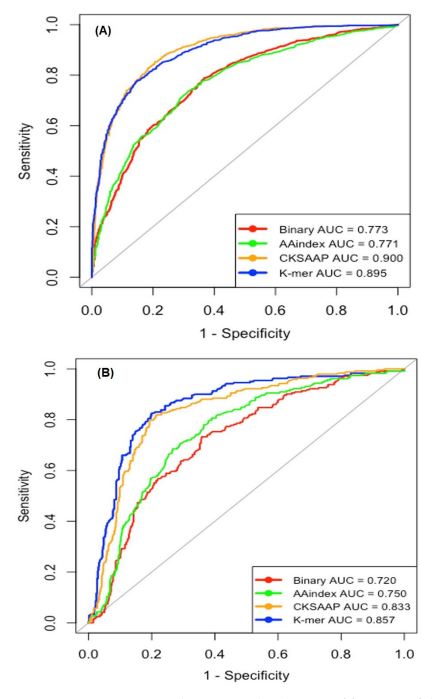


Figure 3. Receiver operating characteristics (ROC) curves of four types of the single encoding-employing models. (A) On the training dataset using five-fold cross-validation test. (B) On the independent test dataset. AUC, area under the curve.

Int. J. Mol. Sci. **2021**, 22, 2704 5 of 11

Table 2. Prediction performance of the single encoding-employing models with feature selection or	ı
the training dataset by five-fold CV.	

Encoding Scheme	Sn	Sp	Acc	MCC
Binary	0.598	0.800	0.699	0.407
	(0.18)	(0.00)	(0.02)	(0.12)
AAindex	0.571	0.809	0.690	0.391
	(0.19)	(0.01)	(0.07)	(0.17)
CKSAAP	0.829	0.809	0.819	0.639
	(0.09)	(0.00)	(0.09)	(0.14)
K-mer	0.811	0.808	0.810	0.619
	(0.06)	(0.00)	(0.11)	(0.08)

Values in parentheses represent the standard error (SE).

2.3. Single Encoding-Employing RF Model on the Independent Dataset

We used an independent dataset to investigate the robustness of the training models. The performances of the models without any feature selection and the models with feature selection were evaluated on the independent dataset, as shown in Tables 3 and 4, respectively. The use of RFE improved the prediction performance of nitrotyrosine sites. As shown in Figure 3B, the CKSAAP and K-mer encodings achieved AUCs of 0.833 and 0.857, respectively, while the binary and AAindex encodings provided AUCs of 0.720 and 0.750, respectively. The CKSAAP and K-mer encodings performed better than the binary and AAindex encodings.

Table 3. Prediction performance of the single encoding-employing models without any feature selection on the independent dataset.

Encoding Scheme	Sn	Sp	Acc	MCC
Binary	0.384	0.806	0.736	0.170
AAindex	0.397	0.800	0.733	0.174
CKSAAP	0.458	0.800	0.743	0.224
K-mer	0.480	0.800	0.747	0.242

Table 4. Prediction performance of the single encoding-employing models with feature selection on the independent dataset.

Encoding Scheme	Sn	Sp	Acc	MCC
Binary	0.445	0.801	0.742	0.214
AAindex	0.438	0.801	0.741	0.209
CKSAAP	0.504	0.805	0.755	0.268
K-mer	0.532	0.804	0.758	0.288

2.4. Prediction Performance of PredNTS

To build the PredNTS, we linearly combined the probability scores generated by the four types of single encoding-employing RF models. We optimized the weight coefficients for the binary, AAindex, CKSAAP, and K-mer encodings as 0.01, 0.01, 0.3, and 0.68, respectively. As shown in Table 5, the PredNTS achieved an AUC of 0.910 on the training dataset by five-fold CV, while it achieved an AUC of 0.860 on the independent dataset. The integration of the four encoding schemes greatly improved the prediction performance. To validate the superiority of the RF employed by the PredNTS, we compared it with the two machine learning algorithms of naïve Bayes (NB) and k-nearest neighbor (KNN). Here, we employed the same number of selected features and the same window size of 41. The performances of the three machine learning algorithms were compared for the combined models without and with feature selection, respectively, as shown in Figure 4A,B. The AUCs of the PredNTS were 3–6% higher than those of the NB and KNN implementing

Int. J. Mol. Sci. **2021**, 22, 2704 6 of 11

combined models. The RF outperformed the NB and KNN, demonstrating the superiority of the RF.

Table 5. Prediction performance of the PredNTS with feature selection.

Dataset	Predictor	AUC
Training dataset	Binary + AAindex + CKSAAP + K-mer	0.910
Independent dataset	Binary + AAindex + CKSAAP + K-mer	0.860

The weight coefficients with respect to the four types (binary, AAindex, CKSAAP, and K-mer) of the single encoding-employing models are 0.01, 0.01, 0.3, and 0.68, respectively.

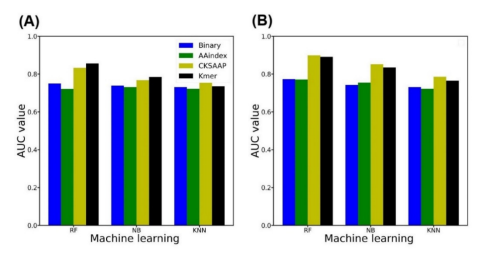


Figure 4. AUC values of the single encoding-employing models with different machine learning algorithms on training datasets. **(A)** Performance comparison of the models without any feature selection. **(B)** Performance comparison of the models with feature selection. NB, naïve Bayes; KNN, k-nearest neighbor; RF, random forest.

2.5. Comparison of PredNTS with Other Existing Predictors

Several computational predictors for nitrotyrosine sites have been developed. We compared the PredNTS predictor with the three existing predictors: GPS-YNO2 [14], DeepNitro [12], and NTyroSite [15]. The comparison was carried using the independent dataset with 203 positive samples and 1022 negative samples. The curated independent datasets were submitted to the GPS-YNO2, DeepNitro, and NTyroSite online servers, then the performance was assessed by the four statistical measures (Sn, Sp, Acc, and MCC). As seen in Table 6, our predictor presented much better performance than the GPS-YNO, DeepNitro, and NTyroSite predictors in terms of Sn, Sp, Ac, and MCC. The PredNTS achieved 0.522 for Sn, 0.809 for Sp, 0.761 for Acc, and 0.286 for MCC. The PredNTS presented significantly higher MCC than the other predictors. This might be caused by the fact that the GPS-YNO2 did not use any independent dataset to evaluate its robustness and that the DeepNitro and NTyroSite did not integrate their encoding schemes.

Table 6. Performance comparison of the PredNTS with the three existing predictors on the independent dataset.

Encoding Scheme	Sn	Sp	Acc	MCC
GPS-YNO2	0.334	0.801	0.724	0.122
DeepNitro	0.339	0.803	0.726	0.128
NTyroSite	0.440	0.793	0.744	0.196
PredNTS	0.522	0.809	0.761	0.286

2.6. Web Server Implementation

The PredNTS webserver was developed to serve potential user communities and is freely available at http://kurata14.bio.kyutech.ac.jp/PredNTS/ (accessed on 1 March

2021). On the main page, users submit a query protein sequence by pasting it into the text box or using the browse button. The server initially calculates the window size based on the number of tyrosine residues. In the meantime, the server generates the PSSM by performing the PSI-BLAST search for the query sequence and encodes the sequence windows. After selecting the important features using the RFE method, the server classifies the feature vectors using the RF algorithm. The webserver returns the predicted results containing the residue positions, request protein name, job ID, and probability scores on the output webpage. The server creates a job ID such as "20200102100011". Users can save this ID for the future enquiry. The PredNTS implementation can handle only FASTA format sequences.

3. Materials and Methods

3.1. Dataset Construction

We collected the datasets from the different public sources including DeepNitro and iNitro-Tyr [12,14]. The experimentally identified nitrotyrosine sites ("Y", tyrosine residue) were considered as positive samples, whereas the resting Y residues were measured as negative samples [19–21]. It contains 796 nitrotyrosine proteins with 1406 experimentally validated nitrotyrosine sites. A sequence window with a length of 2w + 1 was prepared so as to place nitrotyrosine in the centre. We removed redundant sequences by considering a threshold of 40% level by CD-HIT [22]. Finally, randomly selected 20% of the samples (203 positive samples and 1022 negative samples) were considered as the independent dataset to examine the model strength. From the whole remaining dataset, we pooled a 1:1 ratio of positive to negative samples (1191 positive samples and 1191 negative samples) as the training model to avoid possible biased predictions. The independent (203 positive samples and 1022 negative samples) dataset was used to compare the proposed PredNTS model with existing predictors.

3.2. Sequence Encoding Scheme

The binary amino acid encoding scheme was used to encode position information from the sequence windows [23–25]. Here, by adopting the binary encoding, we converted a 41 amino acid sequence, including the gap that is represented as (-), into a 861 (=41 \times 21)-dimensional feature vector.

The physicochemical properties of amino acids have been extracted from the AAindex database 24 (version 9.1) [26]. Herein, we used 15 types of AAindex properties to generate a 615 (= 41×15)-dimensional vector.

The composition of k-spaced amino acid pairs (CKSAAP) encoding is the composition of the k-spaced residue pairs in the window, which is widely used in a protein bioinformatics field [14,20,27]. In this scheme, k represents the gap length of two amino acids. For example, k = 0 provides 400 amino acid residue pairs (i.e., AA, AC, AD, ..., YY). At k = 0, 1, 2, 3, and 4, it generates a 2000-dimensional feature vector. Details of the CKSAAP encoding are described in our previous studies [14,25].

The K-mer encoding is widely used in the field of genomics and bioinformatics [23,28–31]. We employed the K-mer to minimize the impact of an arbitrary starting point. The K-mer encodes a monopeptide into a 20-dimensional feature vector at K = 1. Similarly, at K = 2 and 3, it encodes dipeptides and tripeptides, which generates an 8020-dimensional feature vector.

3.3. Feature Selection

We considered the RFE as a feature selection approach to remove non-essential features from the dataset [32]. This method was classified as a wrapper method, which started from building a learning model for the entire dataset. We calculated the important scores from each predictor and trimmed the least important features out of the current set of features. The procedure is repeated until the number of optimal performance features converges. The 'rfe' function from the 'Caret' R package was adopted to obtain the important features.

3.4. Machine Learning Algorithm

The RF is a supervised and ensemble machine learning classifier that combines multiple tree-based representations to create a more powerful and interpretable model. It is widely used in protein bioinformatics research [33–43]. It performs as a huge assortment of uncorrelated decision trees, and the votes are carried to decide the final classification from the whole trees. The prediction model of our PredNTS was built using the 'RandomForest' R package (https://cran.r-project.org/web/packages/randomForest/ (accessed on 1 March 2021)). In addition, we compared the RF algorithm with the naïve Bayes (NB) and k-nearest neighbor (KNN) algorithms. An R package of an NB algorithm (https://cran.r-project.org/web/packages/naivebayes/ (accessed on 1 March 2021)) was employed to classify the nitrotyrosine proteins, while the R package (https://rpubs.com/njvijay/16444 (accessed on 1 March 2021)) was used to build the KNN model.

3.5. Evaluation Measure

Our PredNTS model is evaluated through five-fold CV. This method starts from randomly selecting and partitioning the training dataset into five sub-folds, then those five sub-folds are divided into the training and test datasets alternately. Four folds are used as the training sets; the remaining one fold is used as the test set. This process is repeated five times in order to measure the entire dataset. Four simple statistical measures of specificity (Sp), sensitivity (Sn), accuracy (Acc), and Matthew's correlation coefficient (MCC) [44–59] are considered to evaluate the prediction performance of the model, as follows:

$$Sn = \frac{TP}{TP + FN} \tag{1}$$

$$Sp = \frac{TN}{TN + FP} \tag{2}$$

$$Ac = \frac{TP + TN}{TP + TN + FP + FN} \tag{3}$$

$$MCC = \frac{(TP \times TN) - (FP \times FN)}{\sqrt{[TP + FN][TN + FP][TP + FP][TN + FN]}}$$
(4)

where TP is the true positive, FP is the false positive, TN is the true negative, and FN is the false negative. The statistical values of (Sn, Sp, Acc) are between 0 and 1 and MCC is -1 to 1. Prediction accuracy is indicated by a high score. In addition, to obtain the area under the curve (AUC), we plot the receiver operating characteristics (ROC) curve, which is used to measure the overall ability of a classifier; the pROC package in the R language (https://cran.r-project.org/web/packages/pROC/ (accessed on 1 March 2021)) is used for this process.

4. Conclusions

We have developed a computational predictor PredNTS that linearly combined the probability scores generated by multiple single encoding-employing RF models. The critical features were selected by the RFE approach using RF. The employed RF algorithm was shown to be superior to other machine learning algorithms. On both the training and independent datasets, the PredNTS achieved excellent prediction performances, outperforming existing state-of-the-art predictors. The PredNTS is a useful computational resource for the prediction of nitrotyrosine sites, and is freely available at http://kurata14.bio.kyutech.ac.jp/PredNTS/ (accessed on 1 March 2021).

Supplementary Materials: The following are available online at https://www.mdpi.com/1422-006 7/22/5/2704/s1, Figure S1: AUC value with respect to selected features by RFE for the four encoding schemes.

Author Contributions: Conceptualization, M.M.H. and H.K.; For analysis, A.N.N., F.N.A., and M.M.H.; Methodology, A.N.N. and M.M.H.; Writing—original draft, F.N.A., M.M.H., and H.K.; Writing—review and editing, A.N.N., M.A.M., W.S., M.M.H., and H.K. All authors have read and agreed to the published version of the manuscript.

Funding: This work was supported by the Grant-in-Aid for Scientific Research (B) (19H04208) and partially supported by the Grant-in-Aid for JSPS Research Fellow (19F19377) from Japan Society for the Promotion of Science (JSPS).

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: All the data are available at http://kurata14.bio.kyutech.ac.jp/PredNTS/.

Conflicts of Interest: The authors declare that they have no conflict of interest.

References

 Zhao, Y.; Zhang, Y.; Sun, H.; Maroto, R.; Brasier, A.R. Selective Affinity Enrichment of Nitrotyrosine-Containing Peptides for Quantitative Analysis in Complex Samples. J. Proteome Res. 2017, 16, 2983–2992. [CrossRef]

- 2. Peng, F.; Li, J.; Guo, T.; Yang, H.; Li, M.; Sang, S.; Li, X.; Desiderio, D.M.; Zhan, X. Nitroproteins in Human Astrocytomas Discovered by Gel Electrophoresis and Tandem Mass Spectrometry. *J. Am. Soc. Mass Spectrom.* **2015**, *26*, 2062–2076. [CrossRef]
- 3. Nuriel, T.; Deeb, R.S.; Hajjar, D.P.; Gross, S.S. Protein 3-nitrotyrosine in complex biological samples: Quantification by high-pressure liquid chromatography/electrochemical detection and emergence of proteomic approaches for unbiased identification of modification sites. *Methods Enzym.* **2008**, *441*, 1–17. [CrossRef]
- 4. Seeley, K.W.; Fertig, A.R.; Dufresne, C.P.; Pinho, J.P.; Stevens, S.M., Jr. Evaluation of a method for nitrotyrosine site identification and relative quantitation using a stable isotope-labeled nitrated spike-in standard and high resolution fourier transform MS and MS/MS analysis. *Int. J. Mol. Sci.* **2014**, *15*, 6265–6285. [CrossRef]
- 5. Lee, S.J.; Lee, J.R.; Kim, Y.H.; Park, Y.S.; Park, S.I.; Park, H.S.; Kim, K.P. Investigation of tyrosine nitration and nitrosylation of angiotensin II and bovine serum albumin with electrospray ionization mass spectrometry. *Rapid Commun. Mass Spectrom.* 2007, 21, 2797–2804. [CrossRef]
- 6. Ghesquiere, B.; Goethals, M.; Van Damme, J.; Staes, A.; Timmerman, E.; Vandekerckhove, J.; Gevaert, K. Improved tandem mass spectrometric characterization of 3-nitrotyrosine sites in peptides. *Rapid Commun. Mass Spectrom.* **2006**, 20, 2885–2893. [CrossRef] [PubMed]
- 7. Gerding, H.R.; Karreman, C.; Daiber, A.; Delp, J.; Hammler, D.; Mex, M.; Schildknecht, S.; Leist, M. Reductive modification of genetically encoded 3-nitrotyrosine sites in alpha synuclein expressed in *E. coli. Redox Biol.* **2019**, *26*, 101251. [CrossRef] [PubMed]
- 8. Zhan, X.; Desiderio, D.M. Mass spectrometric identification of in vivo nitrotyrosine sites in the human pituitary tumor proteome. *Methods Mol. Biol.* **2009**, *566*, 137–163. [CrossRef] [PubMed]
- 9. Moraes, J.B.; Maes, M.; Roomruangwong, C.; Bonifacio, K.L.; Barbosa, D.S.; Vargas, H.O.; Anderson, G.; Kubera, M.; Carvalho, A.F.; Nunes, S.O.V. In major affective disorders, early life trauma predict increased nitro-oxidative stress, lipid peroxidation and protein oxidation and recurrence of major affective disorders, suicidal behaviors and a lowered quality of life. *Metab. Brain Dis.* **2018**, 33, 1081–1096. [CrossRef] [PubMed]
- 10. Xiang, T.; Zhang, S.; Cheng, N.; Ge, S.; Wen, J.; Xiao, J.; Wu, X. Oxidored-nitro domain-containing protein 1 promotes liver fibrosis by activating the Wnt/beta-catenin signaling pathway in vitro. *Mol. Med. Rep.* **2017**, *16*, 5050–5054. [CrossRef]
- Ha, M.N.; Delpeut, S.; Noyce, R.S.; Sisson, G.; Black, K.M.; Lin, L.T.; Bilimoria, D.; Plemper, R.K.; Prive, G.G.; Richardson, C.D. Mutations in the Fusion Protein of Measles Virus That Confer Resistance to the Membrane Fusion Inhibitors Carbobenzoxy-d-Phe-I-Phe-Gly and 4-Nitro-2-Phenylacetyl Amino-Benzamide. J. Virol. 2017, 91. [CrossRef]
- 12. Xie, Y.; Luo, X.; Li, Y.; Chen, L.; Ma, W.; Huang, J.; Cui, J.; Zhao, Y.; Xue, Y.; Zuo, Z.; et al. DeepNitro: Prediction of Protein Nitration and Nitrosylation Sites by Deep Learning. *Genom. Proteom. Bioinform.* **2018**, *16*, 294–306. [CrossRef] [PubMed]
- 13. Hasan, M.M.; Khatun, M.S.; Mollah, M.N.H.; Yong, C.; Dianjing, G. NTyroSite: Computational Identification of Protein Nitrotyrosine Sites Using Sequence Evolutionary Features. *Molecules* **2018**, 23, 1667. [CrossRef] [PubMed]
- 14. Xu, Y.; Wen, X.; Wen, L.S.; Wu, L.Y.; Deng, N.Y.; Chou, K.C. iNitro-Tyr: Prediction of nitrotyrosine sites in proteins with general pseudo amino acid composition. *PLoS ONE* **2014**, *9*, e105018. [CrossRef]
- 15. Liu, Z.; Cao, J.; Ma, Q.; Gao, X.; Ren, J.; Xue, Y. GPS-YNO2: Computational prediction of tyrosine nitration sites in proteins. *Mol. Biosyst.* **2011**, *7*, 1197–1204. [CrossRef]
- 16. Vacic, V.; Iakoucheva, L.M.; Radivojac, P. Two Sample Logo: A graphical representation of the differences between two sets of sequence alignments. *Bioinformatics* **2006**, 22, 1536–1537. [CrossRef]
- 17. Wei, L.; Hu, J.; Li, F.; Song, J.; Su, R.; Zou, Q. Comparative analysis and prediction of quorum-sensing peptides using feature representation learning and machine learning algorithms. *Brief. Bioinform.* **2018**. [CrossRef]
- 18. Hasan, M.M.; Alam, M.A.; Shoombuatong, W.; Kurata, H. IRC-Fuse: Improved and robust prediction of redox-sensitive cysteine by fusing of multiple feature representations. *J. Comput. Aided Mol.* **2021**. [CrossRef]

19. Hasan, M.M.; Rashid, M.M.; Khatun, M.S.; Kurata, H. Computational identification of microbial phosphorylation sites by the enhanced characteristics of sequence information. *Sci. Rep.* **2019**, *9*, 8258. [CrossRef]

- 20. Hasan, M.M.; Zhou, Y.; Lu, X.; Li, J.; Song, J.; Zhang, Z. Computational Identification of Protein Pupylation Sites by Using Profile-Based Composition of k-Spaced Amino Acid Pairs. *PLoS ONE* **2015**, *10*, e0129635. [CrossRef]
- 21. Chen, Z.; Zhou, Y.; Zhang, Z.; Song, J. Towards more accurate prediction of ubiquitination sites: A comprehensive review of current methods, tools and features. *Brief. Bioinform.* **2015**, *16*, 640–657. [CrossRef]
- 22. Fu, L.; Niu, B.; Zhu, Z.; Wu, S.; Li, W. CD-HIT: Accelerated for clustering the next-generation sequencing data. *Bioinformatics* **2012**, 28, 3150–3152. [CrossRef]
- 23. Hasan, M.M.; Schaduangrat, N.; Basith, S.; Lee, G.; Shoombuatong, W.; Manavalan, B. HLPpred-Fuse: Improved and robust prediction of hemolytic peptide and its activity by fusing multiple feature representation. *Bioinformatics* **2020**, *36*, 3350–3356. [CrossRef] [PubMed]
- 24. Khatun, M.S.; Hasan, M.M.; Kurata, H. PreAIP: Computational Prediction of Anti-inflammatory Peptides by Integrating Multiple Complementary Features. *Front. Genet.* **2019**, *10*, 129. [CrossRef] [PubMed]
- 25. Hasan, M.M.; Yang, S.; Zhou, Y.; Mollah, M.N. SuccinSite: A computational tool for the prediction of protein succinylation sites by exploiting the amino acid patterns and properties. *Mol. Biosyst.* **2016**, *12*, 786–795. [CrossRef] [PubMed]
- Kawashima, S.; Pokarowski, P.; Pokarowska, M.; Kolinski, A.; Katayama, T.; Kanehisa, M. AAindex: Amino acid index database, progress report 2008. Nucleic Acids Res. 2008, 36, D202–D205. [CrossRef]
- 27. Hasan, M.M.; Kurata, H. GPSuc: Global Prediction of Generic and Species-specific Succinylation Sites by aggregating multiple sequence features. *PLoS ONE* **2018**, *13*, e0200283. [CrossRef] [PubMed]
- Khatun, M.S.; Hasan, M.M.; Shoombuatong, W.; Kurata, H. ProIn-Fuse: Improved and robust prediction of proinflammatory peptides by fusing of multiple feature representations. J. Comput. Aided Mol. Des. 2020, 34, 1229–1236. [CrossRef]
- 29. Hasan, M.M.; Manavalan, B.; Shoombuatong, W.; Khatun, M.S.; Kurata, H. i6mA-Fuse: Improved and robust prediction of DNA 6 mA sites in the Rosaceae genome by fusing multiple feature representation. *Plant Mol. Biol.* **2020**, *103*, 225–234. [CrossRef]
- Hasan, M.M.; Manavalan, B.; Khatun, M.S.; Kurata, H. i4mC-ROSE, a bioinformatics tool for the identification of DNA N4-methylcytosine sites in the Rosaceae genome. *Int. J. Biol. Macromol.* 2020, 157, 752–758. [CrossRef] [PubMed]
- 31. Hasan, M.M.; Basith, S.; Khatun, M.S.; Lee, G.; Manavalan, B.; Kurata, H. Meta-i6mA: An interspecies predictor for identifying DNA N6-methyladenine sites of plant genomes by exploiting informative features in an integrative machine-learning framework. *Brief. Bioinform.* 2020. [CrossRef]
- Peng, C.; Wu, X.; Yuan, W.; Zhang, X.; Li, Y. MGRFE: Multilayer recursive feature elimination based on an embedded genetic algorithm for cancer classification. *IEEE/ACM Trans. Comput. Biol. Bioinform.* 2019. [CrossRef]
- 33. Wei, L.; He, W.; Malik, A.; Su, R.; Cui, L.; Manavalan, B. Computational prediction and interpretation of cell-specific replication origin sites from multiple eukaryotes by exploiting stacking framework. *Brief. Bioinform.* **2020**. [CrossRef] [PubMed]
- 34. Basith, S.; Manavalan, B.; Hwan Shin, T.; Lee, G. Machine intelligence in peptide therapeutics: A next-generation tool for rapid disease screening. *Med. Res. Rev.* **2020**, *40*, 1276–1314. [CrossRef]
- 35. Manavalan, B.; Basith, S.; Shin, T.H.; Wei, L.; Lee, G. Meta-4mCpred: A Sequence-Based Meta-Predictor for Accurate DNA 4mC Site Prediction Using Effective Feature Representation. *Mol. Ther. Nucleic Acids* **2019**, *16*, 733–744. [CrossRef] [PubMed]
- 36. Hasan, M.M.; Manavalan, B.; Khatun, M.S.; Kurata, H. Prediction of S-nitrosylation sites by integrating support vector machines and random forest. *Mol. Omics* **2019**, *15*, 451–458. [CrossRef] [PubMed]
- 37. Manavalan, B.; Basith, S.; Shin, T.H.; Wei, L.; Lee, G. mAHTPred: A sequence-based meta-predictor for improving the prediction of anti-hypertensive peptides using effective feature representation. *Bioinformatics* **2019**, *35*, 2757–2765. [CrossRef]
- 38. Basith, S.; Manavalan, B.; Shin, T.H.; Lee, G. SDM6A: A Web-Based Integrative Machine-Learning Framework for Predicting 6mA Sites in the Rice Genome. *Mol. Ther. Nucleic Acids* **2019**, *18*, 131–141. [CrossRef]
- 39. Tasmia, S.A.; Faisal, F.F.; Mosharaf, M.P.; Hasan, M.M.; Mollah, M.N.H. An Improved Computational Prediction Model for Lysine Succinylation Sites Mapping on Homo sapiens by Fusing Two Sequence Encoding Schemes with the Random Forest Classifier. *Curr. Genom.* 2021. [CrossRef]
- 40. Auliah, F.N.; Nilamyani, A.N.; Shoombuatong, W.; Alam, M.A.; Hasan, M.M.; Kurata, H. PUP-Fuse: Prediction of Protein Pupylation Sites by Integrating Multiple Sequence Representations. *Int. J. Mol. Sci.* **2021**, 22, 2120. [CrossRef]
- 41. Basith, S.; Manavalan, B.; Shin, T.H.; Lee, D.Y.; Lee, G. Evolution of Machine Learning Algorithms in the Prediction and Design of Anticancer Peptides. *Curr. Protein. Pept. Sci.* **2020**, *21*, 1242–1250. [CrossRef]
- 42. Khatun, S.; Hasan, M.; Kurata, H. Efficient computational model for identification of antitubercular peptides by integrating amino acid patterns and properties. *FEBS Lett.* **2019**, *593*, 3029–3039. [CrossRef] [PubMed]
- 43. Manavalan, B.; Shin, T.H.; Kim, M.O.; Lee, G. AIPpred: Sequence-Based Prediction of Anti-inflammatory Peptides Using Random Forest. *Front. Pharm.* **2018**, *9*, 276. [CrossRef]
- 44. Khatun, M.S.; Shoombuatong, W.; Hasan, M.M.; Kurata, H. Evolution of Sequence-based Bioinformatics Tools for Protein-protein Interaction Prediction. *Curr. Genom.* **2020**, *21*, 454–463. [CrossRef]
- 45. Charoenkwan, P.; Yana, J.; Schaduangrat, N.; Nantasenamat, C.; Hasan, M.M.; Shoombuatong, W. iBitter-SCM: Identification and characterization of bitter peptides using a scoring card method with propensity scores of dipeptides. *Genomics* **2020**, 112, 2813–2822. [CrossRef]

46. Charoenkwan, P.; Yana, J.; Nantasenamat, C.; Hasan, M.M.; Shoombuatong, W. iUmami-SCM: A Novel Sequence-Based Predictor for Prediction and Analysis of Umami Peptides Using a Scoring Card Method with Propensity Scores of Dipeptides. *J. Chem. Inf. Model.* 2020. [CrossRef]

- 47. Charoenkwan, P.; Nantasenamat, C.; Hasan, M.M.; Shoombuatong, W. iTTCA-Hybrid: Improved and robust identification of tumor T cell antigens by utilizing hybrid feature representation. *Anal. Biochem.* **2020**, *599*, 113747. [CrossRef] [PubMed]
- 48. Charoenkwan, P.; Nantasenamat, C.; Hasan, M.M.; Shoombuatong, W. Meta-iPVP: A sequence-based meta-predictor for improving the prediction of phage virion proteins using effective feature representation. *J. Comput. Aided Mol. Des.* **2020**, *34*, 1105–1116. [CrossRef]
- 49. Charoenkwan, P.; Kanthawong, S.; Nantasenamat, C.; Hasan, M.M.; Shoombuatong, W. iDPPIV-SCM: A Sequence-Based Predictor for Identifying and Analyzing Dipeptidyl Peptidase IV (DPP-IV) Inhibitory Peptides Using a Scoring Card Method. *J. Proteome Res.* 2020, 19, 4125–4136. [CrossRef] [PubMed]
- 50. Charoenkwan, P.; Kanthawong, S.; Nantasenamat, C.; Hasan, M.M.; Shoombuatong, W. iAMY-SCM: Improved prediction and analysis of amyloid proteins using a scoring card method with propensity scores of dipeptides. *Genomics* **2020**. [CrossRef]
- 51. Ning, Q.; Ma, Z.; Zhao, X.; Yin, M. SSKM_Succ: A novel succinylation sites prediction method incorprating K-means clustering with a new semi-supervised learning algorithm. *IEEE/ACM Trans. Comput. Biol. Bioinform.* **2020**. [CrossRef]
- 52. Ning, Q.; Yu, M.; Ji, J.; Ma, Z.; Zhao, X. Analysis and prediction of human acetylation using a cascade classifier based on support vector machine. *BMC Bioinform.* **2019**, *20*, 346. [CrossRef]
- 53. Hasan, M.M.; Shoombuatong, W.; Kurata, H.; Manavalan, B. Critical evaluation of web-based DNA N6-methyladenine site prediction tools. *Brief. Funct. Genom.* **2021**. [CrossRef]
- 54. Hasan, M.M.; Khatun, M.S.; Kurata, H. iLBE for Computational Identification of Linear B-cell Epitopes by Integrating Sequence and Evolutionary Features. *Genom. Proteom. Bioinform.* **2020**. [CrossRef] [PubMed]
- 55. Hasan, M.M.; Khatun, M.S.; Kurata, H. Large-Scale Assessment of Bioinformatics Tools for Lysine Succinylation Sites. *Cells* **2019**, 8, 95. [CrossRef]
- 56. Charoenkwan, P.; Chiangjong, W.; Lee, V.S.; Nantasenamat, C.; Hasan, M.M.; Shoombuatong, W. Improved prediction and characterization of anticancer activities of peptides using a novel flexible scoring card method. *Sci. Rep.* **2021**, *11*, 3017. [CrossRef] [PubMed]
- 57. Wei, L.; Su, R.; Luan, S.; Liao, Z.; Manavalan, B.; Zou, Q.; Shi, X. Iterative feature representations improve N4-methylcytosine site prediction. *Bioinformatics* **2019**, *35*, 4930–4937. [CrossRef] [PubMed]
- 58. Charoenkwan, P.; Nantasenama, C.; Hasan, M.M.; Manavalan, B.; Shoombuatong, W. BERT4Bitter: A bidirectional encoder representations from transformers (BERT)-based model for improving the prediction of bitter peptides in Bioinformatics. *Bioinformatics* **2021**. [CrossRef]
- 59. Manavalan, B.; Hasan, M.M.; Basith, S.; Gosu, V.; Shin, T.H.; Lee, G. Empirical Comparison and Analysis of Web-Based DNA N (4)-Methylcytosine Site Prediction Tools. *Mol. Ther. Nucleic Acids* **2020**, 22, 406–420. [CrossRef] [PubMed]