Deep learning application detecting SARS-CoV-2 key enzymes inhibitors

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Abstract
The fast spread of the COVID-19 over the world pressured scientists to find its cures. Especially, with the disastrous results, it engendered from human life losses to long-term impacts on infected people’s health and the huge financial losses. In addition to the massive efforts made by researchers and medicals on finding safe, smart, fast, and efficient methods to accurately make an early diagnosis of the COVID-19. Some researchers focused on finding drugs to treat the disease and its symptoms, others worked on creating effective vaccines, while several concentrated on finding inhibitors for the key enzymes of the virus, to reduce its spreading and reproduction inside the human body. These enzymes’ inhibitors are usually found in aliments, plants, fungi, or even in some drugs. Since these inhibitors slow and halt the replication of the virus in the human body, they can help fight it at an early stage saving the patient from death risk. Moreover, if the human body’s immune system gets rid of the virus at the early stage it can be spared from the disastrous sequels it may leave inside the patient’s body. Our research aims to find aliments and plants that are rich in these inhibitors. In this paper, we developed a deep learning application that is trained with various aliments, plants, and drugs to detect if a component contains SARS-CoV-2 key inhibitor(s) intending to help them find more sources containing these inhibitors. The application is trained to identify various sources rich in thirteen coronavirus-2 key inhibitors. The sources are currently just aliments, plants, and seeds and the identification is done by their names.

Keywords Key enzymes inhibitors · COVID-19 · Deep learning · Plants · Aliments · Identification

1 Introduction
In recent years, the interest in artificial intelligence (AI) has been growing constantly. The need to have smart applications has motivated researchers to develop machine learning-based and then deep learning-based algorithms and programs. They were inspired by the human brain’s amazing abilities to learn. Therefore, they aimed to mimic its learning, thinking, and decision-making abilities through these algorithms to add the smartness trait to

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machines. As a result, various fields bloomed after supporting it with AI such as computer vision [1] where the computers are trained to identify objects from images and videos. Various deep learning (DL) applications appeared, like smart healthcare [2], gaming [3], and social media where the AI helps make an accurate prediction about user preferences [4]. Additionally, its application to health systems where a computer can identify the presence of diseases from MRI and CT images such as breast cancer, COVID-19 [5], etc. The use of deep learning to serve medical purposes is a prosperous and worth investment due to the benefits it brings to humanity. It can boost the accuracy of diagnostics to make it more efficient and fasten the detection process of diseases. It is widely known that early detection and intervention of medical staff increases the chances of survival, especially in critical cases. Profoundly training the computers about medical knowledge can highly assist the doctors. However, this requires transferring their knowledge and expertise in an accurate, organized, and systematic way to the computers. The spread of COVID-19 and the disastrous losses it engendered, with over 252,780,044 cases of infection and over 5,098,342 deaths worldwide reported at the time of writing based on the statistics provided by WHO [6]. Not to mention the drastic negative impact it left on the economy and the financial losses recorded worldwide. Moreover, the influence it left on our daily lives, like the obligatory wear of masks and the need to carry sanitizers everywhere, the travel ban and restrictions as well as the need for health passports and vaccination proofs, etc. Furthermore, to the moment there is no accurate prediction about when will the health safety prevail or when will this virus completely diminish, especially, with new strikes recorded from time to time and the appearance of new variants. This situation pushed researchers to find solutions to it, by finding remediating drugs, and vaccinations or by helping its early detection through the use of DL. Researchers in [7] proposed a framework that embeds a mass surveillance system and multi-model diagnostic system with the B5G (beyond 5G) and deep learning techniques as well as edge and cloud computing to efficiently detect and diagnose COVID-19 and infection like diseases. The need for a prompt response during the pandemic and real-time awareness about the infection spread as well as the exchange of knowledge in a secure and privacy-preserving manner led to the use of multiple technologies such as The Internet of Vehicles (IoV) [8], Unmanned Aerial Vehicles (UAVs) or drones, blockchains, deep learning and computer vision, edge computing, cloud computing, big data, etc. All in all, the need to have fast internet and secure confidential communications were further emphasized as essential needs [9].

Furthermore, the necessity of deploying the smart healthcare systems powered by the internet of medical things and computer-aided diagnostics has once more been highlighted during this pandemic to ensure the safety of healthcare workers, the efficiency of treatment, fast response, and limitation of virus spread [10].

Researchers of [11] used the AI techniques to identify SARS-CoV-2 main protease inhibitors from FDA-approved drugs where they were able to detect 13 positive drugs amongst 20 potentially tested drugs, 6 of which were identified for the first time in their study. Authors of [12] used deep learning pretrained drug-target interaction model to find potential drugs among the commercially available that could act on SARS-CoV-2 viral proteins and inhibit them. In their results, they found multiple potential inhibitor drugs and they highly recommended them to be clinically tested for further confirmation. Similarly, authors of [13] used molecular deep learning training to identify potential inhibitors SARS-CoV-2 Mpro inhibitors. The authors of [14] used generative deep learning models inspired by generative chemistry to design new drug-like inhibitors of 2019-nCoV. Multiple works were published aiming to use DL to find and repurpose drugs from the FDA-approved ones that are commercially available to treat COVID-19 such as in [15]. Since deep learning relies on an accurate dataset, researchers have been focusing on collecting and providing related molecular databases such as CoronaDB-AI [16].

After reading the efforts made by scientists from multiple perspectives in order to elaborate an effective healthcare system against COVID-19. We draw Fig. 1 to summarize all their efforts. The first thing we noticed is the persistent emphasis on the need for a stable, fast, and large bandwidth communication medium for the exchange of medical data represented by the use of 4G, 5G, and B5G, etc. The second high requirement is the secure lightweight communications, the control access of data, and the privacy of patients, leading to the support of edge computing usage. Thirdly, the massive amount of exchanged data and the necessity to share information between medical staff worldwide recommended the use of cloud computing, blockchains, and big data. Fourthly, the need for fast and accurate diagnosis and the processing of massive data to look for remediating drugs or vaccines pushed for the use of deep learning and AI advanced techniques. Lastly, to prevent the spread of the virus in an effective safe way, IoT technology backed by (infrared) cameras, drones, and sensors is recommended to be used. To recap, the made efforts are directed toward four areas which are:

1. Accurately and efficiently diagnosing COVID-19,
2. Finding effective COVID-19 drugs and vaccines,
3. Preventing the COVID-19 from spreading through massive surveillance systems,
(4) Ensure public safety and prevent viruses from crossing borders (health passports).

Motivated by the same reasons, we also aimed in this paper to use DL to discover COVID-19 key enzyme inhibitors. Unlike the mentioned works, our proposed application is original and novel. It aims to use DL techniques to identify the inhibitors from natural resources like aliments, plants, herbs, etc. They are directly edible and they can strengthen the immune system during the COVID-19 infection and prevent virus replications. Encouraging the importance of a healthy diet, and reckoning on food to become a remedy. Therefore, we trained the computer to know which aliments/herbs can be eaten to reduce the negative impacts of the COVID-19 virus on the body of the infected and to reduce its reproductivity rate helping the body to defend itself. Our app was trained based on works published by scientists on key enzyme inhibitors of the COVID-19 virus. We first selected these inhibitors and then researched natural aliments, plants, herbs, and seeds rich with them. Lastly, we fed these data to our developed DL web app. This paper will be organized as follows: Sect. 2 explains the deep learning concept and how it works. Section 3 illustrates the COVID-19 treatment mechanisms. Section 4 describes the proposed application, its role, the dataset collection, and the preparation of training. Section 5 illustrates the testing dataset, phases, and results. Section 6 concludes the paper and highlights our future perspectives.

2 Technical background

Deep learning is a subfield of machine learning that models high-level data abstractions using multiple neurons’ layers consisting of complex structures or non-linear transformations [17]. Noting that in DL these layers are not human-designed rather they are learned from data through general-purpose learning procedures [18]. In DL the high-level features are extracted from low-level features. The DL uses neural networks with multiple hidden layers where the input is processed through the layers to compute the output. There exist various DL architectures each is suitable for a certain type of training datasets such as convolutional neural network (CNN) for images [19], recurrent neural network (RNN) for sequential data, and its extension for variable-length data the long short-term memory (LSTM) [17, 20, 21]. In our app, we used LSTM for the training because the length of our data is variable. The LSTM can learn long-term dependencies and remember them. It is composed of layers and memory blocks noted as cells (see Fig. 2) [22]. The data can be added or removed through a sigmoid gate. Noting that a gate is a layer or a sequence of matrix operations containing various weights.
The initial step in composing the LSTM network is eliminating and excluding useless data through the sigmoid or forget gate. This process is repeated at each inner layer to eliminate unnecessary data and store the important ones in the cells until the final output is calculated [23]. In our application, we used an LSTM network trained using our textual data. The data is a set of aliment/plants/herbs/seeds where the COVID-19 key enzyme inhibitors are found or absent. Our data was stored as json file seeded to our web app that uses LSTM neural networks for the training. The web application is developed using HTML, CSS, and JavaScript. For the deep learning functions and networks, we used brainjs [24] which is a GPU accelerated neural network implementation in JavaScript for use in browsers and via nodejs [25].

3 COVID-19 treatments review

We have screened different databases about the number of published works until the time of writing of this paper (26th, January 2022) on COVID-19 vaccines, we have found more than 342 000 published papers in google scholar, about 19,793 papers in the science direct database [26] and 2875 paper in nature database [27]. Actually, we have different variants of COVID-19, the most spread is Omicron followed by Delta, we have recorded 106 published papers on nature database just for omicron, this number is higher in other databases. This proves the international serious interest in curing this disease. For the natural SARS-CoV-2 inhibitors obtained from plants; we have discovered more than 50, 400 papers on google scholar, nearly 9200 papers on science-direct, with 529 papers on the nature database. We focused on selecting inhibitors abundant in our daily food intakes to facilitate balancing our diet during infection with this virus and boosting our immune system. We have collected these data according to the high potent inhibitors published in previous works with in vitro; in vivo and in silico studies.

Scientists around the world have used common drugs for other viruses like SARS-CoV-1, MERS-CoV, and HIV as well as antimalarial drugs were inspected for activity against SARS-CoV-2 [28]. Including Remdesivir (designed for Ebola virus), Lopinavir / Ritonavir (designed for HIV) has inhibitory activity on SARS-CoV-2 Main protease (Mpro) [29], chloroquine and hydroxychloroquine (designed for malaria), and Tocilizumab (designed for rheumatoid arthritis), in addition, other potential drugs from existing antiviral agents have also been proposed [30, 31]. PAXLOVID™ (PF-07321332) is the last developed drug (one pill against COVID-19) by Pfizer which is an inhibitor of the SARS-CoV-2 main protease (3CLpro); this latter binds covalently to the catalytic amino acid Cys145. In November 2021, Pfizer reported positive Phase 2/3 results, with an 89% reduction in hospitalizations if administered within three days of symptom onset [32, 33, 34].

China was the first country that used Medicinal plants for COVID-19 treatment, more than 23,600 published papers on the most used medicinal plants, especially in China; they have used a well-known mixture in their folk medicine to treat COVID-19 patients [35] as they describe it in the published reviews [36, 37]. According to these reviews, the top ten Chinese herb medicines used were Maxing Shigan Tang, Lianhua Qingwen granule/capsule, Xuebijing injection, Dayuanyn, Shufeng Jiedu capsule, Qingfei Paidu Tang, Xiaochaihu Tang, Ganlu Xiaodu Dan, Liujuanzi Tang and Toujie Quwen granule [36].

Numerous studies identified different proteins of the SARS-CoV-2 as keys to its development and replication; The 3C-Like proteinase (3CLpro) of the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is the first drug target due to its role in the replication of the virus [38]; Following studies have identified different drug targets as these enzymes or known as non-structural proteins NSP, which include NSP1-NSP16. NSP3 (papain-like protease), NSP5 (3-chymotrypsin-like protease “3-CLprotease”), NSP12 (RdRp: RNA-dependent RNA polymerase), NSP13 (helicase), NSP14 (N7-methyltransferase), and NSP16 (2'-O-methyl transferase) are important viral enzymes while NSP7-NSP10 are regulatory proteins [39], with more than 1600 3D structures published in PDB [40]. Meta-analysis was achieved to determine the different published PDB structures of the non-structural proteins; Table 1 groups the major potent natural compounds found until now as potential inhibitors of the 3CLpro through clinical trials describing their possible sources. The results of these meta-analyses were set as the necessary data for training the DL application.

![Fig. 2 LSTM Cell and its operations](image-url)
Table 1 Natural chemical compounds as SARS-CoV-2 3CL-pro potential inhibitors

| Molecule Name | References | Some natural sources |
|---------------|------------|----------------------|
| Folic Acid-vitamin B9 | [41, 42, 43, 44, 45] | Turnip greens, Spinach, Romaine, Lettuce, Asparagus, Broccoli, Brussels sprouts, Peanuts, Sunflower seeds, Strawberry, Bananas, Whole grains, Liver, Salmon, Eggs |
| Hispidin | [43, 46, 47, 48, 49] | Inonotus hispidus, I. obliquus, I. xeranticus, Phellinus linteus, P. ignarius, P. baumii, P. harmala, P. sensulato, Australian fungus Cortinarius, Neoantopanus nambii, horsetail (Equisetum arvense), piper (Piper methysticum) |
| Niacin-vitamin B3 | [50, 51, 52, 53, 54, 55] | Chicken Breast, Tuna, Salmon, Anchovies, Ground Beef, Peanuts, Avocado, Brown Rice, Whole Wheat, Mushrooms, Green Peas, Potatoes, Grilled sesame seed, Almond |
| Curcumin | [56] | Turmeric, Curcuma amada |
| Baicalin and Baicalein | [57, 58] | Scutellaria baicalensis, S. lateriflora, S. galericulata, Oroxyllum indicum |
| Aloe-emodin | [59, 60, 61] | Cassia occidentalis, Rheum palmatum L, Aloe vera, Polygonum multiflorum Thunb, Rhamnus frangula bark, Rhamnus purshiana, Cassia angustifolia leaves, Rheum rhabonticum rhizome |
| Amentoflavone | [62] [63] | Origanum majorana L, Calophyllum rutilare leaves, C. spongiontianum barks, leaves, C. venulosum leaves, Campylospernum callanth fruits leaves, Campylospernum manii leaves |
| Hesperetin | [64, 65] | Origanum majorana L, Calophyllum rutilare leaves, Calophyllum spongiontianum barks, leaves; Campylospernum callanth fruits, Canarium pinela, Casearia clarki |
| Luteolin | [66, 67] | Origanum majorana L, Parsley (Petroselinum crispum), Artichoke (Cynara scolymus L, Basil (Ocimum sanctum L), Celery (Apium graveolens), Cabbage (Brassica oleracea), Broccoli, Rosemary, Onions…etc |
| Quercetin | [68, 69, 70] | Capers, Onions, Elderberries, Kale, Okra, Apple Peels, Aronia Berries, Cranberries, Asparagus, Goji Berries, Lovage leaves, Radish leaves |
| Psoralidin | [71, 72] | Ficus carica, Anise seeds, Caraway seeds, Carrots, Celeriac, Chervil, Cilantro, Coriander seeds, Cumin seeds, Dill, Fenel seeds, Grapefruit, Lemons, Limes, Lovage, Mustard seeds |
| Theaflavin-3’-gallate | [73] | Black tea, Camellia sinensis (L.) Kuntze |
| Tannic acid | [74, 75] | Black tea, Grapes, Green tea, Persimmons, Black beans, Red beans, Apricots, Cherries, Peaches, Dates, English walnuts, Black walnuts, Cashews, Blueberries, Blackberries, Strawberries, Chocolate, Coffee, Cinnamon, Cumin |

4 Proposed application

The implementation of our applications followed multiple steps which are illustrated in Figs. 3 and 4. The most tedious and time-consuming task was searching for COVID-19 inhibitors, with the huge number of papers published about COVID-19 (See Sect. 3), filtering them and selecting those about inhibitors then looking for their natural sources like aliments, plants, seeds, and herbs was effort-demanding. After collecting all this data (Fig. 3. Steps 1 and 2), we had to think of how to represent it and seed it to the neural network. The training dataset is represented as a json file composed of multiple entries; each entry has two fields. The 1st represents the natural source, the 2nd represents whether or not this natural source contains inhibitors (Fig. 3. Step 3). The next step was to select the appropriate learning model to use which is LSTM (Fig. 3. Step 4).

Steps 5 and 6 in Fig. 3 were to implement the backend which contained the brainjs LSTM training network code on the training data composed of 175 aliment/plant containing 13 COVID-19 enzymes inhibitors, the LSTM parameters are resumed in Table 2. Noting that the learning
rate defines the training speed which is slow if close to 0, and quick if close to 1. We checked these parameters from brainjs documentation [24]. The training was done on a quad-core i7 with NVIDIA GeForce GT 750 M graphic card. Upon confirming that the backend works, we started building the front-end, we used HTML, CSS for the interface and its styling, and JavaScript to implement its functioning (Fig. 3. Step 7). The front-end is simple, in the current version, the user (or researcher) needs to provide the name of a natural source and the application will check whether it contains one of COVID-19 key enzymes’ inhibitors that it was trained to recognize. To be able to run the functionalities of the backend from the browser, we used browserify tool [76] like in [77] (Fig. 3. Step 8). Noting that as illustrated in Fig. 4, we refined the implementation and the training parameters multiple times. The refining was done after multiple rounds of testing and result analysis. It aimed to enhance the accuracy of the identification of the app before finally providing it to the users. The testing details and parameters and their impact on the results are illustrated in the next section.

5 Testing and analysis

To test our application, we tried various sources and aliments to check if they inhibit or not the COVID-19 enzymes. The sizes of training and testing datasets are illustrated in Table 2. We tested the application multiple times by changing the training parameters (number of iterations and errorThresh) to get more refined and accurate results. The accuracy level of our application against the testing dataset in each test is recapped in Table 3. We choose to evaluate our app by calculating the accuracy level using Eq. 1 and the error rate using Eq. 2. The average obtained accuracy level varied between 54% in Test 1 to 74% in Test 4. Consecutively, the error rate varied between 36 to 26% respectively. Figure 5 depicts the average obtained results in each test. Overall, it can be observed that the refining process which required tuning the training parameters helped in improving the training accuracy and lowering the error rate.

Table 2 LSTM parameters

| Parameter          | Test 1 | Test 2 | Test 3 | Test 4 |
|--------------------|--------|--------|--------|--------|
| Iterations         | 2000   | 2000   | 5000   | 7000   |
| ErrorThresh        | 5*10⁻² | 10⁻³   | 10⁻⁴   | 10⁻⁴   |
| Learning Rate      | 0.3    | 0.3    | 0.3    | 0.3    |
| Momentum           | 0.1    | 0.1    | 0.1    | 0.1    |
| Callback period    | 10     | 10     | 10     | 10     |
| Training data set  | JSON file of string | | | |
| Training data size | 175    |        |        |        |
| Testing data size  | 814    |        |        |        |

Table 3 Testing results recap

|                     | Test 1 | Test 2 | Test 3 | Test 4 |
|---------------------|--------|--------|--------|--------|
| Accuracy level      | 54%    | 55%    | 72%    | 74%    |
| Error rate          | 46%    | 45%    | 28%    | 26%    |
| False-positive      | 73%    | 75%    | 49%    | 42%    |
| False-negative      | 27%    | 25%    | 51%    | 58%    |
While evaluating the training accuracy and error rates, we decided to further investigate the types of errors given by our trained application. We classified the errors into the false positive and false negative. The false-positive represents the mistake our app gave when it classified an element containing COVID-19 inhibitors as a non-inhibitor-container. Its rate is calculated using Eq. 3. The false negative is the mistake our app made when it classified an element that does not contain COVID-19 inhibitors as an inhibitor-container. Its rate is calculated using Eq. 4.

Table 3 illustrates that as we were refining the results, we were able to reduce the false-positive errors from 73 to 42% in tests 1 and 4 respectively. On the contrary, the false-negative rate increased from 27% in Test 1 to 58% in Test 4. Noting that in Test 2, we obtained the lowest false-negative rate (25%), although the false-positive rate was the highest (75%). Figures 6, 7, 8 and 9 detail the identification accuracy of each inhibitor as well as the error rate (false positive and negative) in each test. The error rate was due to the nature of the dataset which requires further processing. Currently, some of the aliments in the dataset are given by their scientific names, some have simple names while others contain acronyms or special characters and are composed of multiple words. The variable length and the non-uniform format are what are causing the moderate accuracy and engendering errors.

\[
\text{Accuracy} = \frac{\text{Correct identification}}{\text{Size of test data set}} \tag{1}
\]

\[
\text{Error Rate} = \frac{\text{Incorrect identification}}{\text{Size of test data set}} \tag{2}
\]

\[
\text{False Positive} = \frac{\text{NRC}}{\text{Incorrect identification}} \tag{3}
\]

where, \(\text{NRC} : \text{Number of rejected correct values}\)

\[
\text{False Negative} = \frac{\text{NAW}}{\text{Incorrect identification}} \tag{4}
\]

where, \(\text{NAW} : \text{Number of accepted wrong values}\)

6 Conclusion and future perspectives

In this paper, we explained the approach we followed to create a DL-based web application to check the natural sources’ continency of COVID-19 key Enzymes’ inhibitors. As we have explained earlier, the current application replies only with yes or no based on what it was trained to know. This application aims to draw the attention of the scientific community world widely to this approach to exchange more data that could help train the app. It could serve also as a large database for scientists working on COVID-19 enzymes inhibition field to check for more potential aliments rich with them before the lab tests. Moreover, the same plants and aliments have different names depending on the regions and countries, some plants although different may have the same key effective compounds. Therefore, we aim to train the app to know the potential sources for COVID-19 enzyme inhibitors by their composition not only their names. Succeeding in this purpose depends on the availability of data regarding the sources rich with these inhibitors and the key compositions of these sources. Currently, we are working on gathering, filtering, and sorting this dataset for the future release of the second version of this application.
Fig. 6 Detailed Results of Test 1

Fig. 7 Detailed Results of Test 2
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Data availability Data are available upon request to the first author (L. Benarous; email: l.benarous@lagh-univ.dz).

Declarations

Conflict of interest There is no conflict of interest.

Informed consent None.

References

1. Voulodimos, A., Doulamis, N., Doulamis, A., Protopapadakis, E.: Deep learning for computer vision: A brief review. Comput. Intell. Neurosci 2018, 1–13 (2018)
2. Muhammad, G., Hossain, M.S.: COVID-19 and Non-COVID-19 classification using multi-layers fusion from lung ultrasound images. Information Fusion 72, 80–88 (2021)

3. Justesen, N., Bontrager, P., Togelius, J., Risi, S.: Deep learning for video game playing. IEEE Transact. Games 12(1), 1–20 (2019)

4. Wang, J.H., Wu, Y.T., Wang, L.: Predicting implicit user preferences with multimodal feature fusion for similar user recommendation in social media. Appl. Sci. 11(3), 1064 (2021)

5. Muhammad, G., Hossain, M.S.: A deep-learning-based edge-centric COVID-19-like pandemic screening and diagnosis system within a BSG framework using blockchain. IEEE Network 35(2), 74–81 (2021)

6. COVID-19 Coronavirus Pandemic, worldometers. [Online]. Available: https://www.worldometers.info/coronavirus/. [Accessed 12 11 2021]

7. Hossain, M.S., Muhammad, G., Guizani, N.: Explainable AI and mass surveillance system-based healthcare framework to combat COVID-19 like pandemics. IEEE Network 34(4), 126–132 (2020)

8. Chien, W.C., Lai, C.F., Hossain, M.S., Muhammad, G.: Heterogeneous space and terrestrial integrated networks for IoT: Architecture and challenges. IEEE Network 33(1), 15–21 (2019)

9. Masud, M., Gaba, G.S., Choudhary, K., et al.: Lightweight and anonymity-preserving user authentication scheme for IoT-based healthcare. IEEE Internet Things J. 9(4), 2649–2656 (2022)

10. Muhammad, G., Alshehri, F., Alsuheih, S., Alshehri, F., Alshehri, F.: Predicting implicit user preferences with multimodal feature fusion for similar user recommendation in social media. Appl. Sci. 11(3), 1064 (2021)

11. Komatsu, H., Tanaka, T., Ye, Z., Ikeda, K., Matsuzaki, T., Yasugi, M., Hosoda, M.: Identification of SARS-CoV-2 main protease inhibitors from FDA-approved drugs by artificial intelligence-supported activity prediction system. J. Biomol. Struct. Dynamics 2019, 1–9 (2021)

12. Beck, B.R., Shin, B., Choi, Y., Park, S., Kang, K.: Predicting commercially available antiviral drugs that may act on the novel coronavirus (SARS-CoV-2) through a drug-target interaction deep learning model. Comput. Struct. Biotechnol. J. 18, 784–790 (2020)

13. Shaker, N., Abou-Zleikha, M., AlAmri, M., & Mehellou, Y. (2020) A Generative deep learning approach for the discovery of SARS-CoV2 protease inhibitors. https://doi.org/10.26434/chemrxiv.11829102.v1

14. Palavonkorn, A., Alekzinsky, V., Zhebrak, A., Zagribelnyy, B., Terentiev, V., Bezrukov, D.S., Ivanenko, Y.: Potential 2019-nCoV 3CL(pro) protease inhibitors designed using generative deep learning approaches. ChemRxiv. Cambridge: Cambridge Open Engage (2020). https://doi.org/10.26434/chemrxiv.11829102.v1

15. Abdel-Basset, M., Hawash, H., Elhoseny, M., Chakrabortty, R.K., Ryan, M.: DeepH-DTA: deep learning for predicting drug-target interactions: a case study of COVID-19 drug repurposing. IEEE Access 8, 170433–170451 (2020)

16. Keshavarzi Arshadi, A., Webb, J., Salem, M., Cruz, E., Calad-Thomson, S., Ghadiani, N., Yuan, J.S.: Artificial intelligence for COVID-19 drug discovery and vaccine development. Frontiers in Artificial Intelligence 3, 65 (2020)

17. Hao, X., Zhang, G., Ma, S.: Deep learning. Int. J. Semantic Comput 10(03), 417–439 (2016)

18. LeCun, Y., Bengio, Y., Hinton, G.: Deep learning. Nature 521(7553), 436–444 (2015)

19. Muhammad, G., Hossain, M.S.: Emotion recognition for cognitive edge computing using deep learning. IEEE Internet Things J. 8(23), 16894–16901 (2021)

20. Amin, S.U., et al.: Attention-inception and long-短期-term memory-based electroencephalography classification for motor imagery tasks in rehabilitation. IEEE Trans. Industr. Inf. (2022). https://doi.org/10.1109/TII.2021.3132340

21. DiPietro, R., & Hager, G.D. (2020) Deep learning: RNNs and LSTM. In Handbook of medical image computing and computer assisted intervention (pp. 503–519) Academic Press

22. M. Phi, "Illustrated Guide to LSTM’s and GRU’s: A step by step explanation," 24 09 2018. [Online]. Available: https://towardsdatascience.com/illustrated-guide-to-lstms-and-gru-s-a-step-by-step-explanation-44e9eb85ff21. [Accessed 24 12 2021]

23. Muhammad, G., Hossain, M.S., Kumar, N.: EGG-based pathology detection for home health monitoring. IEEE J. Sel. Areas Commun. 39(2), 603 (2021)

24. "Brain.js," [Online]. Available: https://brain.js.org/#/. [Accessed 15 11 2021]

25. "Node.js," [Online]. Available: https://nodejs.org/en/. [Accessed 15 11 2021]

26. "Science Direct," [Online]. Available: https://www.sciencedirect.com/. [Accessed 28 01 2022]

27. "Nature," [Online]. Available: https://www.nature.com/. [Accessed 28 01 2022]

28. Abouelela, M.E., et al.: Identification of potential SARS-CoV-2 main protease and spike protein inhibitors from the genus aloe: an in silico study for drug development. Molecules 26(6), 1767 (2021). https://doi.org/10.3390/molecules26061767

29. Yadav, M., Dhagat, S., Esvari, J.S.: Emerging strategies on in silico drug development against COVID-19: challenges and opportunities. Eur. J. Pharm. Sci. 155, 105522 (2020). https://doi.org/10.1016/j.ejps.2020.105522

30. Gimeno, A., et al.: Prediction of Novel Inhibitors of the Main Protease (M-pro) of SARS-CoV-2 through Consensus Docking and Drug Reposition. Int. J. Mol. Sci. 21(11), 3793 (2020). https://doi.org/10.3390/ijms21113793

31. Ibrahim, B., Önen, E.: Coronavirus disease 2019 (COVID-19): A literature review. Gevher Nesibe Journal IESDR 6(12), 56–72 (2021). https://doi.org/10.46648/jen.200

32. Kneller, D.W., Li, H., Galanie, S., Phillips, G., Labbe, A., Weiss, K.L., Zhang, Q., Arnould, M.A., Coates, L., Louis, J.M., Bonnesen, P.V., Kovalyevsky, A. Desing of hybrid covalent inhibitors of SARS-CoV-2 main protease containing a nitrile warhead, To be published

33. Pfizer begins dosing in Phase II/III trial of antiviral drug for COVID-19. » [archive], Clinical Trials Arena, 2 september 2021

34. Pfizer. Novel COVID-19 Oral Antiviral Treatment Candidate Reduced Risk Of Hospitalization Or Death By 89% In Interim Analysis Of Phase 2/3 EPIC-HR Study [archive], 5 novembre 2021

35. Li, X.Y., Teh, B.P., Tan, T.Y.C.: Medicinal Plants in COVID-19: Potential and Limitations. Fronti. Pharmacol. (2021).https://doi.org/10.3389/fphar.2021.611408

36. Liang, S.-B., et al.: Chinese herbal medicine used with or without conventional western therapy for COVID-19: An evidence review of clinical studies. Fronti. Pharmacol. (2021).https://doi.org/10.3389/fphar.2020.583450

37. Benarba, B., Pandiella, A.: Medicinal plants as sources of active molecules against COVID-19. Fronti. Pharmacol. (2020). https://doi.org/10.3389/fphar.2020.01189

38. Keta, S., Bhujbal, S.P., Cho, S.J.: Rational approach toward COVID-19 main protease inhibitors via molecular docking, molecular dynamics simulation and free energy calculation. Scientific Rep. (2020).https://doi.org/10.1038/s41598-020-74468-0

39. Itfikhar, H., Ali, H.N., Farooq, S., Naveed, H., Shahzad-ul-Husn, S.: Identification of potential inhibitors of three key enzymes of SARS-CoV2 using computational approach. Comput. Biol. Med. 122, 103848 (2020).https://doi.org/10.1016/j.compbiomed.2020.103848
40. Protein Data Bank [Online]. Available: https://www.rcsb.org/. [Accessed 17 11 2021]
41. Acosta-Elias, J., Espinosa-Tanguma, R.: The folate concentration and/or folic acid metabolites in plasma as factor for COVID-19 infection. Front. Pharmacol. (2020). https://doi.org/10.3389/fphar.2020.01062
42. Kumar, V., Kancharla, S., Jena, M.K.: In silico virtual screening-based study of Nutracuticals predicts the therapeutic potentials of folic acid and its derivatives against COVID-19. VirusDisease 32(1), 29–37 (2021)
43. Serseg, T., Benarous, K., Youfi, M.: Hspidin and lepidine E: Two natural compounds and folic acid as potential inhibitors of 2019-novel Coronavirus main protease (2019- nCoVMP). molecular docking and SAR study. Curr. Comput. Aided Drug Des. 17(3), 469–477 (2021)
44. Zapata, M.E., Rovirosa, A., Carmuego, E.: Iron and folic acid: natural, enriched, fortified, and supplements. Analysis of food sources in the Autonomous City of Buenos Aires. Arch. Argent. Pediatr. 118(3), 160–165 (2020)
45. Suslowati, A., Maryati, Y.: and Asyipanto, “Characteristic on infants biscuit for complementary feeding through fortificant paste of natural folic acid as smart food.” IOP Conf. Ser. Earth Environ. Sci. 484, 012131 (2020)
46. Galanakis, C.M., Aldawoud, T.M.S., Rizou, M., Rowan, N.J., Ibrahim, S.A.: Food ingredients and active compounds against the Coronavirus disease (COVID-19) pandemic: A comprehensive review”. Foods 9(11), 1701 (2020)
47. Khanna, K., et al.: Herbal immune-boosters: Substantial warriors of pandemic Covid-19 battle. Phytotherapy 85(135361), 135361 (2021)
48. He, P., Zhang, Y., Li, N.: The phytochemistry and pharmacology of medicinal fungi of the genus Phellinus: a review. Food Funct. 12(5), 1856–1881 (2021)
49. Palkina, K.A., Ipatova, D.A., Shakhova, E.S., Balakireva, A.V., Markina, N.M.: Therapeutic potential of hispidin-fungal and plant polyketide. J. Fungi (Basel) 7(5), 323 (2021)
50. Lotti, M., Hamblin, M.R., Rezaei, N.: COVID-19: Transmission, prevention, and potential therapeutic opportunities. Clin. Chim. Acta 508, 254–266 (2020)
51. Dr, J., Roli and P. Dr Archana,: COVID-19: A Cure and preventive options. J. Immunol. Immunother, Int. 2020. https://doi.org/10.23937/2378-3672/1410052
52. Mengist, H.M., Dilnessa, T., Jin, T.: Structural basis of potential inhibitors targeting SARS-CoV-2 main protease. Front. Chem. 9, 622898 (2021)
53. Chang, X., Mowat, D.N., Mallard, B.A.: Supplemental chromium and niacin for stressed feeder calves. Can. J. Anim. Sci. 75(3), 351–358 (1995)
54. Peikov, P., Obreshkova, D., Draganov, G.: Food supplements containing L-tryptophan and 5-hydroxytryptophan as precursors of serotonin, melatonin and niacin in human health. Acta Medica Bulgarica 36(2), 65–69 (2009)
55. Davis Presley, S.I., KarthikRamanan, V.G., Prasanth, S.M.: Isolation of curcuminoids from turmeric using non chlorinated solvents. J. Crit. Rev. (2020). https://doi.org/10.31838/jcr.07.04.129
56. BaradaranRahimi, V., Askari, V.R., Hosseinzadeh, H.: Promising influences of Scutellaria baicalensis and its two active constituents, baicalin, and baicalein, against metabolic syndrome: A review”. Phytother. Res. 35(7), 3558–3574 (2021)
57. Roy, M.K., et al.: Baicalin, a flavonoid extracted from a methanolic extract of Oxyryum indicum induces proliferation of a cancer cell line in vitro via induction of apoptosis. Pharmazie 62(2), 149–153 (2007)
58. Kumar, S., Yadav, M., Yadav, A., Rohilla, P., Yadav, J.P.: Antiplasmodial potential and quantification of aloin and aloe- emodin in Aloe vera collected from different climatic regions of India. BMC Complement. Altern. Med. 17(1), 369 (2017)
59. Wang, L., et al.: Ultrasonic extraction and separation of anthraquinones from Rheum palmatum L. Ultrason. Sonochem. 15(5), 738–746 (2008)
60. Dong, X., et al.: Aloe-emodin: A review of its pharmacology, toxicity, and pharmacokinetics. Phytother. Res. 34(2), 270–281 (2020)
61. Baiatour, O., et al.: Salt effect on phenolics and antioxidant activities of Tunisian and Canadian sweet marjoram (Origanum majorana L.) shoots: Salt effect on phenolics and antioxidant activities of Origanum majorana L. shoots. J. Sci. Food Agric. 93(1), 134–141 (2013)
62. Oubada, A., Garca, M., BellobalArcon, A., CuestaRubio, O., Monzote, L.: Antiileishmanial activity of leaf extract from Calophyllum rutilare against Leishmania amazonensis. Emir. J. Food Agric. 26(9), 807 (2014)
63. Erenler, R., et al.: Isolation and identification of chemical constituents from Origanum majorana and investigation of antipro liferative and antioxidant activities: Isolation and identification of chemical constituents from Origanum majorana. J. Sci. Food Agric. 96(3), 822–836 (2016)
64. Hazafa, A., Rehman, K.-U., Jahan, N., Jabeen, Z.: The role of polyphenol (flavonoids) compounds in the treatment of cancer cells. Nutr. Cancer 72(3), 386–397 (2020)
65. Haidari, F., Keshavarz, S.A., Mohammad Shahi, M., Mahboob, S.-A., Rashidi, M.-R.: Effects of parsley (Petroselium crispum) and its flavonol constituents, kaempferol and quercetin, on serum uric acid levels, biomarkers of oxidative stress and liver xanthine oxidoreductase activity inOxonate-induced hyperuricemic rats. Iran. J. Pharm. Res. 10(4), 811–819 (2011)
66. Zhang, Q., Zhou, M.-M., Chen, P.-L., Cao, Y.-Y., Tan, X.-L.: Optimization of ultrasonic-assisted enzymatic hydrolysis for the extraction of luteolin and apigenin from celery. J. Food Sci. 76(5), C680–C685 (2011)
67. Magar, R.T., Sohng, J.K.: A review on structure, modifications and structure-activity relation of quercetin and its derivatives. J. Microbiol. Biotechnol. 30(1), 11–20 (2020)
68. Redford, K.E., Abbott, G.W.: The ubiquitous flavonoid quercetin as a target for KCNQ potassium channel activator. Commun. Biol. 3(1), 356 (2020)
69. Sharifi, N., Mahernia, S., Amanlou, M.: Comparison of different methods in quercetin extraction from leaves of Raphanus sativus L. Pharm. Sci. 23(1), 59–65 (2017)
70. Yi, L.-T., et al.: Antidepressant-like effects of psoralidin isolated from the seeds of Psoralea Corylifolia in the forced swimming test in mice. Prog. Neuropsychopharmacol Biol. Psychiatry 32(2), 510–519 (2008)
71. Miao, Y.-H., Hu, Y.-H., Yang, J., Liu, T., Sun, J., Wang, X.-J.: Natural source, bioactivity and synthesis of benzoferan derivatives. RSC Adv. 9(47), 27510–27540 (2019)
72. Glisan, L., Grove, K.A., Yennawar, N.H., Lambert, J.D.: Inhibition of pancreatic lipase by black tea theaflavins: Comparative enzymology and in silico modeling studies. Food Chem. 216, 296–300 (2017). https://doi.org/10.1016/j.foodchem.2016.08.052
73. Gülc¸in, I ˙, Huyut, Z., Elmastas ¸, M., Aboul-Enein, H.Y.: Radical scavenging and antioxidant activity of tannic acid. Arab. J. Chem. 3(1), 43–53 (2010). https://doi.org/10.1016/j.arabjc.2009.12.008
74. Versari, A., du Toit, W., Parpinello, G.P.: Oenological tannins: a review. Aust. J. Grape Wine Res. 19(1), 1–10 (2012). https://doi.org/10.1111/jagw.12002
75. Bordignon-Luiz, M.T., Gauche, C., Gris, E.F., Falcão, L.D.: Colour stability of anthocyanins from Isabel grapes (Vitis labrusca L.) in model systems. LWT Food Sci. Technol. 40(4), 594–599 (2007). https://doi.org/10.1016/j.lwt.2006.02.022
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