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Coronavirus herd immunity optimizer with greedy crossover for feature selection in medical diagnosis

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ABSTRACT

The importance of medical data and the crucial nature of the decisions that are based on such data, as well as the large increase in its volume, has encouraged researchers to develop feature selection (FS)-based approaches to identify the most relevant data for specific medical problems. In this paper, two intelligent wrapper FS approaches based on a new metaheuristic algorithm named the coronavirus herd immunity optimizer (CHIO) were applied with and without the incorporation of a greedy crossover (GC) operator strategy to enhance exploration of the search space by CHIO. The two proposed approaches, CHIO and CHIO-GC, were evaluated using 23 medical benchmark datasets and a real-world COVID-19 dataset. The experimental results indicated that CHIO-GC outperformed CHIO in terms of search capability, as reflected in classification accuracy, selection size, F-measure, standard deviation, and convergence speed. The GC operator was able to enhance the balance between exploration and exploitation of the CHIO in the search and correct suboptimal solutions for faster convergence. The proposed CHIO-GC was also compared with two previous wrapper FS approaches, namely, binary moth flame optimization with Lévy flight (LBFMO_V3) and the hyper learning binary dragonfly algorithm (HLBDA), as well as four filter methods namely, Chi-square, Relief, correlation-based feature selection and information gain. CHIO-GC surpassed LBFMO_V3 and the four filter methods with an accuracy rate of 0.79 on 23 medical benchmark datasets. CHIO-GC also surpassed HLBDA with an accuracy rate of 0.93 when applied to the COVID-19 dataset. These encouraging results were obtained by striking a sufficient balance between the two search phases of CHIO-GC during the hunt for correct solutions, which also increased the convergence rate. This was accomplished by integrating a greedy crossover technique into the CHIO algorithm to remedy the inferior solutions found during premature convergence and while locked into a local optimum search space.

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1. Introduction

It is hard for a human to retrieve essential information from the large volume of data stored and disseminated by numerous health research centers around the world, which adversely affects the capacity of medical staff to extract the necessary knowledge from medical data [1]. Moreover, given that the health and medical care industry is one of the key industries that does not tolerate incorrect actions being taken as a result of inaccurate data processing, many artificial intelligence approaches have been used to improve the consistency of medical information as these approaches are capable of handling the growing amount of data in a highly efficient manner [2,3]. Through the use of artificial intelligence, including machine learning, it is envisaged that it will be possible to create healthcare applications that can perform as well as better than human physicians in certain tasks [4,5]. Machine learning is able to link, analyze and present data in a more intelligible manner, which then enables human medical practitioners to make accurate decisions and take appropriate action [6,7].

Information extracted by artificial intelligence techniques involves the use of a conceptual relationship that expresses the data in a new manner that is more understandable and meaningful to the data owner, without making any assumptions about what the knowledge within that data could be [8,9]. However, physical examination is a crucial task, and failure for it can affect the safety and efficiency of the overall treatment process [10,11]. There is a
lot of data that a classifier must take into consideration, although it is not related to the problem of the study and is not related to it [12,13]. Therefore, the selection of appropriate data or features to which to apply the classifier will increase the efficiency of the results produced by the classifier and at the same time reduce the time consumed by the learning model, especially when the volume of data is large [14,15]. Thus feature selection (FS) is considered a critical process for enhancing the efficiency of a learning algorithm [16].

The FS problem essentially involves finding a way to select the lowest number of relevant features from the original dataset that often comprises a massive variety of features [17]. In a large dataset; certain features are linked to the problem of interest while others are not. If all the features were chosen, this would definitely have an effect on the search results either in terms of time consumed or classification accuracy [18]. Thus, the objective of the FS process is to reduce the dimension of the search space as much as possible, but not at the cost of accuracy [19]. Therefore, the success of the selection task relies on two essential aspects: decreasing the number of features and increasing classification accuracy [20,21].

The FS process involves a generation process, evaluation phase, meeting the termination criterion and completing a validation procedure [22]. These four stages can be achieved by applying a FS method, such as a filter, wrapper, embedded or hybrid method. The wrapper method is distinct from the filter method in that it employs a learning algorithm during the evaluation phase, whereas the filter method evaluates specific features independently of the classification process by using a certain standard threshold [23,24]. The embedded method is similar to the wrapper method as a classifier is used in the selection process in the evaluation step, but the use of the classifier in the embedded method is comparatively less cost-effective than in the wrapper method [25]. On the other hand, the hybrid method sequentially employs a filter and a wrapper method and hence the selection of features involves two iterations. First, the filter is used to produce a subset of features and then the wrapper is used to pick features from a subset obtained from first step [26].

During the generation phase, a set of features is selected from the full dataset to decide whether it matches the solution or not [27]. Basically, each feature is examined to create the best subset, either through a process of forward selection or backward elimination, which increases the degree of complexity by 2^n [28]. To reduce the time it takes to generate feature subsets in this phase, an optimization method is often used as a search strategy. Optimization techniques are estimation processes and the results obtained by these techniques are either optimal or suboptimal. One of the well-known and widely used optimization methods is the metaheuristic algorithm [29–32].

Metaheuristic algorithms are intelligent algorithms that are based on the concept of identifying a particular mathematical tool with the aim of optimizing a specific problem [33]. Improvements are made by several frequent implementation attempts in order to find the correct solution for a particular problem [34]. These intelligent algorithms utilize the knowledge gathered during the search to guide the search process, during which they iteratively create new solutions by integrating one or more good solutions, and they are often also combined with some kind of operator in order to prevent them from becoming stuck in a local optimum [35]. While metaheuristics try to find the optimal solution, they are typically imperfect mechanisms as they do not ensure that the best global solution is found. Rather, they often produce approximate results [36].

Two types of searches are performed by metaheuristic algorithms to find an optimal solution: exploration and exploitation [37]. In the exploration phase, the search traverses numerous sites and different environments to explore and discover more areas for high-quality solution. Population-based metaheuristic algorithms are exploration-oriented [38,39]. On the other hand, in the exploitation phase, existing resources are focused on a particular search area. Single-based metaheuristic algorithms are considered to be exploitation-oriented [40–42].

Numerous metaheuristics have been proposed and are widely used to solve FS problems in different research domains. These include the monarch butterfly optimization algorithm (MBO) [43–45], chaotic dragonfly algorithm (CDA) [46], whale optimization algorithm (WOA) [47], spotted hyena optimizer (SHO) [48], atom search optimization (ASO) [49], chaotic interior search algorithm, equilibrium optimizer algorithm (EOA) [50], and chaotic competitive swarm optimization (CCSO) [51] among many others [52–59].

In this study, a new metaheuristic algorithm named the coronavirus herd immunity optimizer (CHIO), which was developed by Al-Betar et al. [60] in 2020, is implemented to solve FS problems in the medical diagnosis domain. The CHIO simulates herd immunity, which is considered to be a means to combat a viral pandemic. It was inspired by the coronavirus known as SARS-CoV-2 or COVID-19 which caused a global pandemic during 2020. The extent of the spread of coronavirus infection depends on how infected individuals communicate directly with other community members and herd immunity can prevent other people from acquiring the infection. In the current study, the CHIO is implemented in two different ways to select the most effective features in medical datasets. First, it is applied in its original form. Then, the exploration capability of the CHIO is enhanced by using a greedy crossover (GC) operator in an approach named CHIO-GC. The two proposed approaches are applied in a wrapper model using a K nearest neighbor (KNN) classifier, and evaluated using 23 medical benchmark datasets, as well as a COVID-19 dataset as a case of a real-world problem dataset. In addition, the performance of the two proposed approaches is compared against other methods in the literature. The analysis revealed that integrating a greedy crossover technique into the CHIO algorithm produced results that were more accurate than those produced using CHIO in its original form. This indicates that CHIO-GC has the ability to remedy the inferior solutions found during premature convergence and while locked into a local optimum search space.

The remainder of this paper is structured as follows: the works most important to this study are presented in Section 2. The suggested FS methods, CHIO, and CHIO-GC, are discussed in Section 3, 4, and 5, respectively. The tests and the findings are discussed in Section 6. Lastly, the conclusion and some possible directions for study are discussed in Section 7.

2. Related work

Feature selection has been used in a wide range of problems, including image processing, sentiment analysis, intrusion detection, and language identification as well as many other domains [61–67]. However, one of the challenges that still needs to be overcome in respect of the use of FS process is its use in the field of medical diagnosis, which is the focus of this research. Therefore, in this section, the most recent work on the use of FS in medical diagnosis will be reviewed. Several different approaches have been proposed in this regard.

For instance, Li et al. [68] employed a hybrid approach for FS in medical diagnoses using a genetic algorithm (GA) to produce sustainable initial positions and a gray wolf optimization (GWO) to modify the existing population positions in a discrete search area. In experiments, the proposed approach was applied to disease diagnosis problems, and the results demonstrated that the
suggested hybrid approach is superior in terms of classification accuracy as compared to the original GA and the GWO. On the other hand, Zuo et al. [69] suggested using a filter-based FS method that explicitly uses the Menger curvature to rate all the features in an electronic health records dataset. The results showed that after reducing the number of features, high classification accuracy is achieved by this method as compared to previous methods.

Anter and Ali [70] designed a hybrid FS solution that combines the crow search optimization algorithm with chaos theory and fuzzy c-means (CFCSA). The suggested CFCSA uses the global optimization approach to prevent local minima trapping and chaos theory to resolve the lack of CSA convergence. Experiments showed that the CFCSA outperforms in terms of mean fitness and standard deviation as compared to other methods such as bat algorithm and the binary crow search algorithm.

In another work, Wang and Chen [71] developed a hybrid learning system that employs a WOA that blends chaotic and multi-swarm techniques to concurrently tackle parameter optimization and FS, as well as to optimize the efficiency of a support vector machine (SVM) to diagnose various diseases. However, the results indicated that the SVM generated by the proposed approach is actually inferior to other profitable SVM methods based on the original WOA, particle swarm optimization (PSO), bacterial foraging optimization and the GA in terms of both classification accuracy and selection size.

Furthermore, Rostami et al. [72] proposed a FS model that incorporates the idea of using node centrality and the PSO algorithm. The proposed scheme consists of three main processes. In the first step, the initial features are visualized as a graphic representation model. In the next step, the core features of all the nodes in the graph are determined. Finally, the enhanced PSO-based search method is used to pick the final features Experiments were done on five medical datasets and the results showed that the proposed approach improves on the reliability and efficacy of previous related approaches.

In research conducted by Verma et al. [73], a cost-sensitive medical diagnostic is regarded as a FS problem, in which each test provides a feature that is used to predict a prediction model. The aim of their study was to find the optimal balance between accuracy and cost. To this end, the researchers used the “weak dominance” property of a set to create a decision model that defines a collection of features in order to offer an “optimal” trade-off between the cost and accuracy of prediction without including knowledge of the true features of the medical state. The findings confirmed the efficiency of the proposed method in respect of optimization problems generated by real-world datasets. Moreover, the FS process was also applied in [74] on a skin disease dataset by different classification techniques. The FS process in this approach enhances the dermatological prediction accuracy.

Kuppuchamy and Mangayararkarasi [75] concentrated on using fuzzy entropy to assess the importance of the feature in the diagnosis of breast cancer. In their study, a number of FS strategies were implemented to obtain useful subsets of features. In addition, the radial base function network was used as a classifier. The Wisconsin Breast Cancer dataset was used in the experiment and the findings showed that high classification accuracy was achieved with reduced selection size. The Wisconsin Diagnostic Breast Cancer dataset was also utilized by Rahman and Muniyandi [76] in their work on selecting effective features by using a FS technique. They used a 15-neuron neural network to classify the cancer. The results showed a significant improvement of up to 99.4% in classification accuracy in comparison with other methods. Another FS technique was proposed by de Lima et al. [77]. The researchers’ technique was based on a twin-bound support vector machine (FSTBSVM). The experiment revealed that the proposed method is very effective and capable of delivering good results with limited features as compared to using the original datasets.

A metaheuristic algorithm was applied for FS in relation to medical issues by Too and Mirjaliil [78]. Specifically, the researchers implemented a hyper-learning binary dragonfly algorithm (HLBDA) in a wrapper FS approach to find optimum feature subsets from over 21 datasets as well as a COVID-19 dataset. The findings revealed the supremacy of HLBDA in terms of increasing the classification accuracy and reducing the number of features chosen in comparison with eight previous works.

On the other hand, Abu Khurma et al. [79] improved the moth flame optimization (MFO) algorithm in two directions. In the first, eight binary variants are generated using eight transition functions. In the second, the Lévy flight operator is incorporated into the MFO structure in association with the transition functions, and named LBMFO-V3. It was shown that the suggested LBMFO V3 method is able to greatly outpace several well-known wrapper methods in 83% of datasets. Also, the suggested methodology surpasses other approaches in the literature approaches in 75% of the datasets. Also, a comparison with the filter-based methodology indicated that the proposed LBMFO-V3 approach is superior across 70% of the datasets.

From the above overview of related works, it can be seen that many metaheuristic algorithms have been used to solve FS problems in the medical diagnosis domain. The findings of previous research studies have shown that these smart algorithms can identify the best related features that can maximize classification accuracy. The effectiveness of these algorithms is attributed to the consistency of their random search mechanism and to their ability to strike a balance between local and global search processes.

In light of the above, in this study, two intelligent FS wrapper strategies based on a new metaheuristic algorithm called CHIO were applied with a greedy crossover operator strategy to enhance CHIO exploration for FS in the field of medical diagnosis. A crossover strategy permits individuals to exchange genetic information during the development of subsequent generations of individuals. A greedy algorithm is a step-by-step method that guarantees that the following step delivers the greatest possible value on the way to a solution. It has been shown that optimization problems can also be solved by using a greedy method [80]. This is because a greedy algorithm can eventually remove the problem if better judgments can be made at any phase, and then find an optimal solution to the entire problem can be discovered.

3. Coronavirus herd immunity optimizer (CHIO)

The CHIO is a new metaheuristic algorithm that was proposed by Al-Betaret al. in 2020 [60]. Similar to many other metaheuristic algorithms, it mimics the behavior of a natural entity, in this case taking its inspiration from a pathogenic coronavirus. The CHIO imitates the process of achieving natural immunity in a herd through the implementation of herd psychology, which is known to be one of the methods of obtaining immunity from infectious diseases.

In 2020, for the third time in as many decades, a pathogenic coronavirus crossed species to infect the human population. The virus, unofficially labeled 2019-nCoV, was first observed in Wuhan, China, in people who had been exposed to seafood or a wet market [81]. The swift response of the Chinese public health and scientific community contributed to the recognition of the related clinical disease and provided initial awareness of the epidemiology of the infection [81]. Acquired immunity is developed by a human getting a normal infection via a pathogen
or by receiving an injection, often a vaccine. Herd immunity stems from the effect of the extent of human immunity on the larger herd [82]. It can be described as indirect immunity against infection that is given to susceptible persons when a reasonably large proportion of individuals in the population are resistant to the infection [62].

Herd immunity relies upon the period a disease remains inside an infected host and the pace at which the disease spreads. The introduction of a single infected person into a group of already vulnerable individuals would result in the continuous indiscriminate spreading of a disease among any of those who were approached by the infected person before such infected individuals died or recovered. The estimated number of individuals who become infected in such a vulnerable population is the so-called simple reproduction number [83]. The disease would be spread to other susceptible contacts by the persons who had acquired the disease from the original infected person, and this mechanism would repeat itself until the disease infected the whole population [64]. However, the presence of herd immunity could lead to the complete elimination of the disease from a society, and, as long as any member of the population has immunity to the disease, the potential of the disease to spread would decline. The decline in the rate of disease spread would be dependent on the size of the immunized herd. Nevertheless, even if total herd immunity could not be achieved, the effects of the disease could be mitigated by the presence of a “buffer” of resistant individuals [85].

The idea of coronavirus herd immunity was mathematically modeled by Al-Betar et al. to create a theoretical optimization algorithm named the CHIO. The model is based on the finding a way to best protect humanity against disease by converting the bulk of the helpless non-infected population into a robust population [60]. As a consequence, all the remaining vulnerable cases would not be affected and the resistant population would no longer transmit the disease. In the model, the population of herd immunity individuals are classified into three categories: susceptible, contaminated (or confirmed) and immunized (or recovered) persons [60,86]. A susceptible individual is a person who is not born with or afflicted with the virus. However, a vulnerable individual may be contaminated by interaction with infectious individuals who have refused to comply with the recommended social distancing or gap. An infected individual is a person who can spread the virus to susceptible individuals who are in close contact with the psychological distancing factor. An immunized individual is a person who is protected from infection and does not threaten untreated individuals. Therefore, this type of individual can help prevent the spreading of the virus to others and thereby avert the triggering of a pandemic [64].

Fig. 1 provides an illustration of the population hierarchy in the herd immunity scenario and the effect on acquiring immunity on the above-described three categories of individuals in the population.

It can be seen from Fig. 1 that herd immunity can be depicted as a tree in which the infectious individual is at the base or root and the branches correspond to the other individuals contacted. The right-hand portion of Fig. 1 shows that the virus cannot be spread to contacted persons if the root individual is immunized.

The herd immunity strategy can be modeled as an optimization algorithm that consists of six main phases [50]. Each of these phases is discussed in turn below:

### Phase 1: Initialization

The CHIO parameters and the issue of optimization are addressed in this step. In respect of objective functionality, the optimization problem is formulated as shown in Eq. (1):

$$\text{Min } f(x) \quad x \in [Lb, Ub]$$  \hspace{1cm} (1)

where $f(x)$ is the measured objective function (or immunity rate) that is computed for the individual $x_i = (x_1, x_2, \ldots, x_n)$, where $x_i$ is the gene indexed by $i$, and $n$ represents the number of genes in each individual. Notice that each gene’s value range is $x_i \in [lbi, ubi]$, where $lbi$ is located. The highest and lowest boundaries of gene $x_i$ are expressed by $Lbi$ and $Ubi$. The CHIO algorithm has four algorithmic parameters and two operational parameters. The four algorithmic parameters are (1) $C_0$, which is the number of preliminary cases of infection initiated by one individual; (2) HIS, which is the size of the population; (3) $Max_Itr$, which is the actual number of iterations; and (4) $n$, which represents the problem dimensionality.

In this stage, two major control parameters of the CHIO are initialized: (1) the basic reproduction rate ($BRr$), which regulates the operators of the CHIO by propagating the coronavirus among the individuals, and (2) the maximum age of infected cases ($Max_Age$), which determines the classification of the infected cases as either having recovered or died.

### Phase 2: Generate initial herd immunity population

The CHIO produces a set of cases (individuals) that is equal to HIS spontaneously (or heuristically). In the herd immunity population (HIP), the generated cases are stored as a two-dimensional matrix of size $n \times HIS$ as follows:

$$\text{HIP} = \begin{bmatrix}
    x_1^1 & x_2^1 & x_3^1 \\
    x_1^2 & x_2^2 & \cdot \\
    \vdots & \vdots & \ddots \\
    HIS & x_{10}^{HIS} & x_{n}^{HIS}
\end{bmatrix}$$  \hspace{1cm} (2)

in which each row $j$ represents a case $x_j$ that is generated basically. This includes $x_j = Lbi + (Ubi - Lbi) \times U(0, 1)$, $\forall i = 1, 2, \ldots n$. The objective function (or immunity rate) is determined by using Eq. (1) for each situation. In addition, the HIS duration status variable $(S)$ for all HIP cases is initiated by either zero (susceptible case) or one case (infected case). Note that the random initiation of the number of ones in $(S)$ is as many as $C_0$.

### Phase 3: Evolve coronavirus herd immunity

The evolution phase is the CHIO’s primary enhancement loop, where gene $x'_i$ in case $x_i$, according to the proportion of the $BRr$, either remains the same or changes according to the influence of social distancing based on the following three rules for infected, susceptible and immune cases:

$$x_i(t+1) \rightarrow \begin{cases}
    x_i'(t) & r \geq BRr \\
    C \left(x_i'(t)\right) & r < \frac{1}{3} \times BRr \quad (\text{infected}) \\
    N \left(x_i'(t)\right) & r < \frac{2}{3} \times BRr \quad (\text{susceptible}) \\
    R \left(x_i'(t)\right) & r < BRr \quad (\text{immune})
\end{cases}$$  \hspace{1cm} (3)

where $r$ produces a number generator between 0 and 1. The three rules are described below:

1. **Infected case**

   Under the spectrum of $r \in [0, \frac{1}{3}BRr]$ any social gap is caused by the new gene value of $x_i'(t+1)$, which is determined by the discrepancy between the present gene and a gene obtained from a contaminated case $x'_i$, such as

   $$x_i'(t+1) = C(x_i'(t))$$  \hspace{1cm} (4)

   where

   $$C \left(x_i'(t)\right) = x_i'(t) + r \times (x_i'(t) - x_i^c(t))$$  \hspace{1cm} (5)
Notice that the value $x_i^t (t)$ is arbitrarily selected on the basis of a condition vector $(S)$ from every contaminated case $x^r$, so that $c = |i|S(i) = 1$.

2. Susceptible case

The new gene value of $x_i^t (t + 1)$ is influenced by any social gap within the spectrum of $r \in [\frac{1}{2} Br_r, \frac{2}{3} Br_r]$, which is determined by the discrepancy between the present gene and a gene extracted from a compromised case $x^m$, such as

$$x_i^t (t + 1) = N(x_i^t (t))$$

where

$$N (x_i^t (t)) = x_i^t (t) + r \times (x_i^t (t) - x_i^m (t))$$

(6)

Notice that the value $x_i^m (t)$ is distributed from every resistant case $x^m$ randomly, and that it is centered on a vector of status $(S)$ given that $m = |i|S(i) = 0$.

3. Immune case

The new gene value of $x_i^t (t + 1)$ is influenced by any social gap within the spectrum of $r \in [\frac{1}{2} Br_r, Br_r]$, which is determined by the discrepancy between the present gene and a gene extracted from a compromised case $x^r$, such as

$$x_i^t (t + 1) = R(x_i^t (t))$$

where

$$R (x_i^t (t)) = x_i^t (t) + r \times (x_i^t (t) - x_i^r (t))$$

(7)

Notice that the value $x_i^r (t)$ is distributed from every resistant case $x^r$ randomly, and that it is centered on a vector of status $(S)$ given that $r = \arg \min_{|i|S(i)=1} f(x_i^r)$.

4. Update herd immunity population

The immunity rate $f (x^t (t + 1))$ of each case $x^t (t + 1)$ generated is determined and the actual case $x^t (t)$ is replaced by the obtained case $x^t (t + 1)$ if the obtained case is stronger, such that $f (x^t (t + 1)) < f (x^t (t))$. Also, the age vector $A_j$ is increased by a value of 1 if $S_j = 1$. For each event, the state vector $(S_j)$ is modified $x^t$ based on the herd immune criterion that uses the following equation:

$$\begin{align*}
S_j \rightarrow \left\{ \begin{array}{ll}
1 & f (x^t (t + 1)) < f (x^t (t + 1)) < \frac{f (x^t (t + 1))}{\Delta f (x)} \wedge S_j = 0 \wedge \text{is.corona}(x_i^t (t + 1)) \\
2 & f (x^t (t + 1)) < \frac{f (x^t (t + 1))}{\Delta f (x)} \wedge S_j = 1
\end{array} \right.
\end{align*}$$

(9)

where the binary value of is.corona $(x^t (t + 1))$ is equal to 1 when the new value is a value from any infected case that has been inherited by case $x^t (t + 1)$. Also, the $\Delta f (x)$ is the mean significance of the immune population rates such as $\overline{\sum_{i=1}^{n} f(x_i)}$.

Notice that the immunity levels of the individuals in the population are altered depending on the social gap measured earlier. If the newly produced individual immunity rate is better than the population’s average immunity rate, this means that the population is becoming more immune to the virus. If the recently discovered population is sufficiently strong to be immune to the virus, then the threshold of herd immunity has been reached.

Phase 5: Fatal cases

In this phase, if the immunity rate of the current infected case $(S_j=$1) cannot be strengthened as defined by the $Max_Age$ parameter (i.e., $Aj > Max_Age$), then this case is considered dead. However, using $x_i^t (t + 1) = Lbi + (Ubi - Lbi) \times U(0, 1)$, $\forall i = 1, 2, \ldots, n$ is then regenerated from scratch. In addition, $Aj$ and $S_j$ are both set to 0. This phase may be beneficial in diversifying the current population and thereby avoiding local optima.

Phase 6: Stop criterion

The CHIO algorithm repeats step 3 to step 5 until the termination criterion is reached, which normally depends on whether...
the maximum number of iterations is reached. In this case, the population is dominated by the total number of susceptible and immunized cases. Also, the infected cases are passed.

All the above phases of the CHIO algorithm are illustrated as a flowchart in Fig. 2.

The pseudocode of the six CHIO phases is given below:

**CHIO algorithm pseudocode**

1. **Step 1:** Initialize the CHIO parameters \(\text{HIS}, \text{Max_itr}, \text{and Max_Age}\)

2. **Step 2:** Generate herd immunity population

3. \(C_{CHIO} = Lbi + (Ubi - Lbi) \times U(0,1), \forall i = 1, 2, \ldots, n\) and \(\forall j = 1, 2, \ldots, \text{HIS}\)

4. calculate the fitness of each search agent

5. set \(Sj=0 \forall j = 1, 2, \ldots, \text{HIS}\)

6. set \(Aj=0 \forall j = 1, 2, \ldots, \text{HIS}\)

7. **Step 3:** Herd immunity evolution

8. while \((t \leq \text{Max_itr})\) do

9. for \(j = 1 \text{ to } \text{HIS} \) do

10. is Corona\(x'_j(t+1)\) = false

11. for \(i = 1 \text{ to } N \) do

12. if \((r < 1/3 \times BRr)\) then

13. \(x'_j(t+1) = C(x'_j(t))\)

14. if Corona\(x'_j(t+1)\) = true

15. else if \((r < 2/3 \times BRr)\) then

16. \(x'_j(t+1) = N(x'_j(t))\)

17. else if \((r < BRr)\) then

18. \(x'_j(t+1) = R(x'_j(t))\)

19. else

20. \(x'_j(t+1) = x'_j(t)\)

21. end if

22. end for

**Step 4:** Update herd immunity population

23. if \(f(x'_j(t+1)) < f(x'_j(t))\) then

24. \(f(x'_j(t)) = f(x'_j(t+1))\)

25. else

26. \(Aj = Aj + 1\)

27. end if

28. end if

29. if \(f(x'_j(t+1)) < \frac{f(x'_j(t+1))}{f(x'_j(t))}Sj = 0\) is Corona\(x'_j(t+1)\) then

30. \(Sj=1, Aj=1\)

31. end if

32. if \(f(x'_j(t+1)) < \frac{f(x'_j(t+1))}{f(x'_j(t))}Sj = 1\) then

33. \(Sj=2, Aj=0\)

34. end if

**Step 5:** Fatality condition

35. if \(Aj > M \times \text{Max_Age} \) and \(S=1\) then

36. \(x'_j = Lbi + (Ubi - Lbi) \times U(0,1), \forall i = 1, 2, \ldots, n\)

37. \(Aj=0\)

38. \(Sj=0\)

39. end if

40. end for

41. \(t = t + 1\)

42. end while

4. **Greedy crossover (GC) operator**

One of the essential search operators is the crossover operator. The main purpose of using this operator is to generate a new promising optimal solution by merging current parent solutions \([87]\). This is seen as an effective technique as the search process will then theoretically lead to new exploration regions where better solutions can be sought \([88]\). The crossover operator takes two solutions and combines them in order to create a new one that is distinct from the previous solutions because it selects the best features of both solutions to form an optimal solution. There are several forms of crossover operators, each of which relies on a particular mechanism \([89]\). In this study, a methodology based on a greedy approach proposed in \([90]\) was used to find the best solutions in the crossover phase. Here, it was specifically implemented to enhance the exploration capability of the CHIO.

A greedy algorithm is a step-by-step approach as it ensures that the next step offers the maximum potential value on the route to a solution \([91]\). A greedy algorithm can also be used to solve optimization problems \([80]\). If better decisions can be made in any step and an optimal solution to the whole problem can be found, a greedy algorithm can ultimately eliminate the problem.

In this study, the application of a greedy strategy first involves the random selection of two parent solutions \(S_A\) and \(S_B\) from the population by CHIO, the first component of the offspring (\(S_O\)) apoints the values between \(S_A\) and \(S_B\) by performing an intersection process between the values, where \(S_O = S_A \cap S_B\).

Then, the remainder of the offspring is proceeded to a greedy strategy based on the potential \(P_x\), which extends \(S_O\) step by step by assigning one element to it at each step until the offspring includes precisely \(N\) elements. In the following, the process of the greedy strategy is explained in more detail. Fig. 3 describes the idea of the GC strategy used in this study.

Let \(N = \{S_1, S_2, \ldots, S_N\}\) be a set of elements, and let \(d_{XY}\) be the distance between them, where \(S_A\) and \(S_Y\) \((d_{XY} = d_{XY})\) elements with \(d_{XY} > 0\) if \(X \neq Y\) and \(d_{XY} = 0\) otherwise. Then, potential \(P_x\) can be calculated, taking into consideration the objective function \(f\) in Eq. (13) in next section, as the following equation:

\[
P_x = \sum_{S_Y \in S} d_{XY}, \quad S_x \in N
\] (11)

In the first step, all the elements in \(S_A\) are evaluated to define the element with the highest potential relative to \(S_0\) and move it from \(S_A \cap S_0\) to \(S_0\). We then take into account the elements in \(S_A \cap S_0\), specify the element with the greatest potential in \(S_A \cap S_0\) and move it to \(S_0\). Then, at each point of this greedy process, we consider the elements in \(S_A \cap S_0\) and \(S_B \cap S_0\) in turn until \(S_0\) reaches the size of \(N\). The offspring in \(S_0\) is usually a relatively high-quality solution.

The features are swapped into the offspring in specific steps. Let swap \(S_x, S_y\) define a transfer that switches \(S_x \in S\) and \(S_y \in N \backslash S\). Then, as swap \(S_x, S_y\) is used the target variation \([90]\), which can be easily measured by:

\[
Target\ Variation = f(S') - f(S) = P_Y - P_x - d_{XY}
\] (12)

where \(S' = S \cup \{S_x, S_y\}\) while \(P_Y\) and \(P_x\) are, respectively, the potential of \(S_x, S_y\) according to Eq. (15).

The population updating process determines whether to come up with a solution for offspring, which is the crossover should be a part of the population and, if not, the current population solution will be substituted. Population management is a critical issue because updating the population rule actually influences the diversity of the population, which in turn affects the convergence of the CHIO search. Algorithm 2 provides the pseudocode of the GC operator.
Algorithm 2: Greedy crossover operator pseudocode

1. CHIO generalizes two parent solutions $S_A$ and $S_B$
2. Greedy crossover: One offspring solution $S_O$:
   3. $S_O \leftarrow S_A \cap S_B$ /* Establish a partial solution first by retaining mutual common features by $S_A$ and $S_B$ */
   4. while $|S_O| < N$ do
      5. Select from $S_A \setminus S_O$ the element $u$ with the highest potential with respect to $S_O$
      6. $S_O \leftarrow S_O \cup \{u\}$, $S_A \leftarrow S_A \setminus \{u\}$
      7. if $|S_O| = N$ then
         8. Return $S_O$ and Stop
      9. end if
   10. Select from $S_B \setminus S_O$ the element $v$ with the highest potential with respect to $S_O$
   11. $S_O \leftarrow S_O \cup \{v\}$, $S_B \leftarrow S_B \setminus \{v\}$
   12. end while
   13. Return $S_O$

5. CHIO-GC for the FS problem

The FS process is used to delete redundant, obsolete and misleading features in order to obtain the best subset that represents the best outcome, where every feature is relevant if the choice depends on it, otherwise it is irrelevant. A feature is considered to be redundant if it is heavily associated with other features. The FS process is a binary optimization problem where solutions are limited to binary values (0, 1). This implies that any optimization strategy used to solve FS problems needs to be built in binary form, such that solutions are represented as either 0 or 1 in a one-dimensional vector. A feature that is selected is assigned the value 1, otherwise 0. Fig. 4 gives an example of the binary representation of selected features.

In this study, the wrapper FS approach is based on KNN, which determines the accuracy rate of the proposed approaches, CHIO and CHIO-GC, for the FS process in medical diagnosis. According to related works, the KNN classifier has been found to have good classification efficiency when applied to FS problems [4]. In this study, the number of nearest neighbors (K) was five. The 5-NN algorithm was used for the fitness assessment during the training period with internal N-fold cross-validation, where the number of folds was five; the average error rate in the classification procedure was determined for each fold of each equivalent particle. The number of folds (N) and the number of nearest neighbors (K) were chosen based on previous research.

The wrapper FS approach involved a generation, evaluation, a termination criterion and validation phase. In the wrapper
approach, the learning algorithm was used as a part of the evaluation phase. This approach created an interaction between the search subset and the classification algorithm. Therefore the KNN classifier was used twice, one time in the evaluation and one time in the validation phase. During the generation step, a subset of features was chosen from the full dataset for the validation process to decide whether or not it matched the solution. In this phase, the exploration and exploitation capabilities of the CHIO algorithm were used to search for and generate a subset from a given dataset. To increase the exploration efficiency of the CHIO to the maximum possible degree, a crossover operator was used in a greedy manner to find the best solutions before CHIO moved on to the exploitation part of the search process. Fig. 5 describes the CHIO with GC for FS based on a wrapper approach using KNN.

The efficiency of the suggested approaches CHIO and CHIO-GC were assessed according to accuracy, recall, precision, F-measure and the number of features (selection size).

Classification accuracy is calculated by:

\[
\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN} \tag{13}
\]

where:
- True positives (TPs) are cases in which the model predicted true and the actual output was also true.
- True negatives (TNs) are cases in which the model predicted false and the actual output was false.
- False positives (FPs) are cases in which the model predicted true and the actual output was false.
- False negatives (FNs) are cases in which the model predicted false and the actual output was true.

Precision describes how accurate the learning model is in terms of how many of the cases that the model predicted as positive are actually positive. Precision is calculated by:

\[
\text{precision} = \frac{TP}{TP + FP} \tag{14}
\]

Recall represents how many of the actual positives the model captures by labeling them as positive (true positive). Recall is calculated by:

\[
\text{recall} = \frac{TP}{TP + FN} \tag{15}
\]

The F-measure expresses the balance between the ratio of recall and precision; the closer it is the higher and close to the degree of accuracy. The F-measure is calculated by:

\[
F - \text{measure} = 2 \left( \frac{\text{precision} \cdot \text{recall}}{\text{precision} + \text{recall}} \right) \tag{16}
\]

6. Experimental results

This section describes the experimental setup and presents the analysis of the results, as well as comparisons with previous methods in order to assess the performance of the proposed method. The instability that has been generated depends on a variety of criteria, including the accuracy rate, the rate of convergence and certain measurements of central inclination. In order to conduct a fair scientific analysis, similar work environments and conditions were observed throughout the experiments. The experiments were carried out using an Intel® Core™ i7-6006U Processor @ 2.00 GHz (four CPUs), ~2.0 GHz with 8 GB of RAM. The CHIO was introduced using Matlab R2016a. The datasets were divided into 70% for training and 30% for testing. The tests were carried out over 30 runs for each dataset and 100 iterations were used in each run.

6.1. Parameter settings

In the experiments, the input parameters were determined by the results of some initial tests which enabled the proposed method to produce better output. In order for the results of the experiment to be the same, the algorithm configurations were identical throughout. Table 1 provides the CHIO parameter values that were used in all the experiments.

As for the KNN classifier, the input was the nearest training instances in the feature space and the output was a class membership. The labeling method depended on the majority of votes
Fig. 5. Proposed CHIO-GC approach.

| Table 1  | Parameter settings. |
|----------|---------------------|
| **Parameter** | **Value** |
| HIS       | 30 |
| Max_Age   | 100 |
| BRr       | 0.01 |
| Max_Itr   | 100 |
| LB (lower bound) | 0 |
| UB (upper bound) | 1 |

6.2. Description of the datasets

The term medical data refers to health-related data that are used to determine routine patient treatment or as part of a diagnostic trial program. There are several categories of such data, such as administrative data, claims data, patient disease data and clinical trial data, among others. In this study, in order to be able to generalize the results of the experiments, we used two types of medical data: a number of medical benchmark datasets and one real-world COVID-19 dataset.

6.2.1. Medical benchmark datasets

Twenty-three well-known benchmarked datasets of diverse patient data were used in the experiments. They were downloaded from a range of data repositories such as UCI, KEEL, and Kaggle, as well as other well-known websites for FS medical

| Table 2  | Characteristics of the medical benchmark datasets. |
|----------|---------------------------------------------------|
| **Dataset** | **Number of features** | **Number of instances** | **Number of classes** | **Source of dataset** |
| 1  | Diagnostic | 30 | 569 | 2 | UCI |
| 2  | Original  | 9 | 699 | 2 | UCI |
| 3  | Prognostic | 33 | 194 | 2 | UCI |
| 4  | Coimbra  | 9 | 115 | 2 | UCI |
| 5  | BreastEW  | 30 | 596 | 2 | UCI |
| 6  | Retinopathy | 19 | 115 | 2 | UCI |
| 7  | Dermatology | 34 | 366 | 6 | UCI |
| 8  | ILPD-Liver  | 10 | 583 | 2 | UCI |
| 9  | Lymphography | 18 | 148 | 4 | UCI |
| 10  | Parkinsons | 22 | 194 | 2 | UCI |
| 11  | ParkinsonC | 753 | 755 | 2 | UCI |
| 12  | SPECT | 22 | 267 | 2 | KEEL |
| 13  | Cleveland | 13 | 297 | 5 | KEEL |
| 14  | HeartEW | 13 | 270 | 2 | KEEL |
| 15  | Hepatitis | 18 | 79 | 2 | KEEL |
| 16  | SAHear | 9 | 461 | 2 | KEEL |
| 17  | Spectfheart | 43 | 266 | 2 | KEEL |
| 18  | Thyroid0387 | 21 | 7200 | 3 | KEEL |
| 19  | Heart | 13 | 302 | 5 | Kaggle |
| 20  | Pima-diabetes | 9 | 768 | 2 | Kaggle |
| 21  | Leukemia | 7129 | 72 | 2 | Kaggle |
| 22  | Colon | 2000 | 62 | 2 | Kaggle |
| 23  | Prostate_GE | 5966 | 102 | 2 | Kaggle |
As can be seen from Table 2, the 23 datasets include many case studies on medical diagnosis and have different structures. The power and reliability of the CHIO and CHIO-GC can be discovered by studying the improved optimizer on different problems with different characteristics. The datasets were divided into two parts: 70% for training and 30% for testing. The tests were carried out over 30 runs for each dataset and 100 iterations were used in each run.

6.2.2. COVID-19 dataset – a real-world dataset

In March 2020, the World Health Organization confirmed that extreme acute respiratory syndrome coronavirus 2 (SARS-CoV-2) or COVID-19, which had emerged in China in late 2019, had reached pandemic status. At the time of writing it has resulted in the death of millions of people worldwide. Artificial intelligence is increasingly being used in a range of technologies for diagnosis, identification and prevention in the global fight against COVID-19. Hence it seemed appropriate that the CHIO and CHIO-GC were applied to predict the health of COVID-19 patients. For this purpose, a real-world dataset on COVID-19 patients was obtained from https://github.com/AtharvaPeshkar/Covid-19-Patient-Health-Analytics. The dataset consisted of 15 features, as listed in Table 3.

In this study, patient data containing missing values for both “death” and “recovery” were excluded from the key dataset. For the experiments, the data were split evenly into two sets of training and testing data for the evaluation process.

7. Results and discussion

In order to evaluate the efficacy of the proposed approaches, CHIO and CHIO-GC, seven outcomes were taken into account: accuracy, error rate, number of features chosen (selection size), precision, recall, F-measure, boxplot and convergence speed. The CHIO and CHIO-GC were compared in terms of accuracy rate and selection size. The results achieved by the two approaches when applied to each of the 23 datasets and the COVID-19 dataset after 30 runs are provided in Table 4.

As can be seen from Table 4, CHIO-GC achieved higher accuracy in all datasets. This suggests that the CHIO is capable of generating more reliable results if its search mechanism is modified. It can also be seen from the table that the CHIO-GC approach was able to reduce the gap between the min and max accuracy values of the CHIO within 30 runs. In all datasets, the accuracy result was not less than that of the basic CHIO and, at the same time, the maximal values were part of the recommended CHIO-GC approach.

As for the FS size, the CHIO-GC demonstrated an advantage over the CHIO in terms of the number of features selected in 17 out of the 24 datasets, namely, Diagnostic, Original, Prognostic, Coimbra, BreastEW, ILPD-Liver, Lymphography, Parkinsons, ParkinsonC, SPECT, SAHear, Thyroid0387, Heart, Leukemia, Colon, Prostate_GE and COVID-19. These results demonstrate the power of the modification in the CHIO-GC to improve the exploration capability of the CHIO to find the best primitive solutions.

The precision, recall and F-measure values of the two approaches were also evaluated in order to further test the results and the extent of the classifier's ability to provide reliable, correlated and result values are equivalent in all sequences for each dataset. Precision represents the ratio of positive IDs that were actually right, while recall reflects the ratio of true positive IDs that were correctly detected, and the F-measure denotes the equilibrium between the recall and precision ratios. The precision, recall and F-measure values were determined by Eqs. (14), (15), and (16), respectively. The degree to which the efficiency of the CHIO and CHIO-GC approaches is adapted and concentrated in all of the datasets used in the experiment is shown in Table 5.

As shown in Table 5, all the F-measure values were higher than the accuracy values shown in Table 4 in all datasets except BreastEW and HeartEW. A large amount of real negatives, which in most technical situations are not relied upon, could have made a significant contribution to the results of the accuracy test. Even though FNs and FPs usually have market costs (quantifiable and non-quantifiable), the F-measure may be a better assessment to use where equilibrium between accuracy and recall is sought and the distribution of classes is inconsistent.

Precision, recall, and the F-measure give a more precise assessment of a classifier's behavior because they can be used to obtain a more in-depth judgment of the classifier's ability to find the correct results in learning than by assessing its performance based on accuracy alone. When the consequences of false positives are significant, precision becomes a useful assessment metric, whereas when the cost of false negatives is significant, recall can provide further insight into the results. Also, the F-measure is useful for understanding the tradeoff between accuracy and coverage when categorizing positive cases because it provides a more accurate estimate of wrongly categorized instances than the accuracy metric. The F-measure provides an overall assessment of a model's reliability that mixes precision and recall, in the same way as addition and multiplication can mix two components to produce a different result entirely. Hence, a strong F1 score indicates the presence of a small sample size of false positives and false negatives, thus false alarms do not affect the results.

In addition, the T-test was used to compare the efficiency of the CHIO and CHIO-GC approaches. Using these proposed approaches, which rely on the precision of the results specific to each dataset, the findings statistics are carried out. By conducting a T-test, with a 95% spectrum of significance (alpha = 0.05) on the p-values obtained and the classification accuracy the, different corresponding statistics are shown in Table 6.

As can be seen from Table 6, the efficiency of the CHIO-GC is slightly higher than that of the original CHIO, where most of the P-values for the 24 datasets are less than 0.0001. These findings show that the use of the CHIO-GC is effective for the solution of FS problems.

A boxplot is a charting technique for displaying a five-number summary. The interquartile range denotes the location of the data’s middle part. The first quartile (the 25% mark) and third quartile (the 75% mark) are located at the respective ends of the box. The chart’s lowest point is on the far left, while its maximum is on the far right. The median is indicated by a vertical bar in the center of the box. A boxplot indicates how closely grouped the data are and whether they are symmetrical. It also exposes
| Table 4 | Accuracy and feature selection size results for CHIO and CHIO-GC. |
|--------|-------------|
| Dataset | CHIO | CHIO-GC | CHIO | CHIO-GC | CHIO | CHIO-GC | CHIO | CHIO-GC |
| 1 Diagnostic | 0.8540 | 0.9033 | 0.91 | 0.96 | 0.79 | 0.84 | 14.4000 | 13.3700 |
| 2 Original | 0.9233 | 0.9710 | 0.99 | 0.99 | 0.85 | 0.94 | 6.2000 | 5.1040 |
| 3 Prognostic | 0.5293 | 0.6716 | 0.82 | 0.77 | 0.47 | 0.60 | 4.7422 | 4.6202 |
| 4 Crohn's | 0.6866 | 0.8896 | 0.87 | 0.91 | 0.70 | 0.86 | 4.6667 | 3.6007 |
| 5 BreastEW | 0.8993 | 0.9400 | 0.94 | 0.97 | 0.85 | 0.89 | 15.8333 | 13.7303 |
| 6 Retinopathy | 0.4660 | 0.6436 | 0.61 | 0.69 | 0.38 | 0.60 | 7.4667 | 7.2647 |
| 7 Dermatology | 0.6690 | 0.8006 | 0.73 | 0.87 | 0.55 | 0.70 | 18.5000 | 18.4900 |
| 8 ILPD-Liver | 0.6423 | 0.7716 | 0.69 | 0.78 | 0.60 | 0.72 | 4.1098 | 4.0000 |
| 9 Lymphography | 0.7606 | 0.8343 | 0.82 | 0.91 | 0.69 | 0.79 | 10.1667 | 10.0622 |
| 10 Parkinsons | 0.6690 | 0.7903 | 0.73 | 0.85 | 0.57 | 0.68 | 7.4667 | 7.2647 |
| 11 ParkinsonC | 0.6856 | 0.8400 | 0.79 | 0.88 | 0.58 | 0.78 | 4.1098 | 4.0000 |
| 12 SPECT | 0.6073 | 0.6960 | 0.69 | 0.88 | 0.55 | 0.60 | 6.4000 | 7.0105 |
| 13 Cleveland | 0.4896 | 0.5966 | 0.58 | 0.64 | 0.44 | 0.55 | 6.8097 | 6.8097 |
| 14 HeartEW | 0.8540 | 0.9116 | 0.91 | 0.94 | 0.79 | 0.87 | 6.4000 | 7.0105 |
| 15 Hepatitis | 0.6690 | 0.7903 | 0.73 | 0.85 | 0.57 | 0.68 | 4.1098 | 4.0000 |
| 16 GCHear | 0.6420 | 0.7036 | 0.70 | 0.73 | 0.59 | 0.68 | 7.4667 | 7.2647 |
| 17 Spectfheart | 0.6716 | 0.7303 | 0.77 | 0.79 | 0.60 | 0.68 | 7.4667 | 7.2647 |
| 18 Thyroid0387 | 0.8986 | 0.9603 | 0.96 | 0.98 | 0.82 | 0.92 | 10.0314 | 8.0116 |
| 19 Heart | 0.7316 | 0.8126 | 0.79 | 0.87 | 0.70 | 0.77 | 8.1000 | 6.1505 |
| 20 Pima-diabetes | 0.7153 | 0.7956 | 0.86 | 0.87 | 0.61 | 0.68 | 5.4667 | 7.0105 |
| 21 Leukemia | 0.9876 | 0.9900 | 1.00 | 1.00 | 0.91 | 0.93 | 3597.4427 | 3560.5107 |
| 22 Colon | 0.7606 | 0.8343 | 0.82 | 0.91 | 0.69 | 0.79 | 10.1667 | 10.0622 |
| 23 Prostate_GE | 0.5555 | 0.6174 | 0.68 | 0.85 | 0.59 | 0.72 | 3.6000 | 3.6000 |
| COVID-19 dataset | 0.9135 | 0.9370 | 0.9482 | 0.9770 | 0.8402 | 0.8818 | 4.1100 | 3.0500 |

| Table 5 | Precision, recall and F-measure results for CHIO and CHIO-GC. |
|--------|-------------|
| Dataset | CHIO | CHIO-GC | CHIO | CHIO-GC | CHIO | CHIO-GC | CHIO | CHIO-GC |
| 1 Diagnostic | 0.913978 | 0.954436 | 0.850000 | 0.954436 | 0.880829 | 0.954436 |
| 2 Original | 0.945455 | 0.973568 | 0.945455 | 0.977876 | 0.945455 | 0.975717 |
| 3 Prognostic | 0.689655 | 0.821333 | 0.714286 | 0.800000 | 0.701754 | 0.810526 |
| 4 Coimbra | 0.825581 | 0.944444 | 0.855422 | 0.876289 | 0.840237 | 0.909091 |
| 5 BreastEW | 0.604651 | 0.780488 | 0.603774 | 0.782222 | 0.696970 |
| 6 Retinopathy | 0.750000 | 0.843854 | 0.677419 | 0.814103 | 0.711864 | 0.880829 |
| 7 Dermatology | 0.790576 | 0.858586 | 0.848315 | 0.904255 | 0.818428 | 0.880829 |
| 8 ILPD-Liver | 0.720000 | 0.814016 | 0.720000 | 0.924731 | 0.720000 | 0.891192 |
| COVID-19 dataset | 0.912938 | 0.991736 | 0.907452 | 0.937500 | 0.910187 | 0.963855 |

the existence and coordinates of any outliers. Fig. 6 shows the boxplots that describe the distribution of the performance of CHIO and CHIO-GC when applied to the 24 datasets over 30 runs.

Note from Fig. 6 that the greedy approach was able to reduce the gap between the minimum and maximum accuracy values of the CHIO algorithm, and bring them closer to the value of the mean. Also, the minimum and maximum values of the CHIO-GC were higher than that of the basic algorithm. This is a clear indication of the ability of the CHIO-GC approach to improve the balance between exploration and exploitation of the CHIO search to produce accurate results.

Fig. 7 shows how the GC strategy was able to reduce the gap between the maximum and minimum accuracy values, and thus maximize accuracy. The accuracy values of the CHIO-GC approach were never worse than the CHIO values in all datasets. The results in the figure were arrived at by averaging the values of CHIO and CHIO-GC over the 30 runs for all datasets.

It is well known that a stable and rapid rate of convergence will lead to better solutions. Thus, in order to further test the efficiency of the CHIO and CHIO-GC, the convergence speed behavior curves of the two approaches were obtained by applying them to each of the 24 datasets over 30 different iterations. In general, any metaheuristic algorithm may take a large number of iterations to reach the optimum point. Therefore, it is important to employ methods that have a convergence rate that is as fast as possible. An algorithm's rate of convergence is usually measured by the number of iterations and by the number of function evaluations that are needed to obtain an acceptable solution.

The results in Fig. 8 indicate that the CHIO-GC was able to boost the global search of the original CHIO and thus improve...
The above results indicated that the CHIO-GC performed better than the original CHIO. In order to further assess the reliability of the CHIO-GC, and its ability to produce a high degree of classification accuracy while at the same time minimizing the number of attributes, it was compared with six methods in the literature. First, the CHIO-GC was compared with LBMFO-V3 [79] by using the 23 medical benchmark datasets. Then, it was compared against HLBD [78] using the COVID-19 dataset. Finally, its performance was compared with that of four filter methods, namely, Chi-square, Relief, correlation-based feature selection (CFS) and information gain (IG).

7.1. Comparison with previous methods

The above results indicated that the CHIO-GC performed better than the original CHIO. In order to further assess the reliability of the CHIO-GC, and its ability to produce a high degree of classification accuracy while at the same time minimizing the number of attributes, it was compared with six methods in the literature. First, the CHIO-GC was compared with LBMFO-V3 [79] by using the 23 medical benchmark datasets. Then, it was compared against HLBD [78] using the COVID-19 dataset. Finally, its performance was compared with that of four filter methods, namely, Chi-square, Relief, correlation-based feature selection (CFS) and information gain (IG).

7.1.1. Comparison with LBMFO-V3

The CHIO-GC was compared with LBMFO-V3 in terms of average classification accuracy and number of selected features using the 23 medical benchmark datasets. Table 7 shows the results of this comparison.
Fig. 6. Boxplots of CHIO and CHIO-GC for all datasets.
Table 7 shows that the CHIO-GC outperformed LBMFO-V3 in terms of classification accuracy in 16 datasets, namely, Original, Prognostic, BreastEW, Retinopathy, ILPD-Liver, Lymphography, Parkinsons, ParkinsonC, SPECT, Cleveland, Hepatitis, SAHear, Spectheart, Heart, Colon and Prostate_GE. The CHIO-GC approach had an overall accuracy of 0.7983 in all datasets, as compared to LBMFO-V3, which achieved 0.7746.

Moreover, the CHIO-GC also performed better than LBMFO-V3 in terms of selection size in 13 datasets, namely, Diagnostic, Original, Prognostic, BreastEW, Parkinsons, ParkinsonC, SPECT, Hepatitis, SAHear, Thyroid0387, Heart, Leukemia and Prostate_GE. The two approaches achieved the same result in one dataset, namely, ILPD-Liver. The CHIO-GC approach generated an overall selection size of 351.4142 features in all datasets, and it overcame the LBMFO-V3, which got 351.9462 features. Fig. 9 graphically illustrates the average accuracy and selection size achieved by CHIO-GC and LBMFO-V3 in all 23 datasets.

### 7.1.2. Comparison with HLBDA

The CHIO-GC was compared with the HLBDA in terms of classification accuracy and number of selected features using the COVID-19 dataset. In terms of classification accuracy the CHIO-GC had an 0.9370 average accuracy rate over 30 runs as compared to 0.9221 obtained by the HLBDA. On the other hand, the HLBDA generated an average of three features in all runs, whereas the CHIO-GC obtained a rate of four features. The features that were selected most frequently by the CHIO-GC across the 30 runs were location, country, age and symptom2. Fig. 10 shows the average accuracy and selection size results for the CHIO-GC and HLBDA using COVID-19 dataset.

### 7.1.3. Comparison with filter methods

The classification accuracy results of the CHIO-GC, which is a wrapper-based approach, were also compared against those of four general filter-based approaches, namely, Chi-square, relief, CFS and IG. These four filter methods offer a high assurance of the intensity values of the datasets. Although wrapper models require that the predictor is optimized as part of the selection phase, filter models focus on the general characteristics of the training data to choose features that are independent of some predictor. The filters used in this part of the analysis were accessed from the WEKA data mining program [92]. Table 8 shows the average

| Dataset                  | Average accuracy | Selection size |
|--------------------------|------------------|----------------|
| CHIO-GC                  | LBMFO-V3         | CHIO-GC        |
| Diagnostic               | 0.9370           | 10.3700        |
| Original                 | 0.9100           | 13.9991        |
| Prognostic               | 0.6676           | 13.5604        |
| BreastEW                 | 0.9400           | 13.7303        |
| Retinopathy              | 0.8806           | 14.6202        |
| Dermatology              | 0.7776           | 15.0126        |
| CHIO-GC                  | LBMFO-V3         | CHIO-GC        |
| ILPD-Liver               | 0.8343           | 17.5033        |
| Parkinsons               | 0.7933           | 18.7383        |
| ParkinsonC               | 0.8400           | 19.3082        |
| SPECT                    | 0.6960           | 19.6050        |
| Cleveland                | 0.5966           | 20.1799        |
| HeartEW                  | 0.9116           | 20.7383        |
| Hepatitis                | 0.7933           | 21.2011        |
| SAHear                   | 0.7036           | 21.6503        |
| Spectheart               | 0.7303           | 21.7903        |
| Thyroid0387              | 0.9603           | 22.0116        |
| Heart                    | 0.8126           | 22.1503        |
| Pima-diabetes            | 0.7933           | 22.7383        |
| Leukemia                 | 0.9900           | 23.5033        |
| Colon                    | 0.7176           | 24.0000        |
| Prostate_GE              | 0.6010           | 24.7944        |
Fig. 8. Convergence speed of CHIO and CHIO-GC.
Table 8

| Dataset             | CHIO-GC | Chi-square | Relief | CFS     | IG     |
|---------------------|---------|------------|--------|---------|--------|
| Diagnostic          | 0.9033  | 0.5714     | 0.9585 | 0.9533  | 0.9349 |
| Original            | 0.9710  | 0.9091     | 0.6426 | 0.6860  | 0.6759 |
| Prognostic          | 0.6716  | 0.5910     | 0.7727 | 0.7576  | 0.7577 |
| Lymphography        | 0.8896  | 0.3846     | 0.6672 | 0.5763  | 0.5578 |
| BreastEW            | 0.9400  | 0.9365     | 0.8160 | 0.8029  | 0.8128 |
| Retinopathy         | 0.6436  | 0.6349     | 0.7727 | 0.7576  | 0.7577 |
| Dermatology         | 0.7716  | 0.7120     | 0.5119 | 0.5223  | 0.5264 |
| ILPD-Liver          | 0.3843  | 0.8824     | 0.5886 | 0.5533  | 0.5204 |
| Parkinsons          | 0.7903  | 0.7581     | 0.7588 | 0.7360  | 0.7150 |
| ParkinsonC          | 0.8400  | 0.6593     | 0.6590 | 0.6487  | 0.6276 |
| SPECT               | 0.6966  | 0.3940     | 0.1181 | 0.0398  | 0.0826 |
| Cleveland           | 0.9116  | 0.9334     | 0.6153 | 0.5757  | 0.6202 |
| HeartEW             | 0.7903  | 0.7778     | 0.5538 | 0.5857  | 0.6417 |
| Hepatitis           | 0.7036  | 0.6471     | 0.5024 | 0.5115  | 0.5227 |
| SAHear              | 0.7303  | 0.7090     | 0.6079 | 0.6279  | 0.5551 |
| Spectfheart         | 0.9603  | 1.0000     | 0.6379 | 0.6955  | 0.9773 |
| Thyroid0387         | 0.8126  | 0.5333     | 0.6317 | 0.5575  | 0.6114 |
| Heart               | 0.7956  | 0.6905     | 0.5147 | 0.5426  | 0.5264 |
| Pima-diabetes       | 0.9900  | 0.7120     | 0.6883 | 0.6759  | 0.6410 |
| Leukemia            | 0.7176  | 0.5850     | 0.5641 | 0.5116  | 0.5097 |
| Colon               | 0.6010  | 0.5042     | 0.5033 | 0.4786  | 0.4421 |
| Prostate_GE         | 0.7983  | 0.7046     | 0.6133 | 0.5887  | 0.5981 |

It can be observed from Table 8 that the CHIO-GC exceeded IG in all datasets. It also exceeded CFS and Relief in all datasets except Diagnostic and Prognostic, respectively. On the other hand, Chi-square performed better than the three other filter methods, and surpassed the results produced by the CHIO-GC in four datasets, namely, Lymphography, SPECT, HeartEW and Thyroid0387. However, overall, the CHIO-GC defeated all the filter methods in 17 datasets with an accuracy rate of 0.7983. Fig. 11 shows the accuracy rate of the CHIO-GC and the four filter methods.

7.2. Discussion

The results produced by the original CHIO indicate that the algorithm has a suitable balance between exploration and exploitation in its search mechanism. This balance is one of the most important strengths of metaheuristic algorithms as it helps them to find the best solutions during the search process. In the CHIO-GC approach, the exploration capability of the CHIO was modified by applying a greedy crossover operator to select the initial features with the aim of maximizing the solution. This modification contributed to an enhancement of the balance between exploration and exploitation. The stronger balance that was achieved also enabled the CHIO-GC to accelerate the rate of convergence during its search for the best solutions. The selection of the most appropriate features ensures that the identified solutions converge to the maximum. Thus exploration by means of randomization facilitates the search of the solution space from a local point of view and at the same time increases the variety of solutions.

The success of the proposed CHIO-GC approach in achieving a good balance between exploration and exploitation was demonstrated in the experiments in several ways. First, the CHIO-GC outperformed the CHIO in all 24 datasets in terms of classification accuracy, as shown in Table 4. Furthermore, the superiority of the CHIO-GC was also observed in the maximum and minimum accuracy it was able to achieve in each run, as shown in Table 4, Figs. 6 and 7. The CHIO-GC was able to minimize the gap between the maximum and the minimum accuracy and make them converge. Also, the convergence speed results showed the power of modify the exploration search which is made up of two criteria, the number of iteration needed to get the optimal solution and the initial starting point of solution. So, as shown in Fig. 8, the CHIO-GC was better than the CHIO in terms of convergence speed as it had a good initial starting point and it did not need more than 30 iterations to obtain the optimal solution in most datasets.

Moreover, the CHIO-GC outperformed LBMFO-V3 in 16 datasets and HLDBA in the COVID-19 dataset in terms of both classification accuracy and selection size. Hence the inclusion of GC in the CHIO approach was proved to be beneficial in correcting the suboptimal solutions that were reached at premature
convergence and when trapped in a suboptimal search space. Furthermore, the CHIO-GC demonstrated its superiority when applied to large datasets such as ParkinsonC, Leukemia, Colon and Prostate_GE. Therefore, the proposed the CHIO-GC approach can be relied upon to solve FS problems if the problem becomes larger.

8. Conclusion

The FS problem is among the most critical issues facing researchers in many fields including medical diagnosis. In recent years, metaheuristics have been commonly used for FS to try to minimize the number of features required to achieve sufficiently reliable results, with the goal of increasing reliability and enhancing performance. In this study, a new metaheuristic named the coronavirus herd immunity optimizer (CHIO) was implemented to solve FS problems in medical diagnosis.

The CHIO was applied as a basic algorithm and as a modified algorithm using greedy crossover (CHIO-GC) to enhance exploration. Two types of dataset were used to assess the proposed approaches: 23 medical benchmark datasets and a real-world COVID-19 dataset. The evaluation of the two approaches was conducted in respect of several criteria, including classification accuracy, number of selected features, error rate, precision, recall, F-measure, boxplot, convergence speed and T-test. All the obtained results indicated that the CHIO-GC enhanced the exploration capability of the original CHIO.

In comparison experiments, the CHIO-GC outperformed two FS wrapper approaches, LBMFO-V3 and the HLBD. The CHIO-GC surpassed LBMFO-V3 in 16 out of the 23 medical benchmark datasets with an accuracy rate of 0.79 and selection size rate of 351 features. It also outdid the HLBD when applied to the COVID-19 dataset, with a classification accuracy of 0.93. Furthermore, the wrapper-based CHIO-GC surpassed four filter methods, namely, Chi-square, Relief, CFS and IG.

It is considered that these promising results were achieved through the strong balance between the two search phases of the CHIO-GC during the discovery of the right solutions, which also accelerated the convergence rate. This was achieved by incorporating a greedy crossover method into the CHIO algorithm to correct the suboptimal solution reached on premature convergence and when trapped in a local optimum search space.

In future work, researchers may wish to consider hybridizing the CHIO with another single-based metaheuristic algorithm such as simulated annealing to try to enhance its exploitation (local) search capability, or applying it in another FS field such as intrusion detection or image segmentation. Furthermore, many other computational intelligence algorithms can be used to solve medical diagnosis problems, such as monarch butterfly optimization (MBO), the earthworm optimization algorithm (EWA), elephant herding optimization (EHO), the moth search (MS) algorithm, the slime mold algorithm (SMA), and Harris hawks optimization (HHO).

CRediT authorship contribution statement

Mohammed Alweshah: Design and implementation of the research, Analysis of the results, Writing of the manuscript. Saleh Alkhalaileh: Design and implementation of the research, Analysis of the results, Writing of the manuscript. Mohammed Azmi Al-Betar: Design and implementation of the research, Analysis of the results, Writing of the manuscript. Azuraliza Abu Bakar: Design and implementation of the research, Analysis of the results, Writing of the manuscript.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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References

[1] W. Li, G.-G. Wang, A.H. Gandomi, A survey of learning-based intelligent optimization algorithms, Arch. Comput. Methods Eng. (2021) 1–19.
[2] S.-H. Liew, Y.-H. Choo, Y.F. Low, Fuzzy-rough classification for brainprint authentication, Jordan. J. Comput. Inf. Technol. (JJCT) 5 (02) (2019).
[3] K. Dyczkowski, Intelligent medical decision support system based on imperfect information, in: The Case of Ovarian Tumor Diagnosis, Studies in Computational Intelligence, Springer, 2018.
[4] M. Alweshah, S. Alkhalaileh, D. Albashish, M. Mafarja, Q. Bsoul, O. Dorgham, A hybrid mine blast algorithm for feature selection problems, Soft Comput. 25 (1) (2021) 517–534.
[5] S. Sengan, G. Kamalam, J. Vellingiri, J. Gopal, P. Velayutham, V. Subramaniyaswamy, Medical information retrieval systems for e-health care records using fuzzy based machine learning model, Microprocess. Microsyst. (2020) 103344.
[63] M. Lui, T. Baldwin, Cross-domain feature selection for language identification, in: Proceedings of 5th international joint conference on natural language processing, 2011, pp. 553–561.

[64] H. Alazzam, A. Sharieh, K.E. Sabri, A feature selection algorithm for intrusion detection system based on pigeon inspired optimizer, Expert Syst. Appl. 76 (2020) 113249.

[65] A. Alarifi, A. Tolba, Z. Al-Makhadmeh, W. Said, A big data approach to sentiment analysis using greedy feature selection with cat swarm optimization-based long short-term memory neural networks, J. Supercomput. 76 (6) (2020) 4414–4429.

[66] M. Toğaçar, B. Ergen, Z. Cömert, F. Özyurt, A deep feature learning model for pneumonia detection applying a combination of mRMR feature selection and machine learning models, Irbm 41 (4) (2020) 212–222.

[67] S.R. Ahmad, A.A. Bakar, M.R. Yaakub, A review of feature selection techniques in sentiment analysis, Intell. Data Anal. 23 (1) (2019) 159–189.

[68] Q. Li, et al., An enhanced grey wolf optimization based feature selection wrapped kernel extreme learning machine for medical diagnosis, Comput. Math. Methods Med. 2017 (2017).

[69] Z. Zuo, J. Li, N.A. Moubayed, Curvature-based feature selection with application in classifying electronic health records, 2021, arXiv preprint arXiv:2101.03581.

[70] A.M. Anter, M. Ali, Feature selection strategy based on hybrid crow search optimization algorithm integrated with chaos theory and fuzzy c-means algorithm for medical diagnosis problems, Soft Comput. 24 (3) (2020) 1565–1584.

[71] M. Wang, H. Chen, Chaotic multi-swarm whale optimizer boosted support vector machine for medical diagnosis, Appl. Soft Comput. 88 (2020) 105946.

[72] M. Rostami, S. Forouzandeh, K. Berahmand, M. Soltani, Integration of multi-objective PSO based feature selection and node centrality for medical datasets, Genomics 112 (6) (2020) 4370–4384.

[73] A. Verma, M.K. Hanawal, N. Hemachandra, Unsupervised online feature selection for cost-sensitive medical diagnosis, in: 2020 International Conference on Communication Systems & NETworks (COMSNETS), IEEE, 2020, pp. 1–6.

[74] A.K. Verma, S. Pal, S. Kumar, Prediction of skin disease using ensemble data mining techniques and feature selection method—a comparative study, Appl. Biochem. Biotechnol. 190 (2) (2020) 341–359.

[75] R. Kuppuchamy, M. Mangayarkarasi, A threshold fuzzy entropy based feature selection approach for breast cancer diagnosis.

[76] M.A. Rahman, R.C. Muniyandi, An enhancement in cancer classification accuracy using a two-step feature selection method based on artificial neural networks with 15 neurons, Symmetry 12 (2) (2020) 271.

[77] M.B. de Lima, J.C. de Lima, R.M. Barbosa, Medical data set classification using a new feature selection algorithm combined with twin-bounded support vector machine, Med. Biol. Eng. Comput. 58 (3) (2020) 519–528.

[78] J. Too, S. Mirjalili, A hyper learning binary dragonfly algorithm for feature selection: A COVID-19 case study, Knowl.-Based Syst. 212 (2020) 106553.

[79] R.A. Khurmaa, I. Aljarah, A. Sharieh, An intelligent feature selection approach based on moth flame optimization for medical diagnosis, Neural Comput. Appl. (2020) 1–40.

[80] Z. Yang, W. Wang, M. Shi, Algorithms and complexity for a class of combinatorial optimization problems with labelling, J. Optim. Theory Appl. 1–23.

[81] A.F. Henwood, Coronavirus disinfection in histopathology, J. Histotechnol. 43 (2) (2020) 102–104.

[82] T. Britton, F. Ball, P. Trapman, A mathematical model reveals the influence of population heterogeneity on herd immunity to SARS-CoV-2, Science 369 (6505) (2020) 846–849.

[83] D.R. Smith, Herd Immunity, the Veterinary clinics of North America, Food Anim. Pract. 35 (3) (2019) 593–604.

[84] M.G.M. Gomes, et al., Individual variation in susceptibility or exposure to SARS-CoV-2 lowers the herd immunity threshold, 2020, MedRxiv.

[85] V.J. Clemente-Suíárez, et al., Dynamics of population immunity due to the herd Effect in the COVID-19 pandemic, Vaccines 8 (2) (2020) 236.

[86] J.S. Lavine, A.A. King, O.N. Bjerndal, Natural immune boosting in pertussis dynamics and the potential for long-term vaccine failure, Proc. Natl. Acad. Sci. 108 (17) (2011) 7259–7264.

[87] O. Gokalp, E. Yildizi, A. Ugur, A novel wrapper feature selection algorithm based on iterated greedy metaheuristic for sentiment classification, Expert Syst. Appl. 146 (2020) 113176.

[88] B. Koohestani, A crossover operator for improving the efficiency of permutation-based genetic algorithms, Expert Syst. Appl. 151 (2020) 113381.

[89] J. Thomas, N.S. Chaudhari, Selection of efficient crossover operator in metaheuristic approach for 2D strip packing, in: 2013 IEEE International Conference on Systems, Man, and Cybernetics, IEEE, 2013, pp. 415–420.

[90] Q. Wu, J.-K. Hao, A hybrid metaheuristic method for the maximum diversity problem, European J. Oper. Res. 231 (2) (2013) 452–464.

[91] F.B. Ozsoydan, M. Sağir, Iterated greedy algorithms enhanced by hyper-heuristic based learning for hybrid flexible flowshop scheduling problem with sequence dependent setup times: a case study at a manufacturing plant, Comput. Oper. Res. 125 (2021) 105044.

[92] M. Hall, E. Frank, G. Holmes, B. Pfahringer, P. Reutemann, I.H. Witten, The WEKA data mining software: an update, ACM SIGKDD Explor. Newsl. 11 (1) (2009) 10–18.