Machine learning approaches for the mortality risk assessment of patients undergoing hemodialysis

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Abstract
Introduction: Mortality is a major primary endpoint for long-term hemodialysis (HD) patients. The clinical status of HD patients generally relies on longitudinal clinical observations such as monthly laboratory examinations and physical examinations.

Methods: A total of 829 HD patients who met the inclusion criteria were analyzed. All patients were tracked from January 2009 to December 2013. Taken together, this study performed full-adjusted-Cox proportional hazards (CoxPH), stepwise-CoxPH, random survival forest (RSF)-CoxPH, and whale optimization algorithm (WOA)-CoxPH model for the all-cause mortality risk assessment in HD patients. The model performance between proposed selections of CoxPH models were evaluated using concordance index.

Results: The WOA-CoxPH model obtained the highest concordance index compared with RSF-CoxPH and typical selection CoxPH model. The eight significant parameters obtained from the WOA-CoxPH model, including age, diabetes mellitus (DM), hemoglobin (Hb), albumin, creatinine (Cr), potassium (K), Kt/V, and cardiothoracic ratio, have also showed significant survival difference between low- and high-risk characteristics in single-factor analysis. By integrating the risk characteristics of each single factor, patients who obtained seven or more risk characteristics of eight selected parameters were dichotomized as high-risk subgroup, and remaining is considered as low-risk subgroup. The integrated low- and high-risk subgroup showed greater discrepancy compared with each single risk factor selected by WOA-CoxPH model.

Conclusion: The study findings revealed WOA-CoxPH model could provide better risk assessment performance compared with RSF-CoxPH and typical selection CoxPH model in the HD patients. In summary, patients who had seven or more risk characteristics of eight selected parameters were at potentially increased risk of all-cause mortality in HD population.

Keywords: feature selection, hemodialysis, machine learning, risk assessment, survival analysis

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Introduction
Patients with end-stage kidney disease (ESKD) require long-term renal replacement therapy, such as hemodialysis (HD) and peritoneal dialysis. However, several clinical factors have been reported to influence long-term HD treatment outcomes such as serum albumin levels. In addition, the HD treatment also becomes an economic burden to the reimbursement for medical care. Therefore, HD-related diagnosis, care planning, and prevention have become critical research issues. The all-cause mortality is commonly considered as a primary endpoint for patients undergoing long-term HD in the clinical...
In current clinical medical research, it is no longer possible to consider only univariate association; there are complex interactions between multiple clinical factors and biomarkers for survival outcome in HD population. Therefore, it is necessary to explore the relevant risk factors that affect the disease in a multivariate and comprehensive manner. In the era of big data, there may be limits to complete care for all clinical factors and biomarkers using typical statistical methods. Therefore, machine learning methods can be used to quickly find out more significant risk factors for diseases to achieve maintenance and slow down the occurrence of disease exacerbations.

In 2017, Ramspek et al. used Cox proportional hazards (CoxPH) model to predict risk of all-cause mortality in dialysis patients. Multiple machine learning approaches, including artificial neural network, particle swarm optimization, biogeography-based optimization, and other hybrid methods have been widely used in the risk assessment of specific diseases. Comparing with typical statistical approaches, the combination use of machine learning could overcome several limitations faced by statistical methods, including the sample size restriction and computation complexity. In 2022, Radović et al. used kernel support vector machine and K-means to determine the expected mortality rate. Random survival forest (RSF) is an exploratory analysis method used to evaluate survival data. RSF uses the survival splitting rules for growing survival tree to identify highly impacted risk factor of mortality. Garcia-Montemayor et al. evaluated the prediction performance of random forest and logistic regression in mortality of HD patients.

Whale optimization algorithm (WOA) is a meta-heuristic optimization algorithm, which is inspired by the hunting behavior of humpback whales. WOA uses exploration and exploitation capabilities to avoid the local optimum and accelerate the convergence in the optimization procedure of feature selection. WOA is highly compatible with other algorithms and statistical methods. WOA has the advantages of simple process and fast convergence speed, and it was found to be enough competitive with other state-of-the-art meta-heuristic methods. WOA can be used to find the best feature combination in feature selection, select the fewest features to obtain the maximum classification accuracy, and has excellent performance in solving optimization problems and applications widely.

The risk assessment of mortality plays important roles in long-term medical care for HD patients. The combined use of WOA, RSF, and typical survival analysis such as CoxPH model could provide more comprehensive risk assessment outcome. We selected three typical models in each field for comparison. The full-adjusted and stepwise selection model represents the typical statistical method, the RSF selection model represents the permutation method, and the WOA model represents the heuristic optimization algorithm. Therefore, this study aimed to apply the WOA, RSF, and typical statistical selection method in the mortality risk assessment, in purpose to identify the optimal risk characteristics combination for all-cause mortality in HD patients.

Methods

Datasets and statistical analysis

This is a retrospective cohort study. All data were obtained from Kaohsiung Chang Gung Memorial Hospital. A total of 829 patients who received regular outpatient HD therapy (three times a week) before 1 January 2009 were retrospectively included under an approved data protocol (201800595B0). All patients were tracked from 1 January 2009 to 31 December 2013. This study was conducted in accordance with the Declaration of Helsinki. We enrolled all HD patients in the beginning, and then excluded those who have not met the inclusion criteria. A total of 874 patients who received regular outpatient HD treatment (three times a week) were enrolled in the initial phase of study. Then, we selected potential risk factors associated with the risk of mortality in HD patients, and then excluded 45 patients who had loss to follow-up, and 2 patients who had missing values for blood measurements. Finally, a total of 829 patients were included in the analysis, and the study population was divided into two groups according to their mortality status. A total of 633 patients who remained alive within the study period were considered as alive cohort, and 196
patients who expired within study period were considered as died cohort. Hence, the mortality rate of HD cohort was known to be approximately 15–20%. Assume the probability of type I error is 0.05 (α), the power is 80%, the population size of current study cohort is 829, the proportion of mortality of Taiwan population is 0.20, and the proportion of mortality of current study cohort is 0.295. By using to the power estimation formula, the estimated critical Z value for given α is approximately 4.279 (Φ), which is equal to power of 1. Thus, the post hoc power of sample size is 100%, which indicates the sample size for current study was appropriate for later analyses.

The clinical factors were collected at the initial phase of study, and the comorbidity and blood measurements data were collected for each HD therapy. The survival outcome was tracked using the death registry database of our institution. The baseline characteristics and laboratory measurements in HD patients including dialysis vintage, age, sex, and diabetes mellitus (DM) status were collected.\(^\text{29}\) In addition, the baseline blood laboratory measurements, including Hb, blood urea nitrogen (BUN), Cr, K, Ca, P, intact parathyroid hormone (iPTH), ferritin, HD adequacy index, Kt/V urea (Daugirdas), urea reduction ratio (URR), and cardiothoracic ratio by chest x-ray examination were collected at the initial phase of study. The distributions of baseline characteristics and mortality-related risk in HD patients were summarized into median (interquartile range), mean (standard deviation), or frequency (percentage) according to the survival categories. The independent two-sample t-test or chi-square test was used to estimate the difference between alive and died group. Univariate CoxPH regression analysis was used to evaluate the association between all-cause mortality and individual risk factors. In addition, we have sorted out the algorithm and application comparison of mortality risk assessment studies in related HD populations proposed by previous studies. Table 1 is as follows.

### Table 1. Comparison of algorithms and application of mortality risk assessment studies in hemodialysis population.

| Study               | Algorithms                        | Application                                                                                     | Model performance                                                                 |
|---------------------|-----------------------------------|-----------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------|
| Yang et al. (2022)  | 1. CoxPH model                    | Identifying risk factors for mortality in hemodialysis patients using multiple feature selection, including stepwise, WOA, and RSF selection approaches to generate all-cause mortality risk assessment model | Optimal concordance: 1. full-adjusted-CoxPH: 0.7404 2. stepwise-CoxPH: 0.7388 3. RSF-CoxPH: 0.7406 4. WOA-CoxPH: 0.7409 Kaplan–Meier (log-rank test \(p\) value) 1. WOA-CoxPH model: Log-rank test \(p\) value < 0.001 |
| Radović et al.\(^\text{19}\) | 1. Kernel SVM algorithm           | Applied SVM method to assess the expected mortality of hemodialysis patients using nine relevant parameters provided by professional nephrologists | Mortality rate prediction is realized with accuracy up to 94.12% and up to 96.77% |
| Garcia-Montemayor et al.\(^\text{21}\) | 1. Logistic regression analysis 2. Random forest 3. AUC | Prediction of mortality of hemodialysis patients at different time points using random forest algorithm and evaluate the prediction performance using AUC | AUC 1. random forest\([ΔAUC 0.68–0.73]\) 2. logistic regression models \([ΔAUC 0.007–0.046]\) |
| Ramspek et al.\(^\text{13}\) | 1. CoxPH model                    | Predict risk of all-cause mortality in dialysis patients using typical Cox model and Kaplan–Meier methods | C-statistics ranging from 0.710 (interquartile range 0.708–0.711) to 0.752 (interquartile range 0.750–0.753) |

AUC, area under curve; CoxPH, Cox proportional hazards; RSF, random survival forest; SVM, support vector machine; WOA, whale optimization algorithm.
This algorithm simulates the humpback whale predation behavior, including exploitation and exploration. The exploitation phase uses a spiral bubble-net attacking method to find a local optimum result, and the exploration phase simulates the prey searching behavior in order to find a global optimum result. WOA feature selection could accelerate convergence to find the optimal solution for the mortality-related risk assessment in HD patients. WOA feature selection pseudocode is shown in Algorithm 1.

The algorithm first enters the stage of encircling prey, and subsequently the search in two stages, the first phase is the exploitation phase (spiral bubble-net attacking method), and the second phase is the exploration phase (search for prey). The behavior patterns in the two phases are described in detail below.

Encircling prey is the primary task, WOA assumes the current best solution or target prey, tries to advance toward the best search target and update the position, and updates the current position with the current best solution. The following equations (1)–(4) describe this behavior:

\[
\hat{D} = |\hat{C} \hat{\tilde{X}} (t) - \tilde{X} (t)|
\]  

(1)

\[
\hat{X} (\hat{t} + 1) = \hat{X} (\hat{t} + 1) - \hat{A} \hat{D}
\]

(2)

\[
\hat{A} = 2 \tilde{a} \tilde{r} - \tilde{a}
\]

(3)

\[
\hat{C} = 2 \tilde{r}
\]

(4)

where \( \hat{t} \) is the current iteration, \( \hat{X} \) is the current position vector solution, \( \hat{X}^* \) is the best solution of current position vector, \(|.|\) is the absolute value, and \( \hat{A} \) and \( \hat{C} \) are coefficient vectors. \( \hat{A} \) and \( \hat{C} \) are defined in equations (3) and (4). \( r \) is random vector in the range \([0, 1]\). The current position vector is updated according to equation (2). The values of \( \hat{A} \) affect the range of the area where the current position vector can be moved. The calculation of \( \tilde{a} \) is shown in equation (5):

Algorithm 1. Pseudo-code of whale optimization algorithm (WOA)–based feature selection.

| Input: \( N \) number of whales, \( T \) number of iteration with \( F \) size of dimension |
| Output: Optimal whale position. |
| Initialize \( X_i^j \), \( i = 1, 2, \ldots, N; j = 1, 2, \ldots, F \). |
| while \( t < T \) |
| for each whale and dimension \( X_i^j \) |
| if Sigmoid \( X_i^j \) \( \geq 0.5 \) \( Y_i^j = 1 \) |
| else \( Y_i^j = 0 \) |
| end for |
| Convert each binary individual \( Y_i \) into a feature combination. |
| Calculating Cox proportional hazard using C-index for evaluation (fitness). |
| Update \( X^* \) if there is a better fit. |
| for each whale \( X_i \) |
| Calculate and Update \( a, A, C, p \) and \( l \). |
| if \( 1 \) \( p < 0.5 \) then |
| if \( 2 \) \(|A| < 1 \) then |
| Update position by Eq. (2) |
| else \( 2 \) \(|A| \geq 1 \) |
| Choose search agent randomly \( X_{\text{rand}} \) |
| Update position by Eq. (10) |
| end if \( 2 \) |
| else if \( 1 \) \( p \geq 0.5 \) |
| Update position by Eq. (6) |
| else if \( 1 \) |
| end for |
| \( t = t + 1 \) |
| end while |
\[
\tilde{a} = 2 - t \frac{2}{MaxIter}
\]  

(5)

where \(MaxIter\) is the maximum number of allowed iterations. According to the above formula, \(\tilde{a}\) is decreased linearly from 2 to 0 over the iterations. This will reduce the movable range of shrinking encircling mechanism with iteration. \(\bar{X}(t)\) and \(X^*(t)\) can establish a spiral equation to simulate the spiral movement of humpback whales. The mathematical equation (6) is as follows:

\[
\bar{X}(t+1) = \bar{D}^t e^{bl} \cos(2\pi l) + \bar{X}^*(t)
\]

(6)

The local search capability is used in the exploitation phase. This phase is divided into two behavioral modes. The first is shrinking encircling mechanism, and the second is spiral updating procedure.

**Shrinking encircling mechanism.** This behavior pattern is achieved by decreasing the value of \(a\) in equation (3). The fluctuation range of \(A\) is decreased by linear reduction from 2 to 0 during the iteration of \(a\), then \(A\) is a random value in the interval \([-a, a]\). \(A\) in \([-1, 1]\), and a new search position can be updated between the current position and the current best position.

**Spiral updating procedure.** \(\bar{X}(t)\) and \(X^*(t)\) are established as a spiral equation to simulate the spiral movement of humpback whales to update the current best position. The spiral updating procedure are shown in equations (6)–(8):

\[
\bar{D}^t = \left| \bar{X}^*(t) - \bar{X}(t) \right| 
\]

(7)

\[
\bar{X}(t+1) = \begin{cases} 
\bar{X}^*(t) - A\bar{D} & \text{if } p < 0.5 \\
\bar{D}^t e^{bl} \cos(2\pi l) + \bar{X}^*(t) & \text{if } p \geq 0.5 
\end{cases}
\]

(8)

where \(l\) is a random number in the range \([-1, 1]\), \(b\) is a constant defining the spiral’s shape, and \(D\) is the distance between the whale \(X\) and a prey. In order to the behavior pattern of whale shrinking the enclosing mechanism is simulated, the trigger probability of the two behavior patterns (2) and (6) is set to 50% each. The mathematical equation is as follows:

When \(|\bar{A}| < 1\), we use equation (7) to update the position, where \(p\) is a random number in [0, 1]

The exploration phase is used in global search capability, and the search prey behavior pattern is used. When \(|\bar{A}| \geq 1\), it enters the exploration stage, and uses the randomly selected whale method to update the current individual’s position vector. Since \(|\bar{A}|\) must be greater than 1 at this stage, according to equation (10), the updated position will deviate from the reference whale, thereby the purpose of global search is achieved:

\[
\bar{D} = \left| \bar{C} \bar{X}_{\text{rand}} - \bar{X}(t) \right|
\]

(9)

\[
\bar{X}(t+1) = \bar{X}_{\text{rand}} - A\bar{D}
\]

(10)

\(\bar{X}_{\text{rand}}\) is a random whale position vector selected from the current population.

WOA was used to solve the feature selection problem which must be converted into binary classification.26 Therefore, the sigmoid function is added to fit the binary trait of mortality-related risk category in HD patients. The position vector of each feature is converted into binary by \(S\) function to search for the best feature combination.26 The sigmoid function is shown in equation (11):

\[
x_i^d = \frac{1}{1 + e^{-S(x_i^k(t+1))}}
\]

(11)

where \(S\) indicates the sigmoid conversion. This study uses 0.5 to make the threshold for binary. \(x_i^k(t+1)\) in equation (12) represents that the \(i\)th whale is the vector of the \(k\)th dimension at the \((t+1)\) iteration, and the value mapped by the \(S\) function is converted to an integer of 1 or 0 by
equation (12). When the dimension $k$ is 1, it means that the feature is selected. Conversely, when $k$ is 0, this feature is not used.

**RSF feature selection**

Random forest was proposed by Breiman et al. (2001), and mainly applied to classification and regression. In 2008, Ishwaran et al. applied random forest extension to survival analysis, and developed the RSF, which belongs to integrated learning. RSF is a nonparametric tree-based survival analysis algorithm based on the random forest algorithm. RSF feature selection approach evaluates variable importance (VIMP) by simultaneously considering survival time and censor status of study population. RSF uses the survival splitting rules for growing survival tree to identify highly impacted risk factor of all-cause mortality in study population. The pseudocode of RSF feature selection algorithm is shown in Algorithm 2.

In this study, RSF is used in feature selection, and mortality-related risk in HD patients was converted from continuous variables to categorical variable. The purpose is to improve the accuracy of RSF variable selection. VIMP, which was calculated to filter select variables, can be ranked and screened. A larger VIMP indicates that the variable has predictive power. Otherwise, a VIMP of zero or negative indicates that the variable’s unpredictable power needs to be considered for filtering.

**Fitness**

In each iteration, the CoxPH model was used as the objective function to evaluate and update the whale position. In survival analysis, the CoxPH model is a commonly used statistic that uses medical research patients and univariate or multivariate variables to predict associations between survival times. The CoxPH model can be used to assess how specified factors influence the rate of a particular event happening (e.g. infection, death).

The CoxPH model was used to establish the survival objects of HD patients as dependent variables, and other HD influencing factors were used as independent variables to assess whether it was related to the mortality of HD patients, and the hazard ratio (HR) value was calculated for the death risk determination. The model can be written as follows:

\[
\ln h(t) = \ln h_0(t) + b_1x_1 + \cdots + b_px_p \quad (13)
\]

or

\[
\frac{\ln h(t)}{h_0(t)} = b_1x_1 + \cdots + b_px_p \quad (14)
\]

where \( h(t) \) is the risk of time \( (t) \); \( x_1, x_2 \ldots x_p \) are explanatory variables. When HR value is less than 1, more than 1, and equal to 1, the risk of death is reduced, increased, and invalidated, respectively.

Concordance index is used to evaluate the predictive ability of the model. It was first proposed by
Frank E Harrell Jr, a professor of biostatistics at Vanderbilt University in 1996. It is mainly used to calculate the discrimination between the predicted value and the true value of the CoxPH model in survival analysis. It is also called Harrell’s concordance index. The C-index calculation method randomly pairs study objects. For a pair of patients, if A’s actual survival time is longer than B, and the predicted result is the same, it is called concordance. The calculation formula of the concordance index \( c \) is as follows:

\[
c = \frac{K}{M}
\]  

where \( M \) is the number of remaining matches.

**Optimal CoxPH regression model**

This study aimed to assess of mortality-related risk in HD patients using machine learning approach to select features. The typical CoxPH model selection approach including all associated factors in known as full-adjusted-CoxPH, and the stepwise-CoxPH model includes the associated factors which have met the critical \( p \) value (\(<0.2\)).

In addition to the typical statistic selection approaches, RSF and WOA methods were used to assess the optimal risk factors combination for all-cause mortality in CoxPH regression model. RSF model uses VIMP to identify associated risk factors, and WOA model uses local and global optimum search to identify associated risk factors. Taken together, this study performed full-adjusted-CoxPH, stepwise-CoxPH, RSF-CoxPH, and WOA-CoxPH model for the all-cause mortality risk assessment in HD patients, and compared the model performance using concordance index. Then, the survival interval and all-cause mortality status of HD patients were used to create a survival object, and concordance index was used as the objective function. Model comparison between four feature selection CoxPH model was performed to clarify the efficiency of proposed feature selection approaches for mortality risk assessment in HD patients. Harrell’s concordance index value was used to compare the model performance; the higher concordance index represents better combination solution. Finally, the identified associated risk factors will be assessed for relevance to all-cause mortality in HD patients.

The HR and 95% confidence interval (CI) were computed. The significant associated risk factors identified by optimal CoxPH model were illustrated using Kaplan–Meier survival curves, and the survival difference between subgroups was tested using log-rank test. Moreover, the significant factors in optimal model were further used to generate an integrated risk score for mortality risk assessment of study population. Receiver operating characteristics (ROC) analysis was performed to further dichotomize the study population into low- and high-risk subgroups, and the survival difference between integrated risk subgroup was also tested using log-rank test. All \( p \) values were two-sided, and a \( p \) value less than 0.5 was considered statistically significant. All analyses were performed by R software (R Development Core Team 2020, version 4.0.2).

**Results**

**Baseline characteristics and laboratory measurements**

Table 2 showed the distribution and comparison of clinicopathological characteristics between alive and died groups. A total of 633 alive patients and 196 dead patients were analyzed. The died group had significant higher proportion in age \( \geq 65 \) (died *versus* alive: 57.65% *versus* 28.28%, \( p < 0.001 \)) and DM (38.27% *versus* 22.27%, \( p < 0.001 \)). In laboratory measurements, the died group had significant higher proportion in hemoglobin (Hb) <10.64g/dl (59.18% *versus* 46.76%, \( p = 0.003 \)), albumin <3.87g/dl (62.24% *versus* 36.49%, \( p < 0.001 \)), creatinine (Cr) <1.70mg/dl (59.18% *versus* 47.24%, \( p = 0.004 \)), Kt/V urea <1.7 (65.31% *versus* 52.61%, \( p = 0.002 \)), and cardiothoracic ratio >0.5 (72.96% *versus* 46.45%, \( p < 0.001 \)) compared with the alive group.

**Individual risk factors of all-cause mortality**

Table 3 showed the univariate CoxPH regression analysis results for all-cause mortality in HD patients. The univariate analysis results showed patients with age \( \geq 65 \) years (HR = 2.91, 95% CI = 2.19–3.87, \( p < 0.001 \)), DM (HR = 1.98, 95% CI = 1.49–2.65, \( p < 0.001 \)), and cardiothoracic ratio >0.5 (HR = 2.75, 95% CI = 2.01–3.77, \( p < 0.001 \)) were more likely to achieve higher mortality risk. In addition, HD patients
Table 2. Baseline characteristics of mortality categories \( (n=829) \) with two categories.

| Characteristics | Total \((n=829)\) | Alive \((n=633)\) | Died \((n=196)\) | \(p\) |
|----------------|-----------------|-----------------|----------------|------|
| Dialysis vintage (years) | 0.213 |
| \(\geq 5.60\) | 415 [50.06] | 325 [51.34] | 90 [45.92] |
| \(< 5.60\) | 414 [49.94] | 308 [48.66] | 106 [54.08] |
| Age (years) | \(< 0.001\) |
| \(\geq 65\) | 292 [35.22] | 179 [28.28] | 113 [57.65] |
| \(< 65\) | 537 [64.78] | 454 [71.72] | 83 [42.35] |
| Sex | 0.819 |
| Male | 376 [45.36] | 289 [45.66] | 87 [44.39] |
| Female | 453 [54.64] | 344 [45.34] | 109 [55.61] |
| DM | \(< 0.001\) |
| 216 [26.06] | 141 [22.27] | 75 [38.27] |
| Laboratory measurements | | | | |
| Hb (g/dl) | 0.003 |
| \(\geq 10.64\) | 417 [50.30] | 337 [53.24] | 80 [40.82] |
| \(< 10.64\) | 412 [49.70] | 296 [46.76] | 116 [59.18] |
| Albumin (g/dl) | \(< 0.001\) |
| \(\geq 3.87\) | 476 [57.42] | 402 [63.51] | 74 [37.76] |
| \(< 3.87\) | 353 [42.58] | 231 [36.49] | 122 [62.24] |
| BUN (mg/dl) | 0.180 |
| \(\geq 68.91\) | 409 [49.33] | 321 [50.71] | 88 [44.90] |
| \(< 68.91\) | 420 [50.66] | 312 [49.29] | 108 [55.10] |
| Cr (mg/dl) | \(< 0.001\) |
| \(\geq 10.52\) | 403 [48.61] | 340 [53.71] | 63 [32.14] |
| \(< 10.52\) | 426 [51.39] | 293 [46.29] | 133 [67.86] |
| K (meq/l) | 0.004 |
| \(\geq 4.95\) | 414 [49.94] | 334 [52.76] | 80 [40.82] |
| \(< 4.9\) | 415 [50.06] | 299 [47.24] | 116 [59.18] |
| Ca (mg/dl) | 0.064 |
| \(\geq 9.22\) | 384 [46.32] | 304 [48.18] | 79 [40.31] |
| \(< 9.22\) | 445 [53.68] | 328 [51.82] | 117 [59.69] |

(Continued)
with Hb $\geq 10.64$ (HR = 0.62, 95% CI = 0.47–0.83, $p = 0.001$), albumin $\geq 3.87$ g/dl (HR = 0.39, 95% CI = 0.29–0.52, $p < 0.001$), Cr $\geq 10.52$ mg/dl (HR = 0.44, 95% CI = 0.33–0.60, $p < 0.001$), K $\geq 4.95$ meq/l (HR = 0.66, 95% CI = 0.50–0.88, $p = 0.004$), Ca $\geq 9.22$ mg/dl (HR = 0.74, 95% CI = 0.56–0.98, $p = 0.037$), and Kt/V urea $\geq 1.70$ (HR = 0.63, 95% CI = 0.47–0.84, $p = 0.002$) were more likely to achieve lower mortality risk.

**Risk assessment model of all-cause mortality**

The analysis results of full-adjusted-CoxPH, stepwise-CoxPH, RSF-CoxPH, and WOA-CoxPH model for all-cause mortality in HD patients are summarized in Table 4. The full-adjusted-CoxPH model showed patients with age $\geq 65$ years (HR = 1.19, 95% CI = 1.46–2.68, $p < 0.001$), DM (HR = 1.64, 95% CI = 1.19–2.27, $p = 0.002$), Hb $< 10.64$ (HR = 0.66, 95% CI = 0.49–0.89, $p = 0.006$), albumin $< 3.87$ g/dl (HR = 0.62, 95% CI = 0.45–0.85, $p = 0.003$), Cr $< 10.52$ mg/dl (HR = 0.61, 95% CI = 0.43–0.88, $p = 0.008$), K $< 4.95$ meq/l (HR = 0.73, 95% CI = 0.55–0.99, $p = 0.041$), Kt/V urea $< 0.71$ (HR = 0.61, 95% CI = 0.44–0.86, $p = 0.005$), and cardiothoracic ratio $\geq 0.5$ (HR = 1.98, 95% CI = 1.42–2.75, $p < 0.001$) were more likely to associate with the increasing risk of all-cause mortality in HD patients.

### Table 2. (Continued)

| Characteristics       | Total (n = 829) | Alive (n = 633) | Died (n = 196) | $p$ |
|-----------------------|----------------|----------------|---------------|-----|
| Phosphate [mg/dl]     |                |                |               | 0.588 |
| $\geq 4.84$           | 401 [48.37]    | 310 [48.97]    | 91 [45.43]    |
| $<4.84$               | 428 [51.63]    | 323 [51.03]    | 105 [53.57]   |
| iPTH (pg/ml)          |                |                |               | 0.919 |
| $\geq 205.60$         | 415 [50.06]    | 318 [50.24]    | 97 [49.49]    |
| $<205.60$             | 414 [49.94]    | 315 [49.76]    | 99 [50.51]    |
| Ferritin [ng/ml]      |                |                |               | 0.171 |
| $\geq 412.70$         | 415 [50.06]    | 308 [48.66]    | 107 [54.59]   |
| $<412.70$             | 414 [49.94]    | 325 [51.34]    | 89 [45.41]    |
| Kt/V urea             |                |                |               | 0.002 |
| $\geq 1.70$           | 368 [44.39]    | 300 [47.39]    | 68 [34.69]    |
| $<1.70$               | 461 [55.61]    | 333 [52.61]    | 128 [65.31]   |
| URR                   |                |                |               | 1.000 |
| $\geq 0.65$           | 786 [94.81]    | 600 [94.79]    | 186 [94.90]   |
| $<0.65$               | 43 [5.19]      | 33 [5.21]      | 10 [5.10]     |
| Cardiothoracic ratio  |                |                |               | $<0.001$ |
| $\geq 0.50$           | 437 [52.71]    | 294 [46.45]    | 143 [72.96]   |
| $<0.50$               | 392 [47.29]    | 339 [53.55]    | 53 [27.04]    |

BUN, blood urine nitrogen; DM, diabetes mellitus; iPTH, intact parathyroid hormone; TSA, time-averaged serum albumin; URR, urea reduction ratio.
The $p$ value is estimated using independent two-sampled t-test.
$p$ values less than 0.05 were considered statistically significant.
In the stepwise-CoxPH model, the risk factors that met critical $p$ value $< 0.2$ were included. The stepwise-CoxPH model indicates patients with age $\geq 65$ years (HR = 1.92, 95% CI = 1.43–2.59, $p < 0.001$), DM (HR = 1.55, 95% CI = 1.15–2.10, $p = 0.004$), Hb $< 10.64$ g/dl (HR = 0.65, 95% CI = 0.49–0.87, $p = 0.004$), albumin $< 3.87$ g/dl (HR = 0.62, 95% CI = 0.46–0.84, $p = 0.002$), Cr $< 10.52$ mg/dl (HR = 0.64, 95% CI = 0.46–0.88, $p = 0.007$), K $< 4.95$ meq/l (HR = 0.75, 95% CI = 0.56–1.00, $p = 0.048$), Kt/V urea $< 0.71$ (HR = 0.61, 95% CI = 0.45–0.84, $p = 0.002$), and cardiothoracic ratio $\geq 0.5$ (HR = 1.99, 95% CI = 1.44–2.75, $p < 0.001$) were more likely to increased all-cause mortality risk.

The RSF-CoxPH model included dialysis vintage, age, sex, DM, Hb, albumin, BUN, creatinine, K, Ca, phosphate, Kt/V urea, and cardiothoracic ratio. Patients characterized with age $\geq 65$ years (HR = 1.97, 95% CI = 1.46–2.66, $p < 0.001$), with DM (HR = 1.63, 95% CI = 1.18–2.25, $p = 0.003$), Hb $< 10.64$ g/dl (HR = 0.66, 95% CI = 0.49–0.89, $p = 0.006$), albumin $< 3.87$ g/dl (HR = 0.62, 95% CI = 0.45–0.85, $p = 0.003$), Cr $< 10.52$ mg/dl (HR = 0.62, 95% CI = 0.46–0.88,
Table 4. Multivariate Cox proportional hazard for all-cause mortality based on different feature selection approaches.

| Characteristics                  | Comparison         | Full-adjusted | Stepwise (p < 0.2) | RSF       | WOA       |
|----------------------------------|--------------------|---------------|--------------------|-----------|-----------|
|                                  |                    | HR | 95% CI | p            | HR | 95% CI | p            | HR | 95% CI | p            | HR | 95% CI | p                      |
| Dialysis vintage (years)         | ≥5.60 versus < 5.60| 1.19 | 0.86–1.63 | 0.289 | 1.19 | 0.87–1.63 | 0.278 | 1.20 | 0.88–1.64 | 0.256 |
| Age                              | ≥65 versus < 65    | 1.98 | 1.46–2.68 | <0.001 | 1.92 | 1.43–2.59 | <0.001 | 1.97 | 1.46–2.66 | <0.001 | 1.99 | 1.47–2.69 | <0.001 |
| Sex                              | Female versus male | 0.95 | 0.68–1.33 | 0.779 | –     | –     | –     | 0.96 | 0.69–1.35 | 0.829 | –     | –                      |
| DM                               | Yes versus no      | 1.64 | 1.19–2.27 | 0.002 | 1.55 | 1.15–2.10 | 0.004 | 1.63 | 1.18–2.25 | 0.003 | 1.63 | 1.18–2.24 | 0.003 |
| Laboratory measurements          |                    |               |                   |          |          |          |          |          |                   |          |          |                       |
| Hb (g/dl)                        | ≥10.64 versus < 10.64| 0.66 | 0.49–0.89 | 0.006 | 0.65 | 0.49–0.87 | 0.004 | 0.66 | 0.49–0.89 | 0.006 | 0.65 | 0.48–0.88 | 0.005 |
| Albumin (g/dl)                   | ≥3.87 versus < 3.87| 0.62 | 0.45–0.85 | 0.003 | 0.62 | 0.46–0.84 | 0.002 | 0.62 | 0.45–0.85 | 0.003 | 0.62 | 0.45–0.85 | 0.003 |
| BUN (mg/dl)                      | ≥68.91 versus < 68.91| 1.12 | 0.82–1.53 | 0.460 | –     | –     | –     | 1.12 | 0.82–1.52 | 0.482 | 1.06 | 0.78–1.43 | 0.730 |
| Cr (mg/dl)                       | ≥10.52 versus < 10.52| 0.61 | 0.43–0.88 | 0.008 | 0.64 | 0.46–0.88 | 0.007 | 0.62 | 0.43–0.88 | 0.008 | 0.62 | 0.44–0.87 | 0.006 |
| K (meq/l)                        | ≥4.95 versus < 4.95| 0.73 | 0.55–0.99 | 0.041 | 0.75 | 0.56–1.00 | 0.048 | 0.74 | 0.55–0.99 | 0.042 | –     | –                      |
| Ca (mg/dl)                       | ≥9.22 versus < 9.22| 0.90 | 0.65–1.23 | 0.500 | –     | –     | –     | 0.92 | 0.67–1.25 | 0.580 | 0.90 | 0.66–1.23 | 0.508 |
| Phosphate (mg/dl)                | ≥4.84 versus < 4.84| 1.00 | 0.73–1.37 | 0.993 | –     | –     | –     | 1.02 | 0.75–1.39 | 0.905 | 1.00 | 0.73–1.35 | 0.978 |
| iPTH (pg/ml)                     | ≥205.60 versus < 205.60| 1.12 | 0.83–1.51 | 0.452 | –     | –     | –     | –     | –               | –     | –     | –                      |
| Ferritin (ng/ml)                 | ≥412.70 versus < 412.70| 0.99 | 0.74–1.32 | 0.940 | –     | –     | –     | –     | 0.99 | 0.74–1.32 | 0.946 | –     | –                      |
| Kt/V urea                        | ≥1.70 versus < 1.70| 0.61 | 0.44–0.86 | 0.005 | 0.61 | 0.45–0.84 | 0.002 | 0.62 | 0.44–0.86 | 0.005 | 0.60 | 0.44–0.83 | 0.002 |
| URR                              | ≥0.65 versus < 0.65| 1.10 | 0.57–2.10 | 0.784 | –     | –     | –     | –     | 1.10 | 0.57–2.11 | 0.772 | –     | –                      |
| Cardiothoracic ratio             | ≥0.50 versus < 0.50| 1.98 | 1.42–2.75 | <0.001 | 1.99 | 1.44–2.75 | <0.001 | 1.99 | 1.43–2.76 | <0.001 | 1.94 | 1.40–2.70 | <0.001 |
| Optimal concordance              |                    | 0.7404 | 0.7388 | 0.7406 | 0.7409 |

BUN, blood urea nitrogen; CI, confidence interval; DM, diabetes mellitus; HR, hazard ratio; iPTH, intact parathyroid hormone; RSF, Random survival forest; URR, urea reduction ratio; WOA, whale optimization algorithm.
p values less than 0.5 were considered statistically significant.
The WOA-CoxPH model included dialysis vintage, age, DM, Hb, albumin, BUN, Cr, Ca, phosphate, ferritin, Kt/V urea, URR, and cardiothoracic ratio. Patients characterized with age ≥ 65 years (HR = 1.99, 95% CI = 1.47–2.69, p < 0.001), DM (HR = 1.63, 95% CI = 1.18–2.24, p = 0.003), Hb < 10.64 g/dl (HR = 0.65, 95% CI = 0.48–0.88, p = 0.005), albumin < 3.87 g/dl (HR = 0.62, 95% CI = 0.45–0.85, p = 0.003), Cr < 10.52 mg/dl (HR = 0.62, 95% CI = 0.44–0.87, p = 0.006), Kt/V urea < 0.71 (HR = 0.60, 95% CI = 0.44–0.83, p = 0.002), and cardiothoracic ratio ≥ 0.5 (HR = 1.94, 95% CI = 1.40–2.70, p < 0.001) were more likely to increased all-cause mortality risk.

RSF-CoxPH and WOA-CoxPH model obtained similar analysis results. Both models selected 13 out of the 16 factors but differ in three factors. The concordance index for full-adjusted-CoxPH, stepwise-CoxPH, RSF-CoxPH, and WOA-CoxPH model were 0.7404, 0.7388, 0.7406, and 0.7409, respectively. The comparison results showed that WOA-CoxPH model obtained the highest C-index among all models, which indicates that the WOA model could achieve better concordance in all-cause mortality risk estimation for HD patients. Furthermore, the Kaplan–Meier curve of significant risk factors identified in both RSF and WOA model were illustrated in Figure 1. Figure 1(a) presented the overall survival probability of all patients, the overall survival rate of all patients was 76.9% (95% CI = 74.0–79.8). Figure 1(b)–(i) illustrated overall survival probability analysis results of the eight significant parameters obtained from the WOA-CoxPH model including age (log-rank test p < 0.001), DM (log-rank test p < 0.001), Hb (log-rank test p = 0.001), albumin (log-rank test p < 0.001), Ca (log-rank test p < 0.001), K (log-rank test p = 0.004), Kt/V (log-rank test p = 0.016), and cardiothoracic ratio (log-rank test p < 0.001), which have also showed significant survival difference between low- and high-risk characteristics in single-factor analysis. The red solid line indicates the high-risk characteristics, and the blue solid line indicates the low-risk characteristics. The high-risk characteristics of eight selected parameters were scored 1, and the low-risk characteristics were scored 0. An integrated risk score for overall survival was generated based on the summation of eight selected parameters derived using WOA-selection model. An optimal cutoff point for the integrated risk score was determined using ROC analysis according to the mortality status. Thus, the patients were dichotomized into low- and high-risk subgroup by the optimal cutoff points of seven. Figure 1(j) showed that the patients who obtained seven or more risk characteristics of eight selected parameters will achieve significant worse overall survival probability compared with those who obtained less or equal to six risk characteristics (log-rank test p < 0.001).

Overall, the integrated low- and high-risk subgroup showed greater discrepancy compared with each single risk factor selected by WOA-CoxPH model. In summary, patients who obtained seven or more risk characteristics of eight selected parameters could have potentially increased risk of all-cause mortality in HD population.

**Discussion**

In this study, we identified optimal risk factors combination of all-cause mortality in HD patients using WOA-CoxPH model. The identified risk factors were age, diabetes, hemoglobin, serum albumin, serum creatinine, serum potassium, Kt/V urea, and cardiothoracic ratio. The study results are consistent with previous studies in HD patients. Based on clinical practice point, aging is characterized by the progressive decline in function of organs. Consequently, aging contributes to the disease occurrence and disease-related mortality. In our previous investigations, we found that elder age was associated with decline in physical functional performance and cognitive function in HD patients. We believe that these disability circumstances predispose HD patients to death risk. It is well recognized that protein-energy-wasting is strongly associated with mortality in dialysis patients. Serum albumin, creatinine, and hemoglobin are commonly applied to be as surrogates for nutritional status. Our previous observational studies have validated that these clinical factors are associated with poor quality of life and death risk in dialysis patients. Diabetes has become the major etiology of chronic kidney disease. This disease
Figure 1. Kaplan–Meier curve for all-cause mortality in (a) all patients and difference subgroup of (b) age, (c) DM status, (d) hemoglobin, (e) albumin, (f) creatinine, (g) potassium, (h) Kt/V, (i) cardiothoracic ratio group, and (j) integrated risk subgroup derived using WOA-CoxPH model.
involves endothelial dysfunction and eventually organ failure. The association between diabetes and risk of mortality in HD patients also has evidenced via systematic review and meta-analysis. Chronic kidney disease patients commonly have elevated serum potassium levels due to reduced renal clearance. Cardiac conductive activity is influenced by serum potassium levels. Elevated serum potassium levels could increase cardiac arrhythmia events in dialysis patients. Consequently, death risk would increase in the event of serum potassium increment especially in HD patients. Considering HD adequacy, Kt/V-urea calculation is commonly used. This calculation reflects the clearance of small molecular weight uremic toxins. This index has been applied as standard surrogate to evaluate dialysis adequacy either on HD or on peritoneal dialysis for years. Moreover, this adequacy index has been reported with survival in HD patients. Taken together, selected risk factors for death in HD patients via WOA and RSF feature are reasonable via clinical practice point and literature review.

The study results indicate WOA-CoxPH model could perform as better survival model compared with typical statistical survival selection model. WOA used a group-based approach combined with exploration and exploitation procedures to accelerate the optimal search process. WOA algorithm could maintain an appropriate balance between exploration and exploitation by considering both local and global search for optimal combination. The mentioned search strategy made WOA more competitive than other methods, especially in terms of complex and interactive feature issues. Moreover, RSF-CoxPH also provides a similar performance with WOA-CoxPH compared with typical CoxPH models. RSF calculates the VIMP of each variable and ranked the value in purpose to filter the high-impact variables for interest outcome, which could provide lower prediction errors. Both WOA-CoxPH and RSF-CoxPH showed the addition machine learning algorithm in feature selection procedure could improve the performance of CoxPH model.

Several limitations of current study should be noted. First, the inclusion of certain covariates was limited due to the retrospective nature of current study. Second, this is a single institution study. Third, the study findings are restricted in HD population. Although the mentioned limitations could limit the generalisability of current findings, this study still provides an optimal feature selection approach by using RSF and WOA in survival model, which could provide better concordance model performance compared with typical feature selection methods in survival model estimation.

**Conclusion**

The all-cause mortality is commonly considered as primary endpoints for patients undergoing long-term HD in the clinical studies. Previous studies indicated all-cause mortality in HD patient is associated with multiple clinical factors, including comorbidity, medications, nutrition status, and others. This study performed the all-cause mortality risk assessment model of HD patients based on machine learning feature selection procedure. Compared with the typical statistical selection model, RSF-CoxPH and WOA-CoxPH model showed better model performance. Therefore, RSF-CoxPH and WOA-CoxPH model identified risk factors combination might contribute to the more precise risk assessment of all-cause mortality in HD patients, and those identified risk factors could be considered as an important monitoring index in further management for HD patients in order to provide better survival outcome in maintenance HD population.

**Declarations**

*Ethics approval and consent to participate*

The study was employed in accordance with the Declaration of Helsinki and approved by the Chuang Gung Medical Foundation Institutional Review Board (201800595B0). Written informed consent was obtained from all participants.

*Consent for publication*

Not applicable.

*Author contributions*

**Cheng-Hong Yang:** Conceptualization; Funding acquisition; Methodology; Project administration; Supervision; Writing – review & editing.

**Yin-Syuan Chen:** Conceptualization; Data curation; Software.
Sin-Hua Moi: Formal analysis; Validation; Writing – original draft.

Jin-Bor Chen: Data curation; Formal analysis; Investigation; Resources; Visualization.

Lin Wang: Conceptualization; Formal analysis; Validation; Writing – review & editing.

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