Pretransplant Prediction of Posttransplant Survival for Liver Recipients with Benign End-Stage Liver Diseases: A Nonlinear Model

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Abstract

Background: The scarcity of grafts available necessitates a system that considers expected posttransplant survival, in addition to pretransplant mortality as estimated by the MELD. So far, however, conventional linear techniques have failed to achieve sufficient accuracy in posttransplant outcome prediction. In this study, we aim to develop a pretransplant predictive model for liver recipients’ survival with benign end-stage liver diseases (BESLD) by a nonlinear method based on pretransplant characteristics, and compare its performance with a BESLD-specific prognostic model (MELD) and a general-illness severity model (the sequential organ failure assessment score, or SOFA score).

Methodology/Principal Findings: With retrospectively collected data on 360 recipients receiving deceased-donor transplantation for BESLD between February 1999 and August 2009 in the west China hospital of Sichuan university, we developed a multi-layer perceptron (MLP) network to predict one-year and two-year survival probability after transplantation. The performances of the MLP, SOFA, and MELD were assessed by measuring both calibration ability and discriminative power, with Hosmer-Lemeshow test and receiver operating characteristic analysis, respectively. By the forward stepwise selection, donor age and BMI; serum concentration of HB, Crea, ALB, TB, ALT, INR, Na+; presence of pretransplant diabetes; dialysis prior to transplantation, and microbiologically proven sepsis were identified to be the optimal input features. The MLP, employing 18 input neurons and 12 hidden neurons, yielded high predictive accuracy, with c-statistic of 0.91 (P<0.001) in one-year and 0.88 (P<0.001) in two-year prediction. The performances of SOFA and MELD were fairly poor in prognostic assessment, with c-statistics of 0.70 and 0.66, respectively, in one-year prediction, and 0.67 and 0.65 in two-year prediction.

Conclusions/Significance: The posttransplant prognosis is a multidimensional nonlinear problem, and the MLP can achieve significantly high accuracy than SOFA and MELD scores in posttransplant survival prediction. The pattern recognition methodologies like MLP hold promise for solving posttransplant outcome prediction.

Introduction

Orthotopic Liver transplantation (OLT) has become an established treatment approach for patients with benign end-stage liver diseases (BESLD, i.e. non-neoplastic diseases), but the growing scarcity of grafts compared to numbers of waiting patients, coupled with the high cost of this procedure, make it imperative to make difficult decisions about how to distribute such scarce organs [1–3], and highlight the need to identify patients likely to have relatively good outcomes after transplantation [4–6]. This need is particularly acute in the Asia-Pacific region, where the carrier rate of hepatitis B virus (HBV) is estimated at 20%–30% [7,8] and large numbers of BESLD patients with HBV-related cirrhosis and severe hepatitis B need OLT. Under such circumstances, the ideal allocation system would allocate livers to candidates who are most likely to die without a transplant, but who also have a high probability of survival after OLT. The balanced application of a model for liver transplant outcome estimation, in concert with a model for end-stage liver disease (MELD) estimating disease severity, would improve transplant outcomes and maximize patients’ benefit from OLT [9].

In order to incorporate likely posttransplant prognosis into decisions about grafts allocation, and to facilitate informed decision-making by potential transplant recipients and their
It is necessary to accurately assess the likelihood of posttransplant survival based on information that is available before transplantation. Although there have been some attempts to develop a model that meets this requirement, most lacked sufficient discriminating accuracy or simply stratified the prognostic risk [4,6,9,11–14]. One major reason for this is inappropriate choice of modeling method [13]. Survival prognosis is a complex nonlinear relationship affected by many interactive factors, especially for a complicated organ transplantation procedure; however, most current models were developed by linear methods, such as multiple regression.

Artificial neural network (ANN) is a computer-based nonlinear data mining mode that can recognize relationships between a series of independent variables and the corresponding dependent variable. It is more successful than traditional linear methods when the prognostic effect of a variable is influenced by other variables in a complex multidimensional nonlinear function, or when the importance of a given prognostic variable is expressed as a complex unknown function of the value of the variable [15,16]. Thus, ANN is particularly suited to modeling complex multidimensional patterns [17,18], and has had remarkable success in many medical problems that are too complicated for linear models [15,19,20]. To date, there have been a few attempts to use ANN for outcome prediction after organ transplantation [17,21,22], but no reliable ANN model had been developed specifically for BESLD recipients.

We investigated the feasibility of using multi-layer perceptron (MLP), arguably one of the most efficient ANN for prognostic research [22,23], to develop a prognostic model to predict individualized survival probability after deceased donor OLT in recipients with BESLD, employing typically available, objective preoperative characteristics. Furthermore, we evaluated and compared the predictive accuracy of this MLP network with a BESLD-specific prognostic model (MELD) and a general-illness severity prognostic model (the sequential organ failure assessment score, or SOFA score).

## Methods

### Data source

Between February 1999 and August 2009, 386 adults with BESLD received deceased-donor (either no heartbeat or brain dead) liver transplants at the 4300-bed West China Hospital of Sichuan University. We excluded 15 recipients with combined organ transplants or partial organs and 11 recipients with incomplete follow-up records. The remaining 360 transplants were involved in this study and followed up by August 31, 2010. Maintenance immunosuppression initially consisted of a triple-drug regimen that included either tacrolimus or cyclosporine, mycophenolate, and prednisone, and that recipients were eventually weaned to dual or single agent.

We extracted demographic characteristics of donors and recipients, pretransplant clinical records (Tables 1 and Table S1), and recipients’ follow-up information form the electronic database of the liver transplantation center at West China Hospital. Surgical and some donor factors were not included in the model development, since they could not have been known when recipients decided whether to undergo OLT and were ranked on the waiting list. All included data were taken from the most recent examinations prior to transplantation, since they reflected the current medical condition of the candidate at time of transplantation.

All organ donations recorded in the electronic database were contributed voluntarily, and no grafts were obtained from executed prisoners or other institutionalized persons. All of the donors or their families had provided written, valid informed consent for donation before the organs were procured. Each liver donation and transplantation in our center was approved by the Medical Ethics Committee of West China Hospital, Sichuan University, and the study protocol was carried out in accordance with the Declaration of Helsinki.

### Dataset division

A data-splitting approach was used in this study. The recipients were randomly divided into a modeling set (80% of the total sample, 290 recipients) used to construct the MLP network, and a validation set (20% of the total sample, 70 recipients) used to assess the models’ predictive accuracy; the validation samples would not be involved in the model development. The modeling set was randomly re-divided into a general training set (80% of the modeling set, 232 recipients) and a cross validation set (20% of the modeling set, 58 recipients) to perform the internal cross validation in MLP training.

### Statistical analysis

Continuous variables were reported as mean ± standard deviation and compared using Student’s t test; categorical variables were reported as numbers and percentages, modeled as dummy variables, and compared using the chi-square test. A value of $P<0.05$ was considered significant in all the analyses. All analyses, except the MLP development, were carried out using SAS 8.0.

### MELD and SOFA scores calculation

The BESLD-specific illness severity was evaluated by the MELD and MELD-Na$^+$ scores, which were calculated according to the following formulas: $\text{MELD} = 3.78 \times \log_e \text{INR} + 11.20 \times \log_e \text{INR} + 9.57 \times \log_e \text{Crea} + (\text{mg/dl}) + 6.4$ [24]. MELD-Na$^+$ = MELD - Na$^+$ – (0.025 × MELD × (140 – Na$^+$)) + 140 [25].

The general illness severity was assessed by the SOFA score, which is composed of scores from six organ systems (respiratory, coagulation, liver, cardiovascular, renal, and neurological) graded from 0 to 4 points according to normal function or the degree of dysfunction [26] (Table 2).

### The MLP network development

An MLP consists of a densely interconnected set of units. In this study, we developed a three-layer network which not only can approximate any reasonable function to any degree of required precision as long as the hidden layer is large enough, but also has an advantage in computing speed compared to multiple hidden layer networks [27]. The concept of a neuron is a high-level abstraction that encompasses both certain values and a set of operations that are performed on those values, and neurons are tied together with weighted connections. The MLP was developed using STATISTICA 8.0.

### Determination of input neurons

We performed the forwards stepwise selection algorithm to screen and identify the input feature variables from the candidate variables (Table 1 and Table S1), in which quantitative variables were assigned one-to-one to the neurons and each sub-category of every categorical variable was defined as an input neuron. All input quantitative variables were scaled linearly between 0 and 1.0 using the following transformation formula, where $\min\{x_{ij}\}$ and $\max\{x_{ij}\}$ were the minimum and maximum values of the variable. The
input categorical variables were entered as dummy variables.

\[
y_j = \frac{x_j - \min_{0 \leq i \leq n} \{x_i\}}{\max_{0 \leq i \leq n} \{x_i\} - \min_{0 \leq i \leq n} \{x_i\}}
\]

**Determination of output neuron.** The probability of survival at posttransplant one year and two years was entered as continuous output on the interval 0–1, in which 0 represents death and 1 represents survival, so the MLP output values represent the probability of posttransplant recipient survival. Survival was chosen as the outcome endpoint because it is the most reliable and unbiased variable in the prognostic research [28].

**Determination of hidden neurons and network transfer function.** The hidden neurons calculate the weighted sum of inputs from the input neurons and produce the output result through an activation algorithm (i.e. transfer function). The weights are adjusted based on the training data in order to minimize the error estimate function [29]. Therefore, the approximate number of hidden neurons and the corresponding transfer function are closely related to the predictive accuracy of the network. In this study, the number of hidden neurons varied from two to 35, and the alternative transfer functions included identity, logistic, tanh, exponential, gaussian and softmax. We applied the enumerative combinatorial method to exhaustively evaluate all possible combinations of hidden neuron numbers and transfer functions, then identified the combination with the best predictive accuracy.

**Cross-validation.** Experiments have verified that the predictive accuracy of an MLP initially increases with the number of training iterations, but starts deteriorating after a critical point, because the network becomes over-fitted to the training data. In this study, the number of hidden neurons varied from two to 35, and the alternative transfer functions included identity, logistic, tanh, exponential, gaussian and softmax. We applied the enumerative combinatorial method to exhaustively evaluate all possible combinations of hidden neuron numbers and transfer functions, then identified the combination with the best predictive accuracy.

**Table 1.** Baseline quantitative characteristics of the training set and validation set.

| Variables                          | Training set (N = 290) | Validation set (N = 70) | P-Value |
|------------------------------------|------------------------|-------------------------|---------|
| **Donor characteristics**          |                        |                         |         |
| D-Age (yr)                         | 38.12 ± 12.33          | 37.88 ± 11.99           | 0.883   |
| D-BMI                              | 23.95 ± 5.84           | 24.89 ± 5.16            | 0.218   |
| cold ischemia times (hour)*        | 8.65 ± 4.16            | 8.92 ± 4.03             | 0.624   |
| warm ischemia times (min)*         | 9.67 ± 2.78            | 9.26 ± 2.65             | 0.265   |
| **Recipient characteristics**      |                        |                         |         |
| Age (yr)                           | 45.25 ± 10.32          | 44.88 ± 10.18           | 0.787   |
| BMI                                | 20.06 ± 4.41           | 20.18 ± 4.35            | 0.838   |
| HB (g/dl)                          | 9.58 ± 5.36            | 9.03 ± 5.11             | 0.437   |
| WBC (×10⁶)                         | 6.36 ± 4.66            | 7.05 ± 4.69             | 0.268   |
| PLT (×10⁹)                         | 83.86 ± 85.58          | 79.99 ± 81.56           | 0.732   |
| ALB (g/dl)                         | 3.06 ± 0.65            | 2.89 ± 0.55             | **0.044** |
| TB (mg/dl)                         | 13.81 ± 14.66          | 12.39 ± 13.46           | 0.461   |
| ALK (u/l)                          | 161.46 ± 186.61        | 172.11 ± 185.32         | 0.668   |
| GGT (u/l)                          | 102.87 ± 184.35        | 97.56 ± 173.23          | 0.827   |
| AST (u/l)                          | 124.62 ± 176.05        | 149.76 ± 175.69         | 0.284   |
| ALT (u/l)                          | 113.06 ± 191.98        | 143.25 ± 183.21         | 0.234   |
| BUN (mmol/l)                       | 20.55 ± 16.86          | 21.02 ± 16.36           | 0.833   |
| Crea (mg/dl)                       | 1.19 ± 1.16            | 1.26 ± 1.15             | 0.650   |
| APTT (s)                           | 57.93 ± 25.69          | 62.57 ± 25.43           | 0.175   |
| INR                                | 2.15 ± 1.28            | 2.49 ± 1.26             | **0.046** |
| Na⁺ (mmol/l)                       | 130.97 ± 9.65          | 129.56 ± 9.47           | 0.272   |
| SOFA scores*                       | 8.69 ± 1.76            | 8.72 ± 1.68             | 0.897   |

*Variables/score 0 1 2 3 4

| Variables/score | 0 | 1 | 2 | 3 | 4 |
|-----------------|---|---|---|---|---|
| PO₂/FIO₂ (mmHg) | >400 | ≤400 | ≤300 | ≤200 | ≤100 |
| Platelets (×10⁹/L) | >150 | ≤150 | ≤100 | ≤50 | ≤20 |
| Bilirubin (mg/dl) | ≤1.2 | 1.2–1.9 | 2–5.9 | 6–11.9 | >12 |
| Cardiovascular (Hg/kg/min) | MAP<70 | Dop≤5 | Dop>5 | Epi>0.1 | (Epi≤0.1) |
| Glasgow Coma Scale | 15 | 13–14 | 10–12 | 6–9 | <6 |
| Creatinine (mg/dl) | ≤1.2 | 1.2–1.9 | 2–3.4 | 3.5–4.9 | >5 |

**Table 2.** Sequential Organ Failure Assessment (SOFA) score.

- *Ischemia times and recipients’ SOFA scores were not used as candidate factors. BMI = body mass index; HB = hemoglobin; WBC = white blood cell; PLT = platelet; BUN = blood urea nitrogen; Crea = creatinine; ALB = albumin; TB = total bilirubin; ALK = alkaline phosphatase; GGT = γ-glutamyltransferase; AST = aspartate aminotransferase; ALT = alanine aminotransferase; INR = international normalized ratio of prothrombin time; APTT = activated partial thromboplastin time; Na⁺ = serum sodium; N/A = not applicable.

- References to the table and the study [28].

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characteristics [27]. One effective and widely-accepted way to prevent this over-fitting is to use cross-validation to stop the training at the point of maximum generalization.

**Network training process.** The training rule used in this MLP was supervised, feedforward, back-propagation of error, which could adjust the internal parameters of the network over repeated training iterations to improve the overall accuracy, by modifying the weight of the connections between neurons. In detail, once an input variable is applied as a stimulus to the input layer, it is propagated through hidden layer until an output is generated; this output is then compared with the desired output and an error signal is calculated; this error signal is then transmitted backwards across the net and the weight of the connections between neurons is updated to decrease the overall error of the network; as training proceeds, the difference between the network output and the desired output decreases to a minimum [30].

**Model Validation**

The performances of the MLP, SOFA score, and MELD score in predicting survival at posttransplant one year and two years were assessed in a validation set by measuring both calibration and discrimination ability [31]. We chose these two intervals because outcome at posttransplant one year could reflect surgical and perioperative risk [4], and outcome at two years could also capture mortality associated with most transplant complications, such as rejection and biliary stricture. Calibration refers to the degree of correspondence between predicted and actual survival probabilities. In this study, we used goodness-of-fit testing to evaluate calibration by the Hosmer-Lemeshow test [32], in which the \( \chi^2 \) statistic is the sum of the squared differences between actual and predicted survival probability. Discrimination is usually assessed by the area under a receiver operating characteristic (ROC) curve [33], which is equal to the index of concordance (i.e., \( r \)-statistic). The ROC analysis was also performed to measure the sensitivity, specificity, positive predictive value, negative predictive value, and the total accuracy of these three predictive models.

**Results**

**Outcomes of the entire series of recipients**

Of the 360 DDLT recipients, the mean time on the waiting list was 9.16±3.56 months, and the median follow-up period was 56.23±26.46 months. The overall 6-month, 1-, 2-, 3- and 5-year survival rates were 89.6%, 86.1%, 82.9%, 78.2% and 73.1%, respectively. Of the 360 recipients, 89 recipients (24.7%) died during the 5-year follow-up period. Of these, 23 (6.4%) died within the first 3 months after transplantation of various perioperative causes, including severe fungal infection or sepsis (n = 6), multiple organ failure (n = 4), hepatic artery thrombosis (n = 3), acute rejection (n = 3), primary graft dysfunction (n = 2), upper gastrointestinal bleeding (n = 2), graft versus host disease (n = 2), and subarachnoid hemorrhage (n = 1). 57 (15.8%) recipients died for causes, including severe fungal infection or sepsis (n = 6), multiple organ failure (n = 2), respiratory failure (n = 1), cerebral hemorrhage (n = 1).

**Recipients’ baseline characteristics**

Table 1 and table S1 showed the baseline characteristics of the modeling set and validation set. Most of the characteristics between the two sets have no differences, but we also observed significant differences in the percentage of HBV-DNA level, as well as in the mean values of ALB and INR between the modeling and validation set.

**MLP input features selection**

Two donor factors and ten recipient factors were identified as optimal input features by the forwards stepwise selection algorithm: donor age and BMI; serum concentration of HB, Crea, ALB, TB, ALT, INR, Na⁺; presence of pretransplant diabetes; dialysis prior to transplantation, and microbiologically-proven sepsis. As each sub-category of every categorical variable is an input neuron, there are 18 input neurons in the MLP network.

**Training and development of the MLP network**

By enumerative combinatorial method and making many iterations of training and cross-validation in each combination, we identified 12 hidden neurons that optimally delineated the network and produced the best performance in both one- and two-year intervals. The most appropriate transfer functions were Logistic, Gaussian for one-year network, and Exponential, Identity for two-year network (Fig. 1.).

Taking one input variable, HB as an example, Figure 2 represents the relationships between HB and other variables, and the output prognosis of the trained MLP network. In every subgraph, HB, another variable, and the output prognosis (ie., the MLP target) composed a simulated 3-D rendering; the output prognosis of the network is plotted versus HB and another variable, and the curved surface represents the relationship between HB, the other variable, and the output prognosis. In such a simulated 3-D rendering composed of only two input variables (HB and another variable) and the output prognosis, there is a nonlinear relationship between HB, other variables, and the output prognosis. The relationships between multi-variables and the output prognosis would undoubtedly be even much more complex in corresponding multidimensional space.

**Model validation**

With the Hosmer-Lemeshow test, a P-value greater than 0.05 and close to 1.0 is considered to indicate better calibration, and the smaller the \( \chi^2 \) value, the better the calibration ability of a model [34]. The MLP’s calibration ability (\( \chi^2 = 1.56, P = 0.82 \) in one-year prediction; \( \chi^2 = 1.74, P = 0.78 \) in two-year prediction) was higher than that of the SOFA and MELD in both intervals’ prediction (Table 3).

Table 4 and Figure 3 show the discrimination of the MLP, SOFA score, and MELD score for predicting posttransplant 1-year and 2-year survival probability. The c-statistic values range from 0 to 1, with 0.5 corresponding to what is expected by chance alone and 1.0 to perfect discrimination. For a prognostic model, a c-statistic below 0.7 generally suggests poor prediction, while a c-statistic above 0.7 indicates a useful model, and a c-statistic greater than 0.8 indicates excellent predictive accuracy [24]. The MLP had c-statistics of 0.91 (\( P<0.001 \)) and 0.88 (\( P<0.001 \)) in one-year and two-year prediction, respectively (Table 4 and Fig. 3). The c-statistics of the SOFA were 0.70 (one-year) and 0.67 (two-year). MELD yielded the least accurate predictions (Table 4 and Fig. 3).

**Discussion**

The large disparity between patient demand and donated organs is a pressing problem for all transplant surgeons, especially in the Asia-Pacific region. The best solution to this problem is still in dispute, as there are two sometimes-contradictory principles of
organ allocation: urgency of patient need, and efficiency of organ use [35]. Unfortunately, prioritizing extremely sick patients make it likely that patients who are not as sick “will be forced to wait until their condition worsens and their chances for success are also diminished” [36], and patients who are very sick may have worse posttransplant outcomes than healthier patients [37]. Thus, the optimal system would offer grafts to those who are sufficiently sick to justify the transplantation but not too sick to benefit from it [38], that is, the urgency of need should be jointly optimized with the likelihood of satisfactory outcomes so as to avoid “futile transplantation”.

Furthermore, OLT ranks among the most expensive medical interventions [39], so the urgency-based principle has contributed to rising healthcare costs [37,40]. An accurate prognostic model could also help potential transplant recipients and their families make informed decisions by providing them with information on the patient’s posttransplant survival probability [11,13].

With the aforementioned goals, a newly-adopted lung allocation score in the United States has incorporated likelihood of posttransplant survival in addition to lung disease severity [41]. The liver transplantation field would also benefit from a continuously optimized allocation system that prioritizes patients who need grafts most, without sacrificing the overall utility of this scarce resource. Such a system necessitates a strong prognostic model that can identify potential recipients with satisfactory survival prospects.

Over the past decade, MELD [42] has proved to be an excellent marker of BESELD-specific illness severity and corresponding pretransplant mortality risk, but many studies have also shown its poor accuracy in predicting posttransplant survival [43,44], which is consistent with our results. The SOFA score was originally developed to quantitatively describe the degree of organ dysfunction in six organ systems and to evaluate morbidity in intensive care unit septic patients [26], but later studies found that it could be applied equally well in non-septic critically ill patients to measure individual or aggregate organ dysfunction and to describe morbidity risk [45]. Since its introduction, the SOFA score has also been widely applied to prognostic mortality assessment in critically ill patients with good results [46], although it was not developed for this purpose. In recent years, some

Figure 1. Topological architecture of the MLP network constructed in this study. The network consisted of 18 input neurons, 12 hidden neurons, and 1 output neuron. doi:10.1371/journal.pone.0031256.g001

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investigations have applied the SOFA to critically ill cirrhotic patients and have also proven its validity in mortality risk assessment for BESLD patients [47–49]. We believe that because BESLD patients usually display multiple-organ damage or dysfunction, such as the renal failure, coagulopathy, and encephalopathy, the SOFA is an excellent scoring model for assessing BESLD patients’ illness severity and mortality risk. Additionally, several studies have analyzed the predictive power of SOFA on post-liver transplant mortality; although these achieved some encouraging results in short-term prognosis assessment [50,51], its value in long-term outcome prediction still requires study. In this study, SOFA achieved good calibration abilities in both intervals and satisfactory discrimination power in one-year prediction, which is consistent with other studies [50,51], but its accuracy was poor in two-year prediction. Although SOFA encompasses the functions of multiple systems including respiratory, hemostastics, hepatic, circulatory, and brain and kidney, it is not specific enough to BESLD patients and is not tailored to posttransplant outcome prediction. Lack of these specificities may account for its discriminative and calibration inferiority to the MLP network.

Although there have been many attempts to develop a specific model to assess posttransplant prognosis, to date, they have not achieved sufficient accuracy, or have simply categorized the patients into various risk groups [4,11]; even with some of the most comprehensive efforts, the predictive accuracy of these models has always been reported in the 60–70% range [4,9,11–14] with no single model being more accurate than any other. We believe there are several possible explanations for this. First, the effect of prognostic factors depends on the underlying liver disease [11–13]. Thus, effort would be better spent developing disease-specific models targeted to BESLD patients or cancer patients. Second, Existing studies rely heavily on a few specific variables derived from linear regression analyses, rather than from data mining. The omission of many variables may hinder the discovery of underlying relationships between prognosis and related factors, and the interactions among factors. Third, transplant recipients represent a very complex biological system where the relationship between

| Table 3. Calibration for MLP, SOFA, and MELD in posttransplant survival prediction. |
|---------------------------------|-----------------|-----------------|
| **Goodness-of-fit (x²)**        | **P-Value**     |
| **Postransplant one-year survival prediction** |
| MLP                              | 1.56            | 0.82            |
| SOFA                             | 5.26            | 0.26            |
| MELD                             | 6.48            | 0.17            |
| **Postransplant two-year survival prediction** |
| MLP                              | 1.74            | 0.78            |
| SOFA                             | 5.64            | 0.23            |
| MELD                             | 6.98            | 0.14            |

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pretransplant variables and posttransplant prognosis is multidimensional and nonlinear (as shown in Fig. 2) [17,23], so linear methods are inadequate in predicting regression coefficients and constructing risk factor models.

With the development of artificial intelligence in recent years, ANN has been a superior data-mining solution for complex prognostic problems [17,20], and MLP has been proven to perform better than other architectures such as radial basis function, recurrent neural network, and self-organizing map [22]. MLP is a computation system that uses a large number of simple units to process information in parallel, so it is capable of learning arbitrarily complex nonlinear functions to arbitrary accuracy levels [22]. Furthermore, MLP allows a certain degree of flexibility when it comes to handling noise [18]. Most importantly, MLP is a nonparametric dynamic model, which can automatically self-training and readjust the internal parameters by back-propagation when more transplants enter the network [52], thus yielding more accurate responses and becoming progressively more dependable over time; this is what the linear models could not achieve.

In this study, although three characteristics of the recipients in the validation set differed from the training set, the MLP still achieved good calibration ability and high discrimination power in posttransplant survival prediction, with c-statistics around 0.9 and satisfactory sensitivity and specificity in both intervals, as well as the small $\chi^2$ statistics and associated $P$-values around 0.8 in both intervals. These results were not only superior to that of the linear regression models reported in previous studies [4,9,12,13], but also outstripped the performances of SOFA and MELD in this study. We believe that several factors may account for the MLP’s outstanding performance. First, the MLP network, employing 12 variables to make predictions, included more comprehensive information associated with the posttransplant prognosis. Second, the input features of our MLP included not only donor factors and measurements of disease severity, but also some well-recognized variables reflecting the complications and comorbidities (such as sepsis and diabetes) in BESLD patients. Meanwhile, it should be noted that we decided not to include some subjective variables (such as encephalopathy or ascites) in our model development.

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**Table 4.** Discrimination of MLP, SOFA, and MELD in posttransplant survival prediction.

|                      | C-statistic ± SE | 95% CI | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) | Accuracy (%) | P-Value |
|----------------------|------------------|--------|----------------|-----------------|---------|---------|--------------|---------|
| **Posttransplant one-year survival prediction** |                  |        |                |                 |         |         |              |         |
| MLP                  | 0.91 ±0.05       | 0.80–0.97 | 91.3          | 88.6            | 84.0    | 93.9    | 89.7         | <0.001  |
| SOFA                 | 0.70 ±0.08       | 0.54–0.86 | 72.0          | 66.7            | 62.1    | 75.9    | 69.0         | 0.04    |
| MELD                 | 0.66 ±0.10       | 0.47–0.84 | 68.0          | 63.6            | 58.6    | 72.4    | 65.5         | 0.10    |
| **Posttransplant two-year survival prediction** |                  |        |                |                 |         |         |              |         |
| MLP                  | 0.88 ±0.07       | 0.74–0.96 | 88.0          | 84.8            | 81.5    | 90.3    | 86.2         | <0.001  |
| SOFA                 | 0.67 ±0.09       | 0.50–0.84 | 68.0          | 66.7            | 60.7    | 73.3    | 67.2         | 0.07    |
| MELD                 | 0.65 ±0.10       | 0.47–0.84 | 64.0          | 63.6            | 57.1    | 70.0    | 63.8         | 0.11    |

PPV = positive predictive value; NPV = negative predictive value.  
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**Figure 3.** ROC curves for MLP, SOFA score, and MELD score in posttransplant survival prediction.  
(A): Posttransplant one-year prediction.  
(B): Posttransplant two-year prediction.  
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because their classifications are subjective and could therefore be arbitrary. Third, being computer-based, the MLP can process more information about the survival process and model more complex nonlinear multidimensional relationships, thus yielding more accurate prognostic estimations.

In this study, donor age and BMI were identified as input features. These two factors could be obtained before transplantation, and have been proved to be associated with graft quality [53,54] and recipient outcomes [9,14]. Although some other donor factors (such as the graft steatosis and ischemia times) may directly reflect graft quality and contribute to posttransplant prognosis, they would have been difficult or impossible to know when clinicians and patients make transplant acceptance decisions and when candidates are ranked on a waiting list. This problem would seem to be an inherent difficulty in pretransplant prediction. Therefore, in order to maximize the practical applicability of a pretransplant model, we believe that it must be constructed in accordance with actual clinical conditions, and enhancing the model’s performance based on the variables available is the most important goal. Thus, we decided not to include this kind of characteristics in our pretransplant model development.

Meanwhile, we chose posttransplant one-year and two-year as the study endpoints in this study because outcomes within this timeframe could reflect surgical and perioperative risk [4] and mortality associated with most early complications. However, as we know, the recipient’s long-term survival would be affected by not only the pretransplant characteristics, but also many intraoperative and posttransplant factors, such as the graft cold-ischemia time and biliary complications. Thus, in our view, once the appropriate modeling method is identified, development of sequential correction models according to the different variable acquisition phases may be a reasonable way to meet the evaluation requirement in different phases. When certain donor characteristics, operative parameters, and even some posttransplant variables could be available after operation, another posttransplant predictive model that incorporated above features should be developed and used to perform a further corrective assessment. We believe the two kinds of model can provide more comprehensive perioperative evaluation information at different variable acquisition phases, and, most importantly, they are consistent with actual clinical conditions.

In this study, we clarified the complex multidimensional and nonlinear relationship between transplant variables and posttransplant outcomes, and identified the value of MLP in solving this complex prognostic problem. We believe this methodological result is the key point of this study, and is more important than the specific factors and specific study intervals included in the presented model.

We believe that this kind of pretransplant model would provide patients and clinicians with important reference information about their early posttransplant prospects during the initial counseling and evaluation phases of referral [4,11,13]. If used alongside the MELD system, the pretransplant model can also help predict early outcome with and without transplantation. This provides clinicians with a combined tool to identify patients likely to benefit most from transplantation [9].

Meanwhile, how to ethically balance medical urgency with posttransplant survival prospects is an important issue. For instance, it could be argued that the patient with the highest combined MELD score and survival prospects should be given priority. But we expect that in practice, scientifically combining the two conflicting determinants would not be so simple, just as the use of MELD to guide graft allocation has sparked a wealth of studies and discussion. Therefore, we believe that comprehensively considering and weighing urgency and survival prospects will require further evidence-based research. Whatever shape the final system takes, however, it will undoubtedly include a prognostic model with high predictive accuracy as an important component.

Although this MLP model was more sophisticated than conventional linear models, in practical application, its software implementation allowed the creation of a new interface that can be incorporated into a website and be easily used by everyone, as in the UNOS website, where an interface was created for MELD calculation. Thus, we believe the model’s complexity should not present a problem in clinical practice.

Despite our encouraging results, our study has some potential limitations. First, it was developed using data from a single center; we did not validate our model externally with data from different sources. Indeed, we divided our dataset into training and validation sets, and the validation samples were not used in model development. Thus, the proposed MLP network should be further verified with data at other major centers. Fortunately, the dynamic nature of the MLP makes it capable of continuously and automatically adjusting its internal parameters and improving as more transplant data from other centers enter the network [52]. Second, the patient population had a high proportion of HBV infection; therefore, this MLP network may have limited applicability to typical North American and European patients, who tend to have a lower rates of HBV but higher rates of hepatitis C and alcoholism than do Chinese BESLD patients.

In summary, artificial intelligence methodologies such as MLP offer significant advantages over conventional statistical techniques in variable selection and dealing with restrictive assumptions of normality and linearity, and thus hold promise for solving posttransplant outcome prediction. Therefore, in future research we plan to use MLP to develop a posttransplant multi-interval sequential correction model, a step toward establishing a balanced system that considers both pretransplant mortality and expected posttransplant survival.

Supporting Information

Table S1 Baseline categorical characteristics of the training set and validation set.

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Author Contributions

Conceived and designed the experiments: MZ FY YPL BL. Performed the experiments: MZ FY BC LNY TFW BL. Analyzed the data: MZ FY BC LNY TFW BL. Wrote the paper: MZ FY.

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