Personalized Donor-Recipient Matching for Organ Transplantation

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Abstract
Organ transplants can improve the life expectancy and quality of life for the recipient but carry the risk of serious postoperative complications, such as septic shock and organ rejection. The probability of a successful transplant depends on a very subtle fashion on compatibility between the donor and the recipient - but current medical practice is short of domain knowledge regarding the complex nature of recipient-donor compatibility. Hence a data-driven approach for learning compatibility has the potential for significant improvements in match quality. This paper proposes a novel system (ConfidentMatch) that is trained using data from electronic health records. ConfidentMatch predicts the success of an organ transplant (in terms of the 3-year survival rates) on the basis of clinical and demographic traits of the donor and recipient. ConfidentMatch captures the heterogeneity of the donor and recipient traits by optimally dividing the feature space into clusters and constructing different optimal predictive models to each cluster. The system controls the complexity of the learned predictive model in a way that allows for assuring more granular and accurate predictions for a larger number of potential recipient-donor pairs, thereby ensuring that predictions are “personalized” and tailored to individual characteristics to the finest possible granularity. Experiments conducted on the UNOS heart transplant dataset show the superiority of the prognostic value of ConfidentMatch to other competing benchmarks; ConfidentMatch can provide predictions of success with 95% accuracy for 5,489 patients of a total population of 9,620 patients, which corresponds to 410 more patients than the most competitive benchmark algorithm (DeepBoost).

Introduction
Organ transplantation is the therapy of choice for patients with end-stage diseases who are refractory to medical therapies (Shah 2012). Even though organ transplantation can increase the life expectancy and quality of life for the recipient, the operation can entail various complications, including infection, acute and chronic rejection and malignancy (Huynh 2014). The pre-operative anticipation of the risk associated with organ transplantation is a regular task that transplant centers perform in order to determine which patients would benefit from transplantation and accurately identify these for whom the risk of transplantation is too high and would therefore provide no survival benefits. Such a risk assessment task is quite complicated for that post-operative patient survival depends on different types of risk factors: recipient-related factors (e.g. cardiovascular disease severity of heart recipients (Wozniak 2014; Nwakanma 2007; Russo 2006; Silva 2016)), recipient-donor matching factors (e.g. weight ratio and human leukocyte antigen (HLA) (Jayarajan 2013), blood group compatibility (Jawitz 2013), race (Allen 2010), etc), and donor-related factors (e.g. diabetes (Arnaoutakis 2012; Taghavi 2013)). The interactions among all these risk factors make the prognosis problem for the organ transplant outcomes highly complex; the National Heart, Lung and Blood Institute (NHLBI) working group suggests resolving this problem by enhancing the phenotypic compatibility characterization of the pre-transplant recipient-donor population (Mancini 2010; Collins 2015; Shah 2012).

In the light of the above, we seek an enhanced phenotypic characterization for the compatibility of patient-donor pairs via a precision medicine approach (Collins 2015) in which we construct personalized predictive models that are tailored to the individual traits of both the donor and the recipient to the finest possible granularity. The advent of electronic health records (EHR) inspire a data-driven approach for constructing such predictive models in which the complex recipient-donor compatibility patterns are discovered from observational data. To that end, we develop ConfidentMatch: an automated system that learns the recipient-donor compatibility patterns from the EHR data in terms of the probability of transplant success for given recipient-donor pairs. The clinicians can utilize ConfidentMatch as a prognostic tool for managing organ transplantation selection decisions in which information about the donor and recipient are fed to the system, and the output comes as the probability of the transplant’s success. The system can also be used in conjunction with any matching algorithms, such as the Nobel prize-winning algorithm of Shapley and Roth (Shapley 1962; Roth 2003). More specifically, a set of patients is matched to a set of donors using the compatibility score that ConfidentMatch computes as an input to the matching algorithm.

In order to learn the highly complex recipient-donor compatibility patterns, ConfidentMatch adopts a novel learn-
ing framework in which the recipient-donor feature space is partitioned into disjoint subsets, and a separate predictive model (learner) is assigned to each partition. Such a learning approach gives rise to a highly complex overall predictive model for mapping recipient-donor features to transplant success probabilities. Over-fitting is controlled by penalizing the number of partitions in the recipient-donor feature space and the complexity of the learners assigned to the different partitions. Unlike existing meta-learning algorithms, ConfidentMatch solves an optimization problem through which it jointly determines how to partition the recipient-donor feature space, and what predictive model to assign to each partition. Since such an optimization problem is intractable, we propose an efficient greedy algorithm that proceeds iteratively by first fixing the number of partitions in the recipient-donor feature space, optimizing the predictive models assigned to each partition, and then further stratifying each partition and re-assigning more “specialized” predictive models to the new, finer partitions. The algorithm stops stratifying the recipient-donor feature space when further stratification would lead to new partitions with no enough per-partition training data for learning finer predictive models.

Experiments conducted on the United Network for Organ Sharing (UNOS) heart transplant dataset (Cecka 1996) show that ConfidentMatch can provide predictions of success with 95% accuracy for 5,489 patients of a total population of 9,620 patients – 410 more patients than for the best state-of-the-art machine learning algorithm (DeepBoost).

Related Work

We identify three broad categories of learning algorithms that are capable of combining multiple predictive models or stratifying the feature space into fine clusters: ensemble learning algorithms, clustering algorithms and tree-based algorithms. We compare ConfidentMatch with these methods hereunder.

Ensemble learning algorithms

Methods based on ensemble learning, such as Random Forest (Liaw 2002), LogitBoost (Friedman et al. 2000), Adaptive Boosting (Freund 1997) and DeepBoost (Kuznetsov 2014), operate by allocating different sets of training data to different weak learners, and then aggregating the predictions of these weak learners through a weighted sum to issue a final prediction. Among many applications, ensemble learning has also been successfully used in (Dai et al. 2016) and (Tekin et al. 2016). While these methods can learn complex functions through the synergy of multiple weak learners, they do not integrate the allocation of the training data to the different learners (in both the bagging and boosting approaches) as part of their loss minimization problems. Therefore, these methods do not - in principle - learn a granular predictive model that performs well uniformly over the feature space, but rather learn a predictive model that works well “on average”. Contrarily, ConfidentMatch jointly optimizes the partitioning of the feature space together with the predictive model associated with each partition, and hence, it learns a refined recipient-donor phenotypic compatibility characterization in which the predictions are tailored to fine segments of the recipient-donor feature space, leading to an overall improved performance as compared to conventional ensemble methods. We will demonstrate the superiority of ConfidentMatch to ensemble learning algorithms in the “Results and Discussion” Section.

Clustering algorithms

Clustering is a natural approach for identifying phenotypic characterizations by grouping “similar” patients (or recipient-donor pairs) into distinct clusters. Clustering algorithms can be divided into two categories: unsupervised and supervised clustering. Unsupervised clustering algorithms, such as the $k$-means algorithm (MacQueen 1967), utilize the feature space solely to learn the partitions that maximize some given objective, and hence they cannot address the organ transplant prognosis problem since they do not consider the transplant outcomes (i.e. labels) in the clustering process.

Supervised clustering algorithms utilize both the feature space and the label space for constructing clusters (Eick 2004; Finley 2005); however, the predictive models assigned to each partition of the feature space are limited to indicator functions (an example for such algorithms is the regression tree (Strobl 2009)). For the organ transplant setting, the complex interactions between the donor and recipient features create highly complex patterns of recipient-donor compatibility (transplant success probabilities) that would exhibit very high per-partition impurity under conventional supervised clustering or tree learning algorithms. This means that learning such complex medical concepts would face the dilemma of exhibiting large over-fitting errors when adopting a large number of clusters (or very deep decision trees) to resolve the per-partition impurity, and exhibiting a large bias error when restricting the number of clusters (or restricting the depth of decision trees). ConfidentMatch approaches this problem by providing a versatile framework for complexity control where complex predictive models can be assigned to every partition to reduce the per-partition bias error, which enables learning complex functions with fewer partitions and hence utilize the training data more efficiently.

Tree-based algorithms

As we will show in the next Section, ConfidentMatch can be interpreted as a nonparametric method for learning complex functions by constructing a “tree of base learners”. While ConfidentMatch displays a tree structure, the algorithm is not a conventional “decision tree” (Quinlan 1986, DeSalvo and Mohri 2016, Kim 2004) since the feature space partitions do not correspond to different “decisions” or labels, but rather correspond to different “base learners” that introduce complexity in the earlier levels of the tree, allowing for a more flexible complexity control for the learned hypothesis.

Methods

Let the $D$-dimensional recipient-donor feature space be denoted as $X$; every instance in $X$ corresponds to a recipient-
donor pair with certain given characteristics. Denote the corresponding label space which designates the success or failure of an organ transplant for a given recipient-donor pair as $Y$; every label instance can be defined as the specific event of transplant success or failure if $Y = \{0, 1\}$, or the probability of the transplant’s success if $Y = [0, 1]$. Let $T$ be a dataset extracted from the EHR; we split the dataset $T$ into two separate sets: a training set $S = \{(x_1^1, y_1^1), \ldots, (x_m^s, y_m^s)\}$ and a validation set $V = \{(x_1^V, y_1^V), \ldots, (x_n^V, y_n^V)\}$, where $S$ and $V$ are disjoint ($S \cap V = \emptyset$). Every entry in $S$ and $V$ comprise a recipient-donor feature pair and a transplant outcome.

The goal of ConfidentMatch is to construct a predictive model $h \in \mathcal{H}$, $h : \mathcal{X} \rightarrow \mathcal{Y}$ that maps recipient-donor pairs to anticipated transplant outcomes; such a model has to be learned from the dataset $T = \{S, V\}$, and can be used for out-of-sample recipient-donor pair in order to assess the risk of a recipient’s transplant operation. The problem of learning the predictive model $h \in \mathcal{H}$ from the labeled dataset $T$ is a standard supervised learning problem (Shalev-Shwartz 2014).

The expected loss of a predictive model $h$ is defined as $L_F(h) = \mathbb{E}_{F}(l(h(x), y))$ where $(h(x), y)$ is a general loss function, and $F$ is the joint recipient-donor feature-label distribution, which is unknown to the clinicians. The optimal predictive model is defined as $h^* = \arg \min_{h \in \mathcal{H}} L_F(h)$; since $F$ is unknown, we cannot directly find the optimal predictive model, and hence we resort to minimizing the empirical loss as measured over the training and validation sets. The empirical loss for the training set is defined as

$$L_S(h) = \frac{1}{m} \sum_{i=1}^{m} l(h(x_i^s), y_i^s),$$

and it can be defined similarly for the validation set.

Note that as pointed out in the previous section, the true hypothesis is likely to be of a very complex structure as it abstracts a complex medical concept, i.e. the interactions between the recipient and donor features and their effect on the transplant outcomes. A poor initial choice for the space of possible models $\mathcal{H}$, e.g. letting $\mathcal{H}$ be a hypothesis class with a small VC dimension, may lead to a large bias in the loss function of true hypothesis due to the classic bias-complexity trade-off, and hence we need a more versatile learning framework for which the complexity of the predictive model $h$ adapts to the complexity of the underlying medical concept being learned.

ConfidentMatch adopts a novel framework for crafting complex predictive models out of simpler baseline models by creating a phenotypic characterization of the recipient-donor feature space in which separate predictive models are assigned to disjoint partitions of the feature space. That is, ConfidentMatch outputs a set of partitions that cover the entire recipient-donor feature space, together with a set of predictive models, each tailored to a given partition, thereby leading to an overall complex, granular predictive model. Formally, we divide the recipient-donor feature space $\mathcal{X}$ into $k$ disjoint subsets, where $k$ is to be determined based on the given dataset, in such a way that for each subset, we can have a separate optimal predictive model that minimizes the overall expected risk. We write $\{X_1, \ldots, X_k\}$ to denote a partition of the feature space $\mathcal{X}$, where all such partitions are ensured to be disjoint and cover $\mathcal{X}$. The partition $\{X_1, \ldots, X_k\}$ induces a partition of the training set $S$ and validation set $V$, i.e. the training set is partitioned as $\{S_1, \ldots, S_k\}$, where $S_i = \{(x, y) : (x, y) \in S$ and $x \in X_i\}$. From now on, $i$ indicates the partition index, and $j$ indicates the instance index.

Given the above construct, the learning problem becomes a problem of (jointly) finding the optimal partitioning $\{X_1, \ldots, X_k\}$ of the recipient-donor feature space, together with the optimal predictive model $h_i \in \mathcal{H}$ associated with every partition $i$, i.e. assuming that we know the distribution $F$, the optimal predictive model is found by solving the following optimization problem

$$\min_{\{X_1, \ldots, X_k\} \in \mathcal{H}} \sum_{i=1}^{k} \mathbb{E}_{F_i}[l(h_i(x), y)]$$

subject to $\mathcal{X} = \bigcup_{i=1}^{k} X_i$, and $X_i \cap X_j = \emptyset \forall i \neq j$. (1)

We break down the problem in (1) into two nested optimization problems; we first focus on the solution of the inner optimization problem and define its solution (for a given partitioning $\{X_1, \ldots, X_k\}$) as

$$d(\{X_1, \ldots, X_k\}) = \min_{h_1, \ldots, h_k \in \mathcal{H}} \sum_{i=1}^{k} \mathbb{E}_{F_i}[l(h_i(x), y)].$$

Note that given a partition $\{X_1, \ldots, X_k\}$, the solutions to the inner optimizations are separable, i.e. the optimal predictor of one partition can be determined independent of the choice of the predictors for the other partitions. Hence, we can simplify the inner optimization problem as follows.

$$\min_{h_1, \ldots, h_k \in \mathcal{H}} \sum_{i=1}^{k} \mathbb{E}_{F_i}[l(h_i(x), y)] = \sum_{i=1}^{k} \min_{h_i \in \mathcal{H}} \mathbb{E}_{F_i}[l(h_i(x), y)].$$

Since ConfidentMatch has no access to the true distribution $F$, the algorithm has to learn the partitioning $\{X_1, \ldots, X_k\}$ and the corresponding predictive models $\{h_i\}_{i=1}^{k}$ from the dataset $T = S \cup V$ in such a way that it reaches a loss function that is as close as possible to the true loss in (1). To achieve this, we construct a proxy for the objective in (1) by replacing the terms $\mathbb{E}_{F}(l(h(x), y))$ and $\mathbb{E}_{F_i}[l(h_i(x), y)]$, which depend on the unknown $F$, with their sample estimates $\frac{|V_i|}{n}$ and $\frac{|S_i|}{n}$, respectively. Hence, empirical loss minimization over the validation dataset $V$ can be formulated as
ConfidentMatch constructs the hypothesis class $\mathcal{H}$ in (2), from which we select a hypothesis (or a predictive model) $h_i$ for every partition $i$, by combining the outputs of a finite set of $M$ learners $\{A_1, ..., A_M\}$, each of which can learn a predictive model that belongs to some hypothesis class $\mathcal{H}(A)$. That is, for a fixed partition $X_i$, using the corresponding training set $S_i$, the predictive model that is learned by the learning algorithm $A_i$ for partition $i$ is $A_i(S_i)$. Therefore, the set of all predictive models that can be learned by all the learning algorithms operating on data set $S_i$ is given as $\mathcal{H}(S_i) = \{A_1(S_i), ..., A_M(S_i)\}$. ConfidentMatch decides the optimal partitioning and the optimal predictor for each partition $i$ that belongs to a set of learnable predictors $\mathcal{H}(S_i)$ by minimizing the empirical loss with respect to the validation data set as follows

$$
\min_{k, X_1, ..., X_k} \left[ \sum_{i=1}^{k} \min_{h_i \in \mathcal{H}(S_i)} \frac{|V_i|}{n} \right] \frac{1}{|V_i|} \left( \sum_{(x_i^r, y_i^r) \in V_i} l(h_i(x_i^r), y_i^r) \right)
$$

subject to $X = \bigcup_{i=1}^{k} X_i$, and $X_i \cap X_l = \emptyset$ for $\forall i \neq l$. (3)

Note that the formulation in (3) does not account for the out-of-sample error (or over-fitting); to handle that, we reformulate problem (3) by replacing the objective function with a tight upper bound on the true loss that appropriately penalizes over-fitting. Define

$$
d\left(\{X_1, ..., X_k\}\right) = \sum_{i=1}^{k} \min_{h_i \in \mathcal{H}(S_i)} \left[ \frac{1}{n} \left( \sum_{(x_i^r, y_i^r) \in V_i} l(h_i(x_i^r), y_i^r) \right) \right] + \alpha \sqrt{\frac{k^2 \log M}{n}}.
$$

(4)

where $\alpha \geq 0$ is a penalty parameter. The expression in (4) comprises the sample estimate of the objective and a penalty term $\alpha \sqrt{\frac{k \log M}{n/k}}$ that penalizes: the number of partitions $k$, the average size of a partition $n/k$, and the number of predictive model $M$ from which we chose one model to assign to a given partition. It can be shown that if the penalty parameter $\alpha \geq \sqrt{\frac{1}{2} \frac{1}{2 \log M} \log \left( \frac{2}{1 - (1 - \delta)^{1/k}} \right)}$, then the probability that $d(\{X_1, ..., X_k\})$ is bounded above by $d(\{X_1, ..., X_k\})$ is greater than $1 - \delta$, i.e. $P(d(\{X_1, ..., X_k\}) < d(\{X_1, ..., X_k\})) \geq 1 - \delta$ (the proof can be found in the supporting material). By using the upper bound $d(\{X_1, ..., X_k\})$ as the objective, the empirical loss minimization problem becomes

$$
\min_{X_1, ..., X_k} \left[ \sum_{i=1}^{k} \min_{h_i \in \mathcal{H}(S_i)} \frac{1}{n} \left( \sum_{(x_i^r, y_i^r) \in V_i} l(h(x_i^r), y_i^r) \right) \right] + \alpha \sqrt{\frac{k^2 \log M}{n}}
$$

subject to $X = \bigcup_{i=1}^{k} X_i$, and $X_i \cap X_l = \emptyset$ for $\forall i \neq l$. (5)

Solving (5) is computationally intractable with exhaustive evaluation because the number of possible partitions for $n$ points is the Bell number (recursively, it can be defined as $B_{n+1} = \sum_{i=0}^{n} B_k$ (Wilf 1994)). Therefore, to address this problem, ConfidentMatch adopts an efficient greedy algorithm for approximating the solution to (5). As a first step to construct such an algorithm, we reformulate (5) by incorporating two more constraints. First, we restrict the partitions of the recipient-donor feature space to be hypercubes. A hypercubic partition of the feature space $X$ is defined as $\{X_1, ..., X_k\}$ where $X_i = \prod_{l=1}^{D} [a_{il}, b_{il}]$, $a_{il} \leq b_{il}, a_{il} \in \mathbb{R}^*$, $b_{il} \in \mathbb{R}^*$. ($\mathbb{R}^*$ is defined as $\mathbb{R} \cup \{ -\infty, \infty \}$). Second, we restrict the number of partitions to be $\gamma \in \mathbb{N}$. The optimization problem in (5) with these additional constraints can be stated as follows,

$$
\min_{X_1, ..., X_k} \left[ \sum_{i=1}^{k} \min_{h_i \in \mathcal{H}(S_i)} \frac{1}{n} \left( \sum_{(x_i^r, y_i^r) \in V_i} l(h(x_i^r), y_i^r) \right) \right] + \alpha \sqrt{\frac{k^2 \log M}{n}}
$$

subject to $X_i = \prod_{l=1}^{D} [a_{il}, b_{il}]$, $a_{il} \leq b_{il}, a_{il} \in \mathbb{R}^*$, $b_{il} \in \mathbb{R}^*$

$k \leq \gamma$, where $k \in \mathbb{Z}_+$

$X = \bigcup_{i=1}^{k} X_i$, and $X_l \cap X_i = \emptyset$ for $\forall i \neq l$. (6)

Let $Opt(X)$ be the optimal partition that solves the optimization problem in (6) (it can be solved by the exhaustive evaluation method); we construct a greedy algorithm which iteratively solve the optimization problem (6) to achieve the approximate solution for (5) as follows. We write the partition that is generated when $X$ is input to the optimization problem (6) as $Opt(X) = \{X_1, X_2, ..., X_k\}$ where $k \leq \gamma$. We apply the same procedure recursively on each $X_i$ separately up to the point where we do not expect to improve the objective function (i.e. the optimal $k$ is 1). The final partition is the union of the partitions that are generated by applying this procedure recursively to each $X_i$. We write the final partition achieved by the greedy algorithm as $\{X_1^*, ..., X_k^*\}$.

The pseudo-code for ConfidentMatch is given in Fig. 1. The algorithm learns the recipient-donor compatibility patterns in an off-line manner using the procedure described
above: it jointly optimizes the partitioning of the recipient-donor feature space (off-line stage I), and then optimizes the predictive model associated with each partition (off-line stage II). Having learned the recipient-donor compatibilities, the algorithm operates in an on-line stage for new recipients and donors by computing a compatibility score, i.e. the probability of transplant success, for a given recipient-donor pair. The compatibility score is displayed to the clinicians and based on it, the clinicians/patients can make decisions on whether a transplant should be conducted, or whether the recipient should be matched with another donor.

In what follows, we specify the computational complexity of ConfidentMatch. Let the complexity of the algorithm $A_t(S_t)$ for learning a predictive model using the dataset $S_t$ be $T_t(|S_t|, D)$. Based on this, it can be shown that the worst-case complexity for computing $Opt(X)$ in the optimization problem (6) is $O(\gamma \gamma + 1 n^{D - 1} \sum_{i=1}^{M} T_i(n, D))$, and the complexity of the greedy algorithm described in Fig. 1 is $O(\gamma \gamma + 1 n^{D - 1} \sum_{i=1}^{M} T_i(n, D))$ (proofs are provided in the supporting material).

### Results and Discussion

Experiments were conducted using the UNOS database for patients who underwent a heart transplant over the years from 1987 to 2015 (Cecka 1996). We use the “Thoracic DATA” dataset in the UNOS database as our root dataset. In this dataset, all patients were followed-up until death, i.e. the post-transplant survival times for all patients are available in the dataset. Of the 148,512 patients in the “Thoracic DATA” who underwent either heart or lung transplant, we extract 60,516 patients who underwent a heart transplant. Of the 60,516 patients who underwent a heart transplant, we exclude 3,800 patients (6.28%) who are still alive (right-censored), and we only use the 56,716 patients for whom we have the exact survival (lifetime) information.

For each patient in the dataset, a total of 504 features are provided; these include a combination of both the patient’s and the donor’s information. We discard 12 features that are normally obtained after the transplant. Of the remaining 492 features, we extract 70 features for which we have less than 10% missing information in order to reduce the noise of imputation. We use the k-nearest-neighbor (KNN) imputation method to impute the missing data (Hastie 1999).

We compared the performance of ConfidentMatch in predicting the success of transplants with the following benchmark algorithms: logistic regression (Logit), Lasso regularized logistic regression (Lasso), decision tree (DTree), Random Forest (RFForest), AdaBoost (ABoost), and DeepBoost (DBoost). We use the correlation feature selection (CFS) method to discover the relevant features for the predictive models of both ConfidentMatch and benchmarks (Hall 1999). The validation set calibrated all parameters of ConfidentMatch ($\alpha$) and the benchmark algorithms. We adopt the following metric for quantifying the performance of the different algorithms. The transplant’s success probability is quantified via the 3-year post-transplant survival rate (long-term survival rate). We say that an algorithm provides a prediction for the transplant’s success (probability of 3-year post-transplant survival) for a certain recipient-donor pair with an accuracy level $X\%$ if the algorithm’s probability of correct prediction is $X\%$ for that recipient-donor pair. Based on this definition, we define the gain of ConfidentMatch at an accuracy level of $X\%$ as the number of recipient-donor instances in the testing dataset for which ConfidentMatch provides an accurate prediction of the transplant success, whereas the best-competing benchmark does not.

We split the dataset into a training/validation set comprising the recipient-donor instances in which the recipient underwent the transplant before the year 2010 (former patients), and a testing set that comprises recipients who underwent the transplant after 2010 (current patients). Of the 56,716 recipient-donor pairs, 37,677 pairs (66.43%) were used for training, 9,419 pairs (16.61%) were used for validating and 9,620 pairs (16.96%) were used for testing. We varied the accuracy level from 80% to 95% and evaluated the performance within this range of the accuracy levels. The execution time of the ConfidentMatch on this dataset is less than 5 hours on MATLAB R2015a with Intel i5 (1.5GHz) processor with 4 GB RAM.

Table 2 and Fig 2 show that ConfidentMatch consis-
<table>
<thead>
<tr>
<th>Rank</th>
<th>Overall Recipient-donor Population</th>
<th>Group A</th>
<th>Group B</th>
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<tr>
<td>1</td>
<td>Ventilator Assist</td>
<td>Ventilator Assist</td>
<td>ECMO Assist</td>
</tr>
<tr>
<td>2</td>
<td>ECMO Assist</td>
<td>ECMO Assist</td>
<td>Ventilator Assist</td>
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<tr>
<td>3</td>
<td>Other Life Support</td>
<td>Other Vent Support</td>
<td>Donor VDRL Result</td>
</tr>
<tr>
<td>4</td>
<td>Other Vent Support</td>
<td>Other Life Support</td>
<td>Days in State 1A</td>
</tr>
<tr>
<td>5</td>
<td>Days in State 1A</td>
<td>VAD support</td>
<td>Prior Cardiac Surgery</td>
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<td>6</td>
<td>Diagnosis Code</td>
<td>Diagnosis Code</td>
<td>Blood Type Matching</td>
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<td>7</td>
<td>Donor Status</td>
<td>Donor Status</td>
<td>Donor Blood Type (O)</td>
</tr>
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<td>Transplant Type</td>
<td>Malignancy</td>
<td>Donor Status</td>
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<tr>
<td>9</td>
<td>Donor HEP-P Antigen</td>
<td>Transplant Type</td>
<td>Donor HEP-B Antigen</td>
</tr>
<tr>
<td>10</td>
<td>Previous MI History</td>
<td>Previous Transplant</td>
<td>Inhaled Assist</td>
</tr>
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</table>

Table 1: Top 10 relevant features for the prediction of heart transplant success. (ECMO: Extracorporeal membrane oxygenation, VDRL: Venereal disease research laboratory, VAD: Ventricular assist device, HEP: Hepatitis, MI: Myocardial infarction)

<table>
<thead>
<tr>
<th>Accu.</th>
<th>CM</th>
<th>LASSO</th>
<th>RF</th>
<th>AB</th>
<th>DB</th>
<th>DTree</th>
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<td>7721</td>
<td>9057</td>
<td>9063</td>
<td>9067</td>
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<td>7577</td>
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<td>90%</td>
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<td>4431</td>
<td>6085</td>
<td>6076</td>
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<td>5065</td>
<td>5058</td>
<td>5079</td>
<td>3124</td>
</tr>
</tbody>
</table>

Table 2: Predictions for the success of heart transplant by different algorithms (* Accu. = Accuracy, CM = ConfidentMatch, RF = random forest, AB = ABoost, DB = DBoost).

Figure 2: The number of recipient-donor pairs with confident prediction based on ConfidentMatch as compared to DBoost.

ConfidentMatch does not only improve the quality of conducted, and whether the recipient should be matched with another donor.

The performance gains achieved by ConfidentMatch can be attributed to the improved phenotypic characterization of the recipient-donor pairs that the algorithm achieves by stratifying the recipient-donor feature space. The fine and granular phenotypic characterization achieved by ConfidentMatch is restricted by the size of the training data; the more recipient-donor instances are available in the training set, the larger is the number of partitions that ConfidentMatch can construct and cast a specialized predictive model. Fig 3 illustrates the trade-off associated with increasing the complexity of ConfidentMatch’s predictive model by increasing the number of partitions; if the number of partitions increases, the gain of ConfidentMatch also increases as it copes with the underlying complexity of the recipient-donor compatibility patterns, until a certain number of partitions when the gain starts to decrease due to over-fitting.

ConfidentMatch does not only improve the quality of
prognosis, but can also draw clinical insights on the patterns of recipient-donor compatibility. To illustrate this, we list the first two partitions through which ConfidentMatch stratifies the recipient feature space: ConfidentMatch forms two recipient groups, group A comprises patients whose length of stay in status 1A (urgent transplant wait-list) is shorter than 10 days, whereas group B comprises the remaining patients. Table 1 lists the most relevant features that are predictive of the transplant outcome for each group. It can be seen that group B patients are more sensitive to the donor characteristics; the donor’s Venereal disease research laboratory (VDRL) result and blood type are relevant to the transplant outcome, whereas group A patients appear to be less sensitive to these features. Thus, the more the patient waits in status 1A, the more it becomes essential to consider the extent of her compatibility with the donors. As more training data becomes available, ConfidentMatch can reveal finer partitions and identify the relevant features for more granular recipient subgroups.

Conclusions
Organ transplants for patients with end-stage diseases carries the risk of various serious post-operative complications, the pre-operative anticipation of the transplant outcome depends on the compatibility between the donor and the recipient. In this paper, we have developed ConfidentMatch, a data-driven system that learns complex recipient-donor compatibility patterns from the outcomes of previous transplants. ConfidentMatch captures the complexity of such compatibility patterns by optimally dividing the recipient-donor feature space into clusters and assigning different optimal predictive models to each cluster, thereby ensuring that predictions are “personalized” and tailored to individual characteristics of both the donors and the recipients. Experiments conducted on a public heart transplant dataset demonstrate the superiority of ConfidentMatch to other competing benchmark algorithms.

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References


