Discovery and Clinical Decision Support for Personalized Healthcare

Jinsung Yoon, Camelia Davtyan, MD, FACP, and Mihaela van der Schaar, Fellow, IEEE

Abstract— With the advent of electronic health records, more data is continuously collected for individual patients, and more data is available for review from past patients. Despite this, it has not yet been possible to successfully use this data to systematically build clinical decision support systems that can produce personalized clinical recommendations to assist clinicians in providing individualized healthcare. In this paper, we present a novel approach, Discovery Engine (DE), that discovers which patient characteristics are most relevant for predicting the correct diagnosis and/or recommending the best treatment regimen for each patient. We demonstrate the performance of DE in two clinical settings: diagnosis of breast cancer as well as a personalized recommendation for a specific chemotherapy regimen for breast cancer patients. For each distinct clinical recommendation, different patient features are relevant; DE can discover these different relevant features and use them to recommend personalized clinical decisions. The DE approach achieves a 16.6% improvement over existing state-of-the-art recommendation algorithms regarding kappa coefficients for recommending the personalized chemotherapy regimens. For diagnostic predictions, the DE approach achieves a 2.18% and 4.20% improvement over existing state-of-the-art prediction algorithms regarding prediction error rate and false positive rate, respectively. We also demonstrate that the performance of our approach is robust against missing information and that the relevant features discovered by DE are confirmed by clinical references.

Index Terms— Clinical decision support systems, diagnosis decision support systems, relevant feature selection, healthcare informatics, personalized treatment.

I. INTRODUCTION

Challenge of integrating high-dimensional clinical data to recommend the most appropriate clinical decision for a given patient [1]. As the understanding of complex diseases progresses, the types of available tests and treatments diversify and, as a result, the difficulty of recommending the optimal clinical decision for particular patient increases as well. Current clinical decisions continue to rely on clinical practice guidelines which, in cases where scientific analysis and

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evidence is scarce, are mostly based on clinical experience and opinion. Also, current clinical practice guidelines are aimed at a "representative" patient rather than an individual patient who may display other relevant characteristics. Such "representative" guidelines may thus miss the opportunity to consider personal traits when recommending clinical decisions [53-56]. For example, the American Cancer Society (ACS) recently issued new guidelines which suggested that women with an average risk of breast cancer should start having mammograms at an age of 45 (five years later than ACS had previously recommended) [2]. However, women who have certain risk factors (family history of breast cancer, no children, etc.) have a higher risk of developing breast cancer, and they would benefit from having mammograms at an earlier age. In cases such as these, the ACS guidelines recommend that a high-risk patient consult with her physician to determine an appropriate screening age and interval, which is based on that particular physician's experience and opinion. Moreover, statistics show that diagnostic errors result in 10% of patient deaths and represent the most frequent type of medical malpractice claims in the United States [3]. This reality highly underscores the urgent need for building smart clinical decision support systems (CDSS) and diagnosis decision support systems (DDSS) that can assist clinicians in making accurate, personalized clinical recommendations [4]. It has been recently recognized [5] that medical informatics tools and machine learning techniques can be successfully used to provide recommendations for personalized diagnosis and treatment.

The goal of this paper is to develop methods that will enable CDSS and DDSS to personalize their recommendations based on individual patient characteristics. The wealth of information being routinely collected as part of the electronic health record (EHR) provides an unprecedented opportunity to discover appropriate clinical recommendations for patients given historical information about the clinical decisions administered to similar patients and their actual outcomes [6]. However, using this information is difficult precisely because there is so much of it. The solution is to extract only the relevant information for the particular patient and the relevant clinical decisions previously used for similar patients among the wealth of available information. Extracting only the relevant information is important because using irrelevant features can significantly hurt the performance of the system, unnecessarily increase its complexity, and decrease its learning/adaptation speed [7]. Furthermore, the efficient discovery of relevant patient features can help clinicians focus on the relevant

J. Yoon, and M. van der Schaar are with the Department of Electrical Engineering, University of California Los Angeles, Los Angeles, CA 90095, USA (email: jsyoon0823@ucla.edu, mihaela@ee.ucla.edu).

C. Davtyan is with the Department of Medicine, Geffen School of Medicine at UCLA Los Angeles, CA 90095, USA (email: CDavtyan@mednet.ucla.edu).

information available about the patient without having to sift through a large patient record.

In this paper, we present a novel approach called Discovery Engine which optimizes clinical recommendations by identifying the features in the patient record that differentiate the individuals who receive a certain clinical decision and respond positively from those who do not. Our approach utilizes the available contextual information about patients and learns from the large quantities of observational clinical data to inform clinical recommendations and make better decisions by learning from similar patients. We show that our DE approach consistently outperforms existing state-of-the-art machine learning algorithms both regarding matching individual patients to the optimal treatment regimens as well as diagnosis accuracy.

One of the biggest challenges faced by this class of recommendation systems is that the rewards/actual outcomes of clinical decisions (e.g. five-year recurrence-free survival) are usually not available [8, 52]. Moreover, even if rewards/actual outcomes of the clinical decisions were available, the counterfactuals - rewards/actual outcomes of alternative clinical decisions that were not used – are never available [52]. What is available, however, is a large medical literature that reports the results of a wide range of clinical studies, including different types of patients, different patient characteristics, different types of clinical decisions, and the actual outcomes of these decisions. We use the results of these studies to construct transfer rewards, which we use as proxies for rewards. This allows us to train the DE algorithms as well as to evaluate their performance in comparison to existing methods when the actual outcomes cannot be achieved. The four primary contributions of this paper are as follows:

- We describe a novel approach for discovering the most relevant information from the EHR that distinguishes between patients that should receive one particular clinical decision and the patients who should be given another. For instance, premenopausal breast cancer patients are more likely to respond to a specific type of chemotherapy such as CEF [9].
- Using the past records in the EHR and external knowledge from the medical literature, our approach discovers the optimal personalized clinical decision based on the discovered relevant information (e.g. their clinical test results, treatment history, and outcomes).
- In lieu of having actual reward values associated with clinical decisions, we define the transfer rewards, a method for estimating actual outcomes described in external knowledge (published literature and clinical practice guidelines) based on their similarities to individual patients given reported characteristics.
- We apply DE to two medical applications: 1. Personalized treatment recommendations (chemotherapy regimens) for breast cancer patients and 2. Diagnosis of breast cancer. DE is used to discover which features are relevant to make a distinct clinical decision and then uses this knowledge to build a clinical decision recommendation system. We evaluate the performance of DE in the context of

breast-cancer diagnosis and treatment and show that it consistently and significantly outperforms state-of-the-art machine learning algorithms.

II. RELATED WORKS

A. Personalized Clinical Decision Support System

Current medical practice relies on manually curated systematic reviews of the available scientific evidence and clinical guidelines that provide recommendations for large groups of patients rather than personalized recommendations that are tailored to individual patients. Clinical decision support systems have been proposed before, but many of them do not consider the specific characteristics of patients and do not provide personalized clinical recommendations; hence, they are not very accurate and have only limited applicability in practice [10-11, 53-56]. Moreover, clinicians often refer to the medical literature available through Medline/PubMed, VisualDX, and UpToDate to help them associate observed finding with possible conditions and recommended decisions. However, these resources are also not customized to a specific patient's case.

Several CDSSs are currently implemented and used in clinical settings. For instance, WizOrder was developed to help reduce medical errors and support clinical decisions when entering orders [61]. This system is now used in several neonatal intensive care units at Vanderbilt University Hospital. Assessment and Treatment of Hypertension: Evidence-based Automation (ATHENA) is a CDSS used to manage hypertension in primary care [62]. It was developed to recommend drug therapies and assess and control their effect on blood pressure. ATHENA is now used in several clinics in Northern California. TherapyEdge-HIV is a web-based real-time alerting system for the treatment of HIV, which is used in over 42 clinical sites [63]. While several CDSS are now implemented in clinical settings, their evaluations are often mixed. Some papers argue that CDSS significantly improve the effectiveness of clinical practice [59-60] while others conclude that CDSS do not affect mortality and may even moderately increase morbidity outcomes [57]. Furthermore, the cost-effectiveness of CDSS has yet to be demonstrated [58].

The advent of Big Data has been identified as an opportunity to improve the performance of existing CDSS as well as catalyze the development of new CDSS [12]. A large literature has used data-driven approaches to developing representative rather than *personalized* healthcare decision support systems [13]. A smaller literature is dedicated to developing personalized CDSS. However, most existing papers in this strand of literature [14]-[20] only either just discuss opportunities rather than propose a concrete algorithm, or they apply off-the-shelf machine learning techniques to the considered medical problem and do not address the unique characteristics and challenges of developing personalized CDSS. Diagnosis decision support systems have been developed for cardiovascular diseases and diabetes using ensemble learning [21], SVMs [22], artificial neural networks [23] or rule-based algorithms [24]. Although some diagnosis

decision support systems issue accurate diagnostic recommendations for specific diseases, most of them are based on a small number of *manually* selected features [25-26]. Whenever the number of features (contexts) is large, these methods fail to perform well [27].

Most importantly, most of the proposed CDSS solely focus on diagnosis recommendations and do not provide solutions for the equally important problem of treatment recommendations. A small number of studies attempt to propose CDSS for treatment recommendations [28]. However, these CDSS differ significantly from our DE. For instance, [29] proposes antibiotic recommendation systems for representative patients but does not use a data-driven approach and [30] proposes a CDSS for personalized medicine recommendations, but which uses similarity information among drugs and patients that is specific to the study at hand and cannot be easily applied to other diseases - the similarity between patients is solely based on the ICD 9 codes and the similarity between drugs is based on their chemical structure. Hence, the methods in [29-30] are not widely applicable to a diverse set of patients and treatments.

In contrast, although our DE was only verified in the context of breast cancer so far, it is designed to operate in a variety of complex diseases such as breast cancer, lung cancer, prostate cancer, etc. Furthermore, as we will show in our experimental section, it can perform well even when the number of features used to make a decision is large because it adopts a novel method to discover the most relevant features to consider when deciding on certain diagnosis or treatment options for a specific patient. For this, we developed a customized feature selection and decision-making system which significantly outperforms existing off-the-shelf techniques. Importantly, based on the authors' knowledge, DE is the first personalized CDSS able to discover which specific features of a patient are indicative of treatment success in a complex disease such as breast cancer.

Our work is also related to other works in the field of medical informatics dedicated to improving breast cancer diagnosis and treatment. However, our work is distinct from prior works, which only analyzed the impact of specific patient features (such as genetic information or imaging information extracted from mammograms or other imaging) to improve the performance of CDSS [31-33]. Our DE system is based on a novel set of machine learning methods developed especially for personalized diagnosis and treatment discovery which are shown in later sections to significantly outperform existing methods that are generic. Moreover, DE is not only applicable to breast cancer, but can be generally applied to discover personalized treatment for other complex diseases.

B. Relevant Feature Selection

Another strand of literature related to this work is relevant feature selection algorithms including Correlation Feature Selection (CFS) and Mutual Information Feature Selection (MIFS) [34-35]. These are related to our feature selection approach. However, our clinical decision dependent feature selection algorithm (CDFS) is very different from existing feature selection algorithms which focus on the patients' characteristics and not on how these characteristics distinctively impact different clinical decisions. Our approach is capable of discovering which different features are relevant to each different clinical decision. This makes CDFS similar to our prior work [36-37] - the RELEAF algorithm. However, unlike RELEAF, which is very slow because it must compare all possible combinations of features, CDFS can discover the relevant features in a very fast and efficient manner because it adopts a sequential feature selection approach. This sequential approach significantly reduces the sample and computational complexity of the RELEAF algorithm.

C. Machine Learning Techniques

Our method also exhibits similarities to the contextual multi-armed bandit problem (MAB). However, contextual MABs are very inefficient when the number of contexts (in our case patient features) is large (see the experimental section of [37]). DE is able to successfully deal with the curse of dimensionality by discovering what information is relevant and making clinical decisions based only on relevant contexts rather than the entire set of contexts that can be extracted from

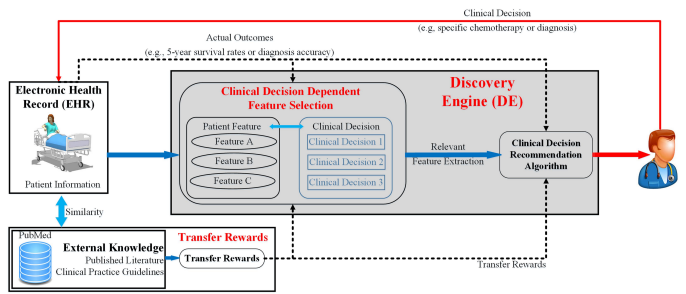


Fig. 1. Personalized clinical decision support system using discovery engine (DE)

the EHR, much of which is irrelevant to the decisions of whether to administer a specific treatment or not.

III. SYSTEM MODEL

In this section, we introduce our method - Discovery Engine (DE). The DE discovers/learns which features/characteristics of a patient are most informative in predicting the success of a clinical decision. For instance, the tumor grade may be found to be relevant for predicting the success of a certain type of chemotherapy in a patient, but not the success of another type of chemotherapy. Thus, different features may be discovered to be relevant for different decisions. Then, when a clinician requests the recommendation from DE for a specific patient, DE decides the best clinical recommendation for the patient which has the best-estimated outcomes. The outcome of a decision is estimated based on the values of the relevant features (i.e. the features found to be relevant for that decision) of the patient. For instance, if the tumor grade was found to be relevant for a certain chemotherapy that chemotherapy will or will not be recommended to that patient depending on that patient's tumor grade.

Fig. 1 depicts the proposed system, which issues a personalized clinical recommendation to the physician about certain patients. The outcomes of certain clinical decisions are used as a reward to train the DE. While the proposed system is applicable in general, we illustrate its use in the context of breast cancer. (A nomenclature table summarizing the variables used and their definitions can be found in the online appendix [64].)

Let $\mathbf{x} = (x_1, x_2, \dots, x_D)$ denote the patient information where D is the total number of patient features such as age, tumor size, estrogen receptor information etc.; $a \in A \triangleq \{a_1, a_2, \dots, a_K\}$ denotes the clinical decision (e.g. chemotherapy regimens or breast cancer diagnosis) that is recommended to the patient. Each patient feature is denoted as $f \in F \triangleq \{f_1, f_2, \dots, f_D\}$. The reward y is derived based on the actual patient outcomes (e.g. five-year survival rates or recurrence rates). Let $\mathbf{x}(n), a(n), y(n)$ be the patient information, clinical decision and reward of n-th patient and $\mathbf{H}_N = (\mathbf{x}(n), a(n), y(n))_{n=1}^N$ be the information available for the N previously seen patients. This represents the training set.

The outcomes of a clinical decision a do not depend on all the features [39]: we assume that the outcomes of a clinical decision a rely only on a subset of features $\mathcal{R}(a) \subseteq F$ which we call the relevant features. Let $\mathcal{R} = \bigcup_{a \in A} \mathcal{R}(a)$ be the set of all relevant features. The DE approach is capable of discovering different features that are relevant to different clinical decisions. We say that $\mathcal{R}(a)$ is relevant/informative for clinical decision a if the expected reward only depends on the information contained in $\mathcal{R}(a)$.

Our goal is to discover the relevant features of each clinical decision a (this may be different for each decision) and recommend the optimal clinical decision that corresponds to the discovered relevant patient information. The optimal recommended clinical decision is given by

$$a^*(\boldsymbol{x_R}) \triangleq arg \max_{\boldsymbol{a}} \mathbb{E}_{\boldsymbol{y}|\boldsymbol{a},\boldsymbol{x_{\mathcal{R}(a)}}}(\boldsymbol{y}|\boldsymbol{a},\boldsymbol{x_{\mathcal{R}(a)}}),$$

where $\mathbb{E}(\cdot)$ is the expectation of the random variable. Therefore, $a^*(x_R)$ is defined as the clinical decision that yields the best

expected patient outcome for a patient characterized by the relevant features x_R .

IV. ALGORITHMS

Discovery engine (DE) consists of two algorithms: a clinical decision dependent feature selection algorithm (CDFS) and a clinical decision recommendation algorithm. As it can be seen in Fig. 1, DE discovers different relevant features for different clinical decisions using CDFS. The detailed steps of each algorithm are described in the following subsections, and the pseudo-code of DE can be found in the online appendix [64].

A. Clinical Decision Dependent Feature Selection (CDFS)

To describe CDFS, we start by introducing a few notations. Let \hat{y}_a and N_a be the sample mean rewards estimate and the number of patients who received the clinical decision a, respectively. Similarly let $\hat{y}_a^S(x_S)$ and $N_a^S(x_S)$ be the sample mean rewards estimate and the number of patients (whose feature information contains x_S and was provided clinical decision a), respectively. We formalize these variables as:

$$N_a = \sum_n \mathbb{I}\{a(n) = a\},\tag{1}$$

$$\hat{y}_a = \frac{1}{N_a} \sum_n \mathbb{I}\{a(n) = a\} \times y(n), \qquad (2)$$

$$N_a^S(\mathbf{x}_S) = \sum_n \mathbb{I}\{\mathbf{x}_S \subset \mathbf{x}(n)\} \mathbb{I}\{a(n) = a\},$$
(3)

$$\hat{y}_a^S(\boldsymbol{x}_S) = \frac{1}{N_a^S(\boldsymbol{x}_S)} \sum_n \mathbb{I}\{\boldsymbol{x}_S \subset \boldsymbol{x}(n)\} \times \mathbb{I}\{\boldsymbol{a}(n) = a\} \times \boldsymbol{y}(n) \,,\, (4)$$

where $\mathbb{I}\{\cdot\}$ is the indicator function.

We define the relevance metric $h_f^r(a)$ as the variance of the rewards for a certain action a if a given feature x_f is considered, when selecting the action: $|\hat{y}_a^f(x_f) - \hat{y}_a|$. This is weighted by $\frac{N_a^f(x_f)}{N_a}$ which represents the frequency with which feature x_f is present when action a is selected. We formalize this as:

$$h_f^r(a) \triangleq \sum_{x_f} \frac{N_a^f(x_f)}{N_a} |\hat{y}_a^f(x_f) - \hat{y}_a|$$
 (5)

Also, we define a redundancy metric $h_{f,s}^d(a)$ which measures how the expected reward made for a given patient is affected by considering an additional feature x_f when clinical decision a is recommended. We formalize this as:

$$h_{f,s}^{d}(a) = -\sum_{x_f, x_s} \frac{N_a^{f,s}(x_f, x_s)}{N_a} [\hat{y}_a^{f,s}(x_f, x_s) - \hat{y}_a^{s}(x_s)]$$
 (6)

Then, we define $\mathcal{U}_f(a)$ as the utility obtained if feature x_f is additionally selected as a relevant feature for clinical decision a. Let $\widehat{\mathcal{R}}(a)$ be defined as the previously discovered relevant features set for clinical decision a. Then, the utility function $\mathcal{U}_f(a)$ is defined as:

$$\mathcal{U}_f(a) = h_f^r(a) - \frac{1}{|\hat{\mathcal{R}}(a)|} \sum\nolimits_{s \in \hat{\mathcal{R}}(a)} h_{f,s}^d(a) \tag{7}$$

where $1/|\hat{\mathcal{R}}(a)|$ is used as a normalization factor. The main steps of the CDFS are outlined below:

Step 1: For each clinical decision a, initialize $\widehat{\mathcal{R}}(a)$ as the empty set (i.e. \emptyset) and its complementary set ($\widehat{\mathcal{R}}^c(a)$) as the set of all features (i.e. F).

Step 2: The algorithm selects the first relevant feature which maximizes the relevance metric $(h_f^r(a))$, i.e.,

$$G = arg \max_{f \in \hat{\mathcal{R}}^c(a)} h_f^r(a)$$
$$\hat{\mathcal{R}}(a) \leftarrow \hat{\mathcal{R}}(a) \cup G$$

Step 3: The algorithm finds the subsequent relevant feature that maximizes utility function $(U_f(a))$, i.e.,

$$H = \arg \max_{f \in \hat{\mathcal{R}}^{c}(a)} \mathcal{U}_{f}(a)$$
$$\hat{\mathcal{R}}(a) \leftarrow \hat{\mathcal{R}}(a) \cup H$$

Step 4: The algorithm iteratively runs Step 3 until the utility function $\mathcal{U}_f(a)$ is less than threshold cost C, where C is an input parameter for the algorithm which can adjust the number of relevant features for each clinical decision, i.e.,

If
$$\max_{f \in \hat{\mathcal{R}}^c(a)} \mathcal{U}_f(a) < C$$
,
then, $\hat{\mathcal{R}}(a) = \hat{\mathcal{R}}(a)$

Discovered relevant features are used to recommend the optimal clinical decision in clinical decision recommendation algorithm.

B. Clinical Decision Recommendation Algorithm

The proposed clinical decision recommendation algorithm recommends the optimal clinical decision which maximizes the estimated patient outcome only based on the relevant contexts (features) discovered by CDFS. The main steps of the recommendation algorithm are outlined below:

Step 1: Find the set of unresolved clinical decisions (U) for the patient with information vector $\mathbf{x}_{\widehat{\mathcal{R}}(a)}$:

$$U = \left\{ a \in A \middle| N_a^{\hat{\mathcal{R}}(a)} \left(\boldsymbol{x}_{\hat{\mathcal{R}}(a)} \right) < C_{TH} \cdot \log(n) \right\}$$
 (8)

where $C_{TH} \cdot \log(n)$ is a control function. C_{TH} is an input parameter which can adjust the trade-off between the confidence of the clinical recommendations and the learning speed of DE.

If a set of unresolved clinical decisions exists $(U \neq \emptyset)$, DE abstains from making clinical decision recommendations and only updates $N_a^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(a)})$ and $\hat{y}_a^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(a)})$ based on the obtained rewards. In other words, DE only issues recommendations when it is sufficiently confident about its clinical recommendations and it abstains otherwise.

Step 2: If there are no unresolved clinical decisions (i.e., $U = \emptyset$) for the patient with information vector $\mathbf{x}_{\widehat{\mathcal{R}}(a)}$, the optimal clinical decision with respect to the relevant feature set $\widehat{\mathcal{R}}(a)$ is determined as

$$\hat{a}(\mathbf{x}) = \arg\max_{a} \hat{y}_{a}^{\hat{\mathcal{R}}(a)} \left(\mathbf{x}_{\hat{\mathcal{R}}(a)} \right) \tag{9}$$
 This optimization selects the clinical decision with the

This optimization selects the clinical decision with the maximum estimated reward for the patient with relevant information vector $\mathbf{x}_{\hat{\mathcal{R}}(a)}$.

After the rewards of the recommended clinical decision are obtained, $N_{\hat{a}}^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(\hat{a})})$ and $\hat{\mathcal{Y}}_{\hat{a}}^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(\hat{a})})$ are updated as follows:

$$N_{\hat{a}}^{\hat{\mathcal{R}}(a)}\left(\boldsymbol{x}_{\hat{\mathcal{R}}(a)}\right) = \sum_{n} \mathbb{I}\left\{\boldsymbol{x}_{\hat{\mathcal{R}}(a)} \subset \boldsymbol{x}(n)\right\} \mathbb{I}\left\{a(n) = \hat{a}\right\}$$
(10)

$$\hat{y}_{\hat{a}}^{\hat{\mathcal{R}}(a)}(\mathbf{x}_{\hat{\mathcal{R}}(a)}) = \frac{y(n)}{N_{a}^{\hat{\mathcal{R}}(a)}(\mathbf{x}_{\varsigma})} \sum_{n} \mathbb{I}\{\mathbf{x}_{\hat{\mathcal{R}}(a)} \subset \mathbf{x}(n)\} \mathbb{I}\{a(n) = \hat{a}\} \quad (11)$$

The computational complexity of DE is $O(ND^2)$; hence, DE has a relatively low run-time complexity with high dimensional datasets.

C. DE with Missing Information

Electronic health records, more often than not, may have missing information for some patients [52]; hence, DE must be able to operate properly even with missing information.

Suppose that the dataset contains missing information. We can divide the feature information vector \mathbf{x} into two components: the available features (\mathbf{x}^{av}) and the missing features (\mathbf{x}^m) . Thus, $\mathbf{x} = \{\mathbf{x}^{av}, \mathbf{x}^m\}$. First, the relevance metric of CDFS is solely computed based on the available information:

$$h_f(a) \triangleq \sum_{x_f^{av}} \frac{N_a^f(x_f^{av})}{N_a} |\hat{y}_a^f(x_f^{av}) - \hat{y}_a|$$

Therefore, if the feature f is frequently missing, $h_f(a)$ decreases, and as a result the feature f is rarely selected as a relevant feature.

Second, it should also be noted that we can estimate the reward $(\hat{y}_a^{\hat{\mathcal{R}}(a)}\left(x_{\hat{\mathcal{R}}(a)}^{av}\right))$ with missing information based on a given patient's available relevant information, $x_{\hat{\mathcal{R}}(a)}^{av}$, for each clinical decision a. More specifically, it can be estimated as:

$$\hat{y}_{a}^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{av}) = \mathbb{E}(\hat{y}_{a}^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{av}, \boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{m}) | \boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{av})$$

$$= \sum_{\boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{m}} \hat{y}_{a}^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{av}, \boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{m}) \cdot P(\boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{m} | \boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{av})$$

$$= \sum_{\boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{m}} \hat{y}_{a}^{\hat{\mathcal{R}}(a)}(\boldsymbol{x}_{\hat{\mathcal{R}}(a)}) \cdot P(\boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{m} | \boldsymbol{x}_{\hat{\mathcal{R}}(a)}^{av})$$

We can estimate the conditional probability, $P(\mathbf{x}_{\mathcal{R}(a)}^m | \mathbf{x}_{\mathcal{R}(a)}^{av})$, based on the probability distribution of the features in the training set. Based on this estimation rule, we can robustly identify the optimal clinical decision even if there is missing information.

V. TRANSFER REWARDS

As discussed in the introduction, the most valuable rewards for most clinical decision support systems are, in theory, the actual patient outcomes (e.g. 5-year survival rates or recurrence rates in the case of breast cancer). However, these outcomes are very difficult to obtain in practice [52]. Instead, we use a proxy for outcomes based on external knowledge which consists of published literature and clinical practice guidelines. We refer to all external knowledge simply as *references* in the remainder of the paper.

The idea is to match patients to appropriate relevant references. For each patient and each reference, we define the term similarity as the amount of information that reference provides about that patient. The similarity is computed by calculating the posterior probability of that patient feature belonging to the population demography from the reference. Then we aggregate the actual outcomes of certain clinical decisions for each reference according to the similarity (posterior probability) and use this as a transfer reward for that clinical decision when applied to that patient. The system model for transfer reward estimation is illustrated in Fig. 2.

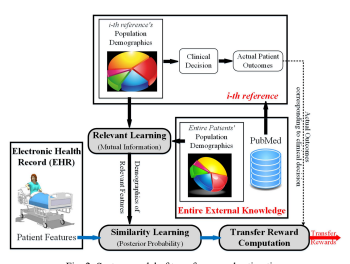


Fig. 2. System model of transfer reward estimation

To compute the transfer rewards, we first estimate the similarity between a patient and a reference. The first step of estimating this *similarity* is to find the relevant patient features for each reference; we do this using a sequential feature selection algorithm based on the mutual information in order to deal with population demography [35]. The mutual information between the *i-th reference* (E_i) and the *k-th* feature (f_k) is defined as:

$$I(f_k; E_i) = \sum_{x \in \chi_k} P(x|E_i) \log \frac{P(x|E_i)}{P(x)}$$
 (12)

where $P(x|E_i)$ is the probability of feature x in reference i, P(x) is the probability of feature x across the entire set of references, and χ_k is context space of f_k . Let $\hat{\mathcal{R}}(E_i)$ be defined as the discovered relevant feature set for reference E_i and the utility function $\mathcal{U}_f(E_i)$ is determined as:

$$U_f(E_i) = I(x_f; E_i) - \frac{1}{|\widehat{\mathcal{R}}(E_i)|} \sum_{s \in \widehat{\mathcal{R}}(a)} I(x_f | E_i; x_s | E_i)$$
 (13)

where $1/|\hat{\mathcal{R}}(E_i)|$ is used as a normalization factor. This utility function measures an increment of mutual information between relevant feature set and the reference when feature f is the additionally selected as relevant feature.

The algorithm selects the first relevant feature which maximizes the mutual information with E_i , i.e.

$$G = arg \max_{f \in \hat{\mathcal{R}}^c(E_i)} I(x_f; E_i)$$
$$\hat{\mathcal{R}}(E_i) = \hat{\mathcal{R}}(E_i) \cup G$$

Then, the algorithm finds the subsequent relevant feature that maximizes utility function $U_f(E_i)$, i.e., $H = arg \max_{f \in \hat{\mathcal{R}}^c(E_i)} U_f(E_i)$

$$H = arg \max_{f \in \hat{\mathcal{R}}^{c}(E_{i})} \mathcal{U}_{f}(E_{i})$$
$$\hat{\mathcal{R}}(E_{i}) = \hat{\mathcal{R}}(E_{i}) \cup H$$

The algorithm iteratively adds new relevant features in $\widehat{\mathcal{R}}(E_i)$ until the maximum utility function $\mathcal{U}_f(a)$ becomes less than zero.

The second step is to compute a posterior probability of a patient feature set belonging to the population demography from the *reference*. Given the *n-th* patient, characterized by the feature vector $\mathbf{x}_n = \{x_1(n), \dots, x_D(n)\}$, we compute the posterior probability that the given patient belongs to the

population demography of the *reference*; we express this value as $P(E_i|X_1 = x_1(n), ..., X_D = x_D(n))$. We compute this via Bayes rule; it is computationally convenient to take logarithms:

$$\log(P(E_i|X_1 = x_1(n), ..., X_D = x_D(n))$$

$$= \log\frac{(P(X_1 = x_1(n), ..., X_D = x_D(n)|E_i)) \cdot P(E_i))}{P(X_1 = x_1(n), ..., X_D = x_D(n))}$$

$$\cong \log P(E_i) + \sum_{l \in \hat{R}(E_i)} \log\frac{P(X_l = x_l(n)|E_i)}{P(X_l = x_l(n))}$$

where we write $P(E_i)$ as the probability of selecting the *i-th* reference as the best clinical decision for the entire population. We define this approximation of posterior probability as a similarity between *n-th* patient and *i-th* reference ($Sim_{E_i}(x_n)$).

Second, we compute the estimated transfer reward of each clinical decision a for n-th patient as a weighted sum of actual outcomes of clinical decisions in each reference according to the *similarity* ($Sim_{E_i}(\mathbf{x})$). i.e.,

$$\widehat{Sim}_{a|E_i}(\mathbf{x}) = \frac{Sim_{E_i}(\mathbf{x})}{\sum_{i:a \in A(E_i)} Sim_{E_i}(\mathbf{x})}$$

$$tr_a(\mathbf{x}) = \sum_{i:a \in A(E_i)} \widehat{Sim}_{a|E_i}(\mathbf{x}) \cdot r(a|E_i)$$

where $r(a|E_i)$ is an actual patient outcome for clinical decision a in *i-th reference*, $(\widehat{Sim}_{a|E_i}(x))$ is a normalized similarity for clinical decision a, and $A(E_i)$ is the set of clinical decisions considered in i-th reference. We define this estimated reward as the transfer reward $(tr_a(x))$. These provide a complete ranking of each clinical decision for each patient. The clinical decision with the highest transfer reward is the optimal clinical decision for the given patient. The pseudo-code for estimating transfer rewards is given in the online appendix [64].

VI. EXPERIMENT I: CHEMOTHERAPY RECOMMENDATION FOR **BREAST CANCER PATIENTS**

While our DE algorithm can be applicable in general clinical decision support systems, in this section, we illustrate its use in the context of a personalized recommendation system of chemotherapy regimens for breast cancer patients in this section. Fig. 3. shows the system model of this application.

A. Data Description

From an initial set of 2.353 references (performing a narrow search of breast cancer chemotherapy regimens using PubMed Clinical Queries), 32 references were selected for further analysis. (The complete list of the 32 references is provided in the online appendix [64].) The list was compiled based on the following two criteria: (1) References contain the clinical outcomes for at least one of the six standard chemotherapy regimens for breast cancer patients [7, 42-43]; (2) References include the demographic information of the breast cancer patients enrolled in the randomized trials.

The sample size of reported references ranged from 50 to 3,934 individuals. There was no crossover of individual subjects between these references. A summary of the population demographics, chemotherapy regimens and actual outcomes in *references* is provided in the online appendix [64].

We evaluate our DE algorithm and benchmarks on the de-identified database of 10,000 breast cancer patients which

was created based on the patients participating in the National Surgical Adjuvant Breast and Bowel Project (NSABP) in the United States and Canada. The patients were diagnosed with operable and palpable breast cancer by core needle biopsy or fine needle aspiration. The patient data is characterized by 15 features summarized in Table II, and those are also corresponding with the patient features in 32 references.

TABLE I CODE FOR EACH CHEMOTHERAPY REGIMEN

Code	Specific Chemotherapy Regimen		
AC	Doxorubicin+ Cyclophosphamide		
ACT	Doxorubicin+ Cyclophosphamide+ Taxanes		
AT	Doxorubicin+ Taxanes		
CAF	Cyclophosphamide+ Doxorubicin+ 5-Fluorouracil		
CEF	Cyclophosphamide+ Epirubicin+ 5-Fluorouracil		
CMF	Cyclophosphamide+ Methotrexate+ 5-Fluorouracil		

We iteratively evaluated the performance of the algorithms based on 10 rounds with 10 different training sets and reported the average performance as a final performance of each algorithm. In each round, we used a randomly selected training set of 4,000 patients among 10,000 entire patients and a disjoint testing set of 6,000 patients. In other words, no training data were used during testing of the model, but 10 different models were used to derive the average performance. We select 4,000 patients to be in the training set since the performance of all algorithms (besides ACL) saturated beyond this number of patients.

TABLE II SUMMARY OF PATIENT INFORMATION FEATURES

SUMMART OF TATIENT INFORMATION TEATORES				
Feature	Range	Features	Range	
Age	30s ~ 60+	PLNC*	0 ~ 10+	
Menopausal	Pre/Post	Lymph Node Status	Pos/Neg	
Race	White/Black/ Other	WHO Score*	0 ~ 5	
Estrogen Receptor	Pos/Neg	Surgery Type	BCT*/MRM*/ No	
Progesterone Receptor	Pos/Neg	Prior Radiotherapy	Exp / No	
HER2NEU*	Pos/Neg/Neu	Prior Chemotherapy	Exp / No	
Tumor Stage	$T1 \sim T4$	Histology	Ductal / Mix / Lobular	
Tumor Grade	G1 ~ G3			

^{*}PLNC: positive axillary lymph node count

B. Benchmarks

We compare the performance of DE with 9 state-of-the-art classification algorithms, ensemble learning algorithms, and feature selection algorithms which are widely used in CDSS and also commonly used benchmarks in medical and machine learning references:

- Correlation Feature Selection (CFS): a well-known feature selection algorithm [34];
- All Contextual Learning (ACL): a well-known contextual learning algorithm which uses all features. This is a

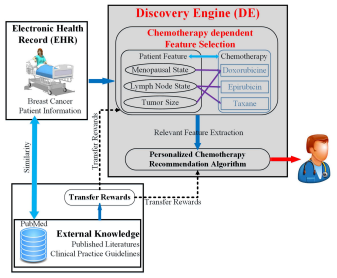


Fig. 3. Personalized chemotherapy recommendation system for breast cancer patients using DE algorithm

modified offline version of the contextual bandit algorithm of Slivkins [40];

- Multivariate Logistic Regression (Logit);
- Linear Regression (Linear);
- Multivariate Support Vector Machines (SVM); we use a radial basis function (RBF) kernel SVM;
- Support Vector Machines with Feature Selection (SVMs-f)
- Adaptive Boosting (AdaBo);
- Classification Tree (CTree);
- Regularized Multivariate Logistic Regression using Lasso (ReLog);
- Regularized Linear Regression using Lasso (ReLin);

C. Success of the optimal chemotherapy recommendation for breast cancer patients

Given a patient, both our algorithm and the benchmark algorithm recommend a chemotherapy regimen corresponding to particular *references*. If the recommended chemotherapy has the highest estimated transfer reward for the patient among all six chemotherapy regimens, we regard the algorithm in question as making the correct recommendation for that patient; i.e. it has recommended the best course of treatment. (Notice that the best course of treatment may not promise a good outcome: some cancers are not treatable.) We take the fraction/percentage of correct recommendations to be the success rate for the algorithm in question.

Given the success rate for the algorithm, we apply two performance metrics: the simple percent agreement and the Cohen's kappa coefficient [44]. Simple percent agreement (p_0) is the success rate (the fraction of times the personalized treatment prediction coincides with the recommendation provided in the medical literature for the patient). Cohen's kappa coefficient (κ) is a metric which measures inter-rater agreement. It is usually considered a more robust measure than a simple percent agreement (p_0) , because κ measures the improvement over chance agreements. If p_e is the probability of agreement by chance, then, kappa coefficient is defined as $\kappa = \frac{p_0 - p_e}{}$

^{*}BCT: breast conservative therapy. MRM: modified radical mastectomy

^{*}HER2NEU: human epidermal growth factor receptor 2

^{*}WHO score: Eastern Cooperative Oncology Group (ECOG) score which is widely used in publications by the World Health Organization (WHO)

^{*}Features with categorical values are changed mutually exclusive binary indicator for the evaluation.

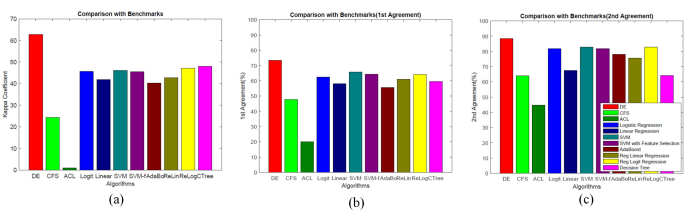


Fig. 4. Performance analysis with benchmark algorithms (a) Kappa coefficient, (b) 1st simple percent agreement, (c) 2nd simple percent agreement

The bar graphs in Fig. 4(b) show that the first chemotherapy recommendation of DE is successful (as defined above) 73.4% of the time and Fig. 4(c) also shows that one of the first two recommendations is successful 88.4% of the time. This is 7.7% better than the second-best approach (SVM) regarding selecting the optimal chemotherapy on its first choice, and 5.6% better regarding matching the optimal chemotherapy within the first 2 choices. This is already a significant improvement; however, regarding kappa coefficients, the improvement is even greater: DE works 16.6% better than SVM, which is described in Fig. 4(a). This is because SVM indiscriminately recommends the popular chemotherapies and is not robust when classifying the less popular chemotherapies. Given robustness considerations, which are essential in medical treatment recommendations, kappa coefficients are more often used as a performance metric in medical informatics.

When comparing our algorithm with other algorithms that rely on feature selection, we again see a significant improvement. Again, note that while other algorithms use feature selection, they do not select relevant features for specific chemotherapies, and it is through this selection that our algorithm achieves improvement. CFS achieves only a 48% of simple percent agreement because it cannot use the efficacy of the chemotherapy to discriminate the relevant features and hence the technique is entirely unsupervised. ACL succumbs to the "curse of dimensionality" because there are 15 features with different ranges, resulting in over 7 million combinations to explore. Logistic regression, linear regression, and SVM perform worse than DE because they do not consider the relevant features for selecting chemotherapies at all.

D. Relevant Features for Each Chemotherapy

Table III shows the top 4 ranked relevant features discovered by CDFS - tumor stage, positive axillary lymph node number (PLNC), estrogen receptor, etc.- for recommending AC, ACT, AT, CAF, CEF and CMF chemotherapy. As it can be seen from Table III, CDFS can discover the different relevant features that are relevant for different chemotherapy regimens.

It is important to note the features discovered by DE are indeed confirmed to be relevant by clinical studies. Firstly, note that the six considered chemotherapy regimens are commonly recommended to node positive breast cancer patients, i.e. patients where cancer has been found in the lymph nodes [43]. It is critical to know whether lymph nodes are positive or

negative. PLNC tells us both the number of nodes and whether lymph nodes are positive or negative. For instance, zero PLNC implies node negative breast cancer, while otherwise indicates node-positive breast cancer. Hence, PLNC is selected as a relevant feature by CDFS. Secondly, the menopausal status is considered important because medications affect cancer differently in premenopausal and postmenopausal women [9]. specifically, the CEF chemotherapy is only recommended for premenopausal women. Although the menopausal status is not included in this relevant feature set, women over the age of 50 are usually considered postmenopausal [9]. Therefore, age was correctly identified by DE to be a discriminative feature for selecting among chemotherapy regimens. Thirdly, tumor stage is another important feature to consider when deciding among chemotherapy regimens as described in reference [42]. Medications A(Doxorubicin), T(Taxotere), E(Epirubicin) are recommended for advanced breast cancer and our top six chemotherapy regimens include more than one of these medications. Therefore, DE has correctly discovered that the features that are relevant for these chemotherapy regimens contain tumor stage information. Finally, the medication T(Taxotere) is usually recommended to breast cancer patients who do not respond to their current chemotherapy. Thus, the prior chemotherapy information is correctly discovered by DE to be relevant for AT and ACT therapies.

TABLE III
DISCOVERED RELEVANT FEATURE FOR EACH CHEMOTHERAPY

DISCOVERED RELEVANT PEATURE FOR EACH CHEMOTHERAFT				
Chemotherapy	1 st Relevant	2 nd Relevant	3 rd Relevant	t 4 th Relevant
Code	Feature	Feature	Feature	Feature
AC	PLNC	Tumor Stage	Estrogen Receptor	Age
ACT	Tumor Stage	Prior Chemotherapy	PLNC	Estrogen Receptor
AT	Prior Chemotherapy	PLNC	Surgery Type	Age
CAF	Surgery Type	Tumor Stage	Age	Tumor Grade
CEF	PLNC	Estrogen Receptor	Tumor Stage	Age
CMF	Estrogen Receptor	PLNC	Radio therapy	Tumor Stage

E. Performance when Patient Information is missing

As explained before, patient information is often missing from the EHR. Moreover, studies have shown that the missing information is often not random [45]. For example, the age of the patient is easy to record, and blood pressure is often verified several times by a nurse when a patient is seen by a clinician, so it is typically neither missing nor incorrect. However, HER2NEU may not be recorded depending on the diagnostic tests ordered and capabilities of a medical center.

Fig. 5. describes the performance of DE when features are missing with various rates. It shows the performance degradation of DE and the benchmark algorithms as a function of the average degree of incompleteness [45]. We did not use the percentage of missing features as a metric since the features are not randomly missing. The percentages of missing features corresponding to each degree of incompleteness are described in the online appendix [64]; these percentages were computed based on statistics extracted from medical records of patients. Fig. 5. shows that the performance of DE degrades from 73.4% to 63.0% (when the average degree of incompleteness is 50%). However, even with missing information, DE continues to outperform the other methods. DE discovers relevant features with low missing probability and can estimate the missing feature information based on the available feature information. As a result, the impact of missing information is minimized. In fact, DE performs better than most other algorithms even when DE misses significant amounts of information from the EHR while the other algorithms make their decision with full information. Hence, we can indeed see that the performance of DE is robust even when information is missing.

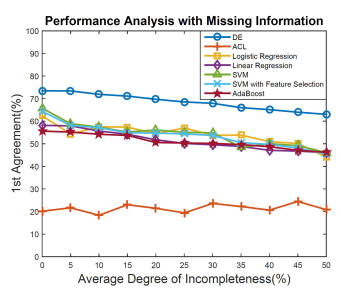


Fig. 5. Performance analysis with missing information

VII. EXPERIMENT II: DIAGNOSIS DECISION SUPPORT SYSTEM FOR BREAST CANCER PATIENTS

In this section, we illustrate how the DE algorithm can be used for breast cancer diagnosis. In this case, we can directly use patients' actual outcomes as the rewards. Fig. 6 describes the system model of this application.

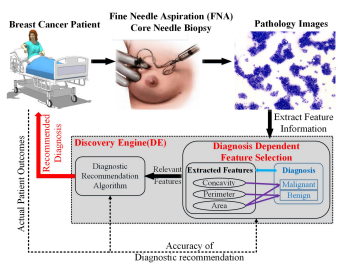


Fig. 6. Personalized diagnostic recommendation system for breast cancer patients using DE algorithm

A. Data Description

TABLE IV SUMMARY OF THE FEATURE INFORMATION

Information Type		Explanation
1	Radius	Mean of distance from center to points on
	Kaulus	the perimeter
2	Texture	Standard deviation of gray-scale values
3	Perimeter	The perimeter of tumor cell nucleus
4	Area	The area of tumor cell nucleus
5	Smoothness	Local variation in radius lengths
6	Compactness	Perimeter^2 / area -1
7	Concavity	Severity of concave portions of the contour
8	Concave Points	Number of concave portions of the contour
9	Symmetry	Symmetricity of tumor cell nucleus
10	Fractal Dimension	Coastline approximation – 1

^{*}Features consist of mean, standard errors and worst of above 10 info types.

In this section we evaluate the performance of DE for breast cancer diagnosis using the UCI *Diagnostic Wisconsin Breast Cancer Database* [46]. The dataset contains 30 patient features extracted from needle biopsy features such as radius, compactness, or smoothness of tumor cell nucleus. Table IV summarizes the details of 30 patient features. The number of instances in this dataset is 569 and the diagnosis (label) for each instance is either malignant or benign.

B. Benchmarks

We compare the performance of DE in this clinical setting with all the benchmarks describes in Section VI-B. Also, we add three existing state-of-the-art feature selection algorithms as the benchmarks to compare the performance of our feature selection algorithm (CDFS) separately.

- Mutual Information Feature Selection (MIFS): a well-known feature selection algorithm based on mutual information [35];
- Relevance Learning with Feedback (RELEAF): an action dependent relevance learning algorithm based on the expected rewards [36-37];
- Principal Component Analysis (PCA): a statistical procedure to discover linearly uncorrelated variables based on orthogonal transformation [47]

^{*}Each info type is computed real value feature for each tumor cell nucleus.

C. Experiments Setup

First, we compare our DE algorithm against state-of-the-art machine learning algorithms: logistic/linear regressions, rbf kernel SVM, adaptive boosting algorithms, classification tree and regularized logistic/linear regressions. We use 10-fold cross-validation to evaluate the performance of algorithms. We performed 10 independent cross-validation runs and reported the average performance of 10 runs.

To highlight the impacts of CDFS, we performed two additional sets of experiments. In the first set, we compared the performance of our DE system using CDFS with the performance of the DE system where CDFS was replaced with one of the four different feature selection algorithms: CFS, MIFS, RELEAF, and PCA. This comparison shows the impact of CDFS on the overall performance of the DE. Other experiment settings are the same as the first experiment.

In the second set of additional experiments, we use our feature selection algorithm CDFS in conjunction with the diagnostic recommendation made by the benchmark algorithms - linear regression, logistic regression, SVM - to highlight the specific impact of our feature selection algorithm. Other experiment settings are the same as the first experiment.

D. Performance of diagnostic recommendation for breast cancer patients

Given a patient, DE and the other benchmark algorithms classify the tumors as malignant or benign. To quantify their performance, we apply three different performance metrics: the prediction error rate (*PER*), the false positive rate (*FPR*), and the false negative rate (*FNR*). *PER* is defined as the fraction of times the recommended diagnosis of our algorithm is different from the actual label. *FPR* and *FNR* are defined as the diagnosis error rate for benign instances and malignant instances, respectively. The main goal of DDSS is to minimize the FPR given an allowable threshold for FNR as selected by the clinicians. In practice, this is often set to be below 2% [25]. Therefore, in this experiment, the FNR threshold is also set to be 2%. Using this threshold, we can characterize our performance metrics as follows.

minimize FPR subject to $FNR \le 2\%$

Comparison with machine learning algorithms:

TABLE V
COMPARISON WITH STATE-OF-THE-ART MACHINE LEARNING TECHNIQUES

COMPINION	VIIII BIAIL OF THE I	HET WITTETHINE EETHE	TECHNIQUES
%	PER	FPR	FNR
DE	<mark>2.23</mark>	<mark>2.62</mark>	1.92
Logit	11.77	18.3	1.96
Linear	8.47	13.55	1.98
SVMs	4.41	6.82	1.99
SVMs-f	4.52	7.03	1.98
AdaBo	9.12	14.86	1.91
CTree	11.45	18.64	1.95
ReLog	6.71	10.11	1.92
ReLin	5.51	9.15	1.93

As the Table V shows, our DE algorithm has 2.23% prediction error rates and 2.62% false positive rates which is 2.18% and 4.20% better than the second-best algorithm (SVM)

when the tolerable threshold of FNR is set to below 2%. There are two reasons for the outstanding performance of the DE approach. First, our diagnostic recommendation algorithm yields high accuracy for classification, because it can provide personalized diagnosis while other comparable algorithms apply the same model for all patients. Second, DE can discover different relevant features for different diagnosis based on CDFS, while the other algorithms (Logistic/Linear Regression, SVMs, AdaBoost and Classification Tree) base their decisions on all the features without relevant feature discovery.

Comparison with feature selection algorithms:

In this subsection, we demonstrate the impact of the CDFS algorithm on the DE system. We compare the performance of the DE using CDFS with the performance of DE using different feature selection algorithms.

TABLE VI
PERFORMANCE OF DE WITH OTHER FEATURE SELECTION METHODS

%	CDFS	RELEAF	CFS	MIFS	PCA
PER	<mark>2.23</mark>	18.37	2.76	3.19	3.94
FPR	<mark>2.62</mark>	24.11	3.90	3.99	6.44
FNR	<mark>1.92</mark>	1.96	1.98	1.90	1.94

As seen in Table VI, CDFS outperforms all other feature selection algorithms when the tolerable threshold of FNR is set to below 2%. This is because CDFS is capable of discovering diagnosis relevant features based on their impact on the expected diagnosis accuracies. Although RELEAF also considers the dependence between diagnosis accuracy and feature selection, it is extremely slow when the number of features is large, as is the case for this and many other medical datasets. Furthermore, combinatorial approach (RELEAF) requires a relatively large amount of training sets to accurately discover the relevant feature, which is not the case for the medical dataset available to us.

TABLE VII
IMPACT OF THE CDFS IN CONJUNCTION WITH ALTERNATIVE MACHINE
LEARNING ALGORITHMS

EE III (II (O I IE COILI III II)				
	PER (%)		FPR (%)	
	CDFS w/o CDFS		CDFS	w/o CDFS
Linear	<mark>5.49</mark>	8.47	<mark>8.88</mark>	13.55
Logit	<mark>7.84</mark>	11.77	<mark>12.7</mark>	18.3
SVMs	<mark>4.01</mark>	4.41	<mark>5.51</mark>	6.82

Next, we replace DE's recommendation algorithms with various existing machine learning algorithms to demonstrate the impact of the CDFS component of DE on the diagnostic decisions. As seen in Table VII, CDFS improves the performance of all benchmark algorithms because it can accurately discover and select the (different) features that are relevant for different diagnosis.

E. Relevant Features for Diagnostic Decision

Table VIII shows the top 5 ranked relevant features discovered by CDFS – worst perimeter, concave points concavity, radius, and area, etc.- for diagnosing malignant or benign cancer among all of the features summarized in Table IV. As seen in Table V, CDFS can discover the different features that are relevant for different diagnoses.

It should be noted that the relevance of the features discovered by DE confirms findings of prior clinical studies. For instance, studies of breast biopsies [48] state that the 3 most important factors to diagnose tumor cell nuclei as malignant or benign are the relative size ratio between nucleus and cytoplasm, irregular shape, and irregular chromatin. However, because chromatin feature information is not available in our dataset, only the relative size and irregular shape can be potential candidates as the relevant features. The size-related features are radius, perimeter, and area, and the shape-related features are the concavity and concave points [48]. The top 5 features found to be relevant by DE to classify malignant and benign tumor cell nuclei are all related to the tumor shape and relative size, which is in accordance to reference [48]. Features such as texture, smoothness, compactness, symmetry, and fractal dimension are not found to be relevant by DE and are not mentioned as important features in reference [48]. Hence, we can conclude that DE can discover the relevant features for making a correct diagnosis without prior medical knowledge.

TABLE VIII
DISCOVERED RELEVANT FEATURES FOR EACH DIAGNOSIS

	Malignant	Benign	
1st Relevant	Worst perimeter	Worst concave points	
Feature	worst perimeter	worst concave points	
2 nd Relevant	Worst concave points	Mean concave points	
Feature	worst concave points	Mean concave points	
3 rd Relevant	Worst radius	Worst perimeter	
Feature	worst radius		
4 th Relevant	Mean concavity	Worst radius	
Feature	Mean concavity	worst radius	
5 th Relevant	Worst area	Maan aanaaritu	
Feature	w orst area	Mean concavity	

VIII. DISCUSSION AND FUTURE WORKS

We proposed a novel approach that discovers the most relevant information from the EHR to determine which clinical decision to recommend for a patient. Furthermore, our approach uses this information to provide personalized recommendations to assist physicians in their decision making process. Our results demonstrate that DE outperforms existing machine learning, prediction, and feature selection methods in both CDSS and DDSS applications. These superior recommendations are extremely important because they have the potential to prevent medical errors and thus improve the quality of medical care. We also showed that our method is robust against missing information, which is important in numerous clinical settings.

Future work will consider that feature information (such as the tumor size, PLNC number and tumor radius in the case of a breast tumor) may change over time. The time dependence may also influence the duration of a therapy and the selection of future therapies. Another DE extension will consider the global sequence of treatment decisions that optimizes long-term outcomes (e.g. overall survival rate or 5-year recurrence rate).

In conclusion, we believe that our proposed contextual learning approach demonstrates promise towards providing useful personalized clinical recommendations. As new types of treatment are evaluated and approved for medical use, clinicians will have an increasingly difficult time determining which clinical decisions are most effective for individual patients. DE provides a pathway towards providing computational methods for personalized clinical decision recommendations.

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Jinsung Yoon received the B.S. degree in electrical engineering from Seoul National University, Seoul, Korea in 2014 and the M.S. degree in electrical engineering from University of California, Los Angeles, USA, in 2016. He is currently pursuing his Ph.D. degree in electrical engineering at

University of California, Los Angeles. His research interests include machine learning algorithms, medical informatics, data mining, recommendation systems, and prediction systems. He was a recipient of the Graduate Study Abroad Scholarship from the National Research Foundation of Korea in 2014.



Camelia Davtyan is a Clinical Professor of Medicine in the Division of General Internal Medicine at UCLA and Faculty at the Iris Cantor UCLA Women's Health Center, one of the 14 Centers of Excellence in Women's Health in the United States. She serves as the Director of

Women's Health of the Comprehensive Health Program at UCLA and Associate Medical Director of the UCLA Medical Hospitality Program. Women's health has been the primary focus of her practice since 1996. She has published and lectured on Women's Health topics for health professionals and the general public and was featured on Lifetime Television and CNN. Dr. Davtyan is Board certified as a Diplomate in Internal Medicine. Dr. Davtyan has consulted for International Women's Health projects in Romania and Armenia. She serves as a member of the UCLA Healthcare Utilization Review Committee and Quality Management Committee. She is a member of the American Medical Women's Association and the American College of Women's Health Physicians and has been elected Fellow of the American College of Physicians.



Mihaela van der Schaar is Chancellor Professor of Electrical Engineering at University of California, Los Angeles. Her research interests include network economics and game theory, online learning, dynamic multi-user networking and communication, multimedia

processing and systems, real-time stream mining. She is an IEEE Fellow, a Distinguished Lecturer of the Communications Society for 2011-2012, the Editor in Chief of IEEE Transactions on Multimedia and a member of the Editorial Board of the IEEE Journal on Selected Topics in Signal Processing. She received an NSF CAREER Award (2004), the Best Paper Award from IEEE Transactions on Circuits and Systems for Video Technology (2005), the Okawa Foundation Award (2006), the IBM Faculty Award (2005, 2007, 2008), the Most Cited Paper Award from EURASIP: Image Communications Journal (2006), the Gamenets Conference Best Paper Award (2011) and the 2011 IEEE Circuits and Systems Society Darlington Award best Paper Award. She received three ISO awards for her contributions to the MPEG video compression and streaming international standardization activities and holds 33 granted US patents.