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Machine Learning of Human Pluripotent Stem Cell-Derived Engineered Cardiac Tissue Contractility for Automated Drug Classification

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SUMMARY

Accurately predicting cardioactive effects of new molecular entities for therapeutics remains a daunting challenge. Immense research effort has been focused toward creating new screening platforms that utilize human pluripotent stem cell (hPSC)-derived cardiomyocytes and three-dimensional engineered cardiac tissue constructs to better recapitulate human heart function and drug responses. As these new platforms become increasingly sophisticated and high throughput, the drug screens result in larger multidimensional datasets. Improved automated analysis methods must therefore be developed in parallel to fully comprehend the cellular response across a multidimensional parameter space. Here, we describe the use of machine learning to comprehensively analyze 17 functional parameters derived from force readouts of hPSC-derived ventricular cardiac tissue strips (hvCTS) electrically paced at a range of frequencies and exposed to a library of compounds. A generated metric is effective for then determining the cardioactivity of a given drug. Furthermore, we demonstrate a classification model that can automatically predict the mechanistic action of an unknown cardioactive drug.

INTRODUCTION

Drugs in late-stage development as well as those with market approval are often withdrawn due to previously undetected drug-induced cardiotoxicity. Unpredicted druginduced cardiotoxicity jeopardizes patients' lives, erodes public trust in the regulatory process, and financially burdens the pharmaceutical industry. For example, cisapride, a gastrointestinal drug intended to treat heartburn, was reported to have caused serious ventricular arrhythmias and sudden deaths prior to withdrawal (Ferriman, 2000). The associated pharmaceutical company agreed to settle lawsuits for a total of \$90 million US dollars for 300 deaths and 16,000 injuries (Harris and Koli, 2005). Progress has been made to facilitate better safety through the adoption of US Food and Drug Administration guidelines that recommend screening new drugs with the human ethera-go-go-related gene (hERG) inhibition assay (FDA, 2005). Nonetheless, within the past decade, an assortment of market-approved therapeutics (e.g., clobutinol, sibutramine, and tegaserod) had to be withdrawn due to unpredicted drug-induced cardiotoxicity (FDA, 2016; Li et al., 2016).

The need for more accurate and faster pre-clinical detection methods has led to the emergence of a variety of screening platforms in recent years. A majority of these detection systems have begun to utilize human pluripotent stem cell-derived cardiomyocytes (hPSC-CMs). These cells are more physiologically similar to human myocardium than laboratory animals or genetically transformed noncardiac cell lines (e.g., human embryonic kidney cells) (Dick et al., 2010). Aside from the commonality of using hPSC-CMs, these platforms vary greatly in their setup, including differences in tissue geometry. Certain platforms employ hPSC-CMs in a two-dimensional manner (e.g., monolayer), while others attempt to recapitulate aspects of the three-dimensional environment of native myocardium by modeling the cells as cardiac muscle fibers or heart chambers (Chen et al., 2014; Huebsch et al., 2016; Lee et al., 2008). These systems can also differ drastically in their methodology of quantifying changes in hPSC-CMs exposed to cardioactive compounds. Some systems examine the electrophysiological properties while others focus on the calcium transients or measurements of contractility (e.g., shortening, force, pressure) generated by the cardiomyocytes (Lu et al., 2015; Maddah et al., 2015; Navarrete et al., 2013; Zhang et al., 2014).

While the experimental platforms may vary, quantitative readouts generally characterize individual contractile events. Combining this with the number of experimental conditions (e.g., various pacing frequencies or drug concentrations) can yield large multidimensional datasets that make it difficult to draw thorough conclusions. Researchers often simplify the raw data by preselecting a limited number of parameters as an attempt to







comprehend the complexity of the data, albeit losing information in the process. Without full interpretation of such a rich dataset, there is a risk of not detecting information that differentiates the behavior of normal hPSC-CMs from those exposed to cardioactive compounds. In addition, as these platforms are meant for high-content screens, the analysis of the datasets needs to be automated.

Holistic approaches must be developed to optimize the utility of datasets generated from screening platforms. Machine learning has been shown to handle such multidimensional datasets in an automated fashion (Lee et al., 2015). We previously demonstrated that support vector machine (SVM), a supervised learning algorithm, can be used to consolidate 12 parameters, which characterize the contractile behavior of hPSC-CMs exposed to cardioactive compounds, into a singular quantitative index that expresses the level of induced cardioactivity (Lee et al., 2015). Machine learning can be further leveraged into a suite of tools that provide more in-depth details of hPSC-CM behavior when exposed to cardioactive compounds. In this study, we hypothesize that multiclassification algorithms can be implemented to create a model to define drug classes and subsequently predict an unknown compound's mechanistic action. Such information would assist in streamlining the drug discovery pipeline, allowing for the rapid identification of select compounds for more in-depth follow-up assays. In addition, this information coupled with knowledge of predicted class can guide scientists to efficiently and selectively screen for specific drug-to-drug interactions that prompt cardiotoxicity (e.g., disruption of Ca²⁺ handling when sofosbuvir and amiodarone are combined) instead of relying on the traditional brute force approach (Millard et al., 2016). Furthermore, drugresponse relationships between the unknown compound and the library can be determined.

To test our hypothesis, we examined a database (not yet published) containing drug screens of various compounds on twitch force measurements from human ventricular cardiac tissue strips (hvCTS) engineered from hPSC-CMs embedded in a 3D collagen-based matrix (Turnbull et al., 2014). A unique aspect of these screens was that the hvCTSs were electrically paced at four different frequencies from 0.5 to 2.0 Hz, spanning a physiological range. These measurements interrogated the influence of cardioactive compounds on the hvCTS force-frequency relationship and contributed to a multidimensional dataset. We selected a total of 12 compounds with acute cardiac effects that represented five drug classes (1, Ca²⁺ channel blockers; 2, adrenergic agonists; 3, cardiac glycosides; 4, hERG K⁺ channel blockers; and 5, angiotensin-converting enzyme [ACE] inhibitors) along with one reference compound (aspirin). In this study, we report the use of machine learning to establish a drug classification model based on hvCTS contractile behavior (using half of the selected compounds) and subsequently demonstrate predictive capabilities by having the model classify unknown compounds, which were withheld from the machine during training.

RESULTS

Formation of the Drug Classification Model

To form the drug classification model, the screens of 12 compounds (Table 1) acquired on the hvCTS platform were used. Each of the compounds, with the exception of aspirin, belonged to one of five classes with each class comprising a minimum of two compounds. Aspirin functioned as a reference for a cardiac-neutral compound. To quantify the cardioactive effects of these compounds, a total of 17 parameters were derived from each contractile event recorded in the hvCTS twitch force versus time tracings (Figure S1). Once the parameters characterizing each contraction were calculated, establishing the library with machine learning consisted of two primary steps. The first step was determining the degree of cardioactivity for each compound at a given dosage by calculating a singular quantitative index generated by a binary SVM approach (Figure 1A). The second step involved multiclass SVM to establish the boundaries that represent each drug class (Figure 1B).

The binary SVM is capable of summarizing all parameters and providing a simple metric that expresses a compound's degree of cardioactivity at a given dosage (Lee et al., 2015). Specifically, the machine is tasked with creating a decision boundary that separates two groups (data from untreated hvCTSs and those from hvCTSs exposed to a concentration of a compound) as seen in Figure 1A. The decision boundary is evaluated for generalizability by classifying withheld data, referred to as a test set. As a result, an SVM accuracy metric is calculated to reflect the machine's ability to identify a cardioactive effect (e.g., 75% accuracy means that out of 100 data points, the machine could correctly classify 75 of them). To account for variations within the dataset, multiple runs are performed to calculate the mean SVM accuracy. A value of approximately 50% mean SVM accuracy suggests non-cardioactivity as the machine cannot distinguish between treated hvCTS data from control data, and the classification becomes random. Prominent cardioactive effects allow the space between the data from treated and control conditions to become more distinguishable, leading to a higher SVM accuracy with 100% being the maximum.

The second step was the utilization of multiclass SVM to create and evaluate a model. The 11 compounds (excluding aspirin) were divided into two groups (Figure 1B). The first group was referred to as a "drug library" and used to train



Compound Name	Class	Description	Test Range (M)	Number of hvCTS (<i>n</i>)
Nifedipine	Ca ²⁺ channel blocker	an L-type Ca ²⁺ channel blocker known to shorten action potential duration (Harris et al., 2013)	10^{-8} to 3.0 × 10^{-5}	10
Mibefradil	Ca ²⁺ channel blocker	a tetralol derivative that blocks both L- and T-type Ca ²⁺ channels with higher affinity for T-type (Martin et al., 2000)	10^{-9} to 3.0 × 10^{-6}	6
Isoproterenol	adrenergic agonist	a mixed β-adrenergic agonists; compound is non-selective in terms of βreceptors (Steinberg, 1999)	10^{-8} to 10^{-4}	10
Norepinephrine	adrenergic agonist	mixed adrenergic agonist that stimulates both α and $\beta receptors (Yang et al., 2014)$	10 ⁻⁹ to 10 ⁻⁵	8
Digoxin	cardiac glycoside	a cardiac glycoside that inhibits Na ⁺ /K ⁺ -ATPase, resulting in higher intracellular Na ⁺ ; higher Na ⁺ concentration suppresses the Na ⁺ /Ca ²⁺ exchanger causing the accumulation of intracellular Ca ²⁺ (Katz et al., 2010)	10 ⁻⁸ to 10 ⁻⁴	9
Duabain	cardiac glycoside	a cardiac glycoside that affects Na ⁺ /K ⁺ -ATPase, which consists of both α and β subunits; has a lower affinity for α subunits than digoxin (Katz et al., 2010).	10 ⁻⁸ to 10 ⁻⁴	10
Flecainide	hERG K⁺ channel blocker	a mixed hERG K ⁺ blocker that also inhibits Na ⁺ channels, causing effects on action potential repolarization and conduction (Harris et al., 2013)	10 ⁻⁸ to 10 ⁻⁴	8
E-4031	hERG K ⁺ channel blocker	a pure hERG K ⁺ channel blocker known for its pro-arrhythmic potential (Ziupa et al., 2014)	10 ⁻⁸ to 10 ⁻⁴	8
Cisapride	hERG K ⁺ channel blocker	a serotonin (5-HT4) receptor agonist that also inhibits the hERG K ⁺ channel (Wong et al., 2010)	10^{-8} to 10^{-4}	9
Lisinopril	ACE inhibitor	an ACE inhibitor, which reduces vasoconstriction and lowers blood pressure in patients (Williams, 1988)	10^{-8} to 10^{-4}	8
Ramipril	ACE inhibitor	an ACE inhibitor; it does not block ACE until it is converted by the liver (Williams, 1988)	10 ⁻⁹ to 10 ⁻⁵	7
Aspirin	non-cardioactive reference	non-steroidal anti-inflammatory drug that has been shown to have no cardioactive effects in screening platforms (Maddah et al., 2015)	10 ⁻⁸ to 10 ⁻⁴	6

n refers to independent biological replicates. hvCTS, human ventricular cardiac tissue strips; hERG, human ether-a-go-go-related gene; ACE, angiotensin-converting enzyme.

a model that defined the boundaries of each drug class. The second group was completely withheld from the computer throughout the entire training and tuning of the model and was referred to as the "unknown" compounds. To normalize for the varying degrees of cardioactivity among compounds, the concentration of a compound that achieved a metric closest to 85% mean SVM accuracy (see Supplemental Experimental Procedures) was used in the formation and evaluation of the model. To ensure that the

library was generalizable or had the capability to classify itself, a subset of the first group's data, a test set, was randomly withheld from the computer prior to training. The library was evaluated on its performance to accurately identify this test set. Afterward, the library was asked to predict the unknown compounds, and its performance was evaluated for predictive capabilities. To account for the variation and random selection of training and test sets, the creation and evaluation of the models were performed 50 times.



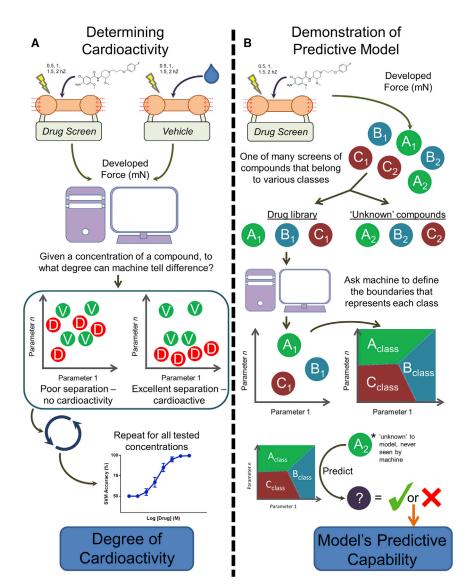


Figure 1. Determining Cardioactivity and Predicting Mechanistic Action

(A) To determine if a concentration of a compound was cardioactive, parameters describing the force waveforms were compared with those of vehicle study with the binary SVM approach. If the compound does not modulate the contractile behavior of hvCTSs, the generated data points would be similar to those of vehicle, yielding poor separation and an SVM accuracy of approximately 50%. However, if a compound's cardioactive effects become more distinquishable, separation between the two groups becomes more feasible and results in a higher SVM accuracy (100% as maximum distinguishability). This binary SVM approach generates a singular quantitative index that describes the degree of cardioactivity of a given concentration for a drug compound.

(B) To create a model for the prediction of mechanistic action, data from screened compounds were divided into two groups: library and unknown. Using data from the library group, the machine defined boundaries that represent various drug families. The model was evaluated for its predictive capabilities by having the machine classify the unknown compounds.

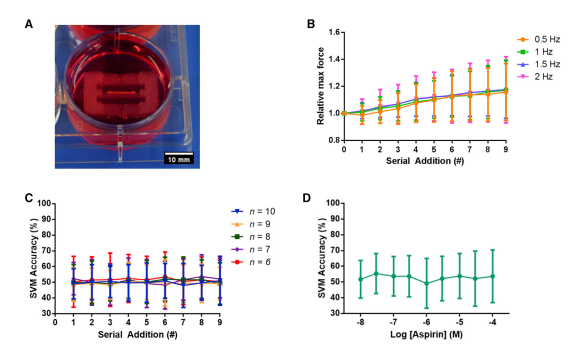
Control Experiments

Although the hvCTSs (Figure 2A) were examined under temperature-controlled conditions, there was an observable drift in contractile behavior of the vehicle-treated hvCTSs. For example, the relative measured maximum developed force increased for all pacing frequencies by a cumulative average of $16.96\% \pm 0.83\%$ upon the ninth serial addition (Figure 2B). To account and normalize for baseline drift, each drug condition was compared with its respective vehicle condition via binary SVM (e.g., measurements of the seventh serial drug addition were compared with those of vehicle-treated hvCTSs at the seventh serial addition). To establish a benchmark of non-cardioactivity, a subset of the vehicle-treated hvCTSs was randomly selected to model a non-cardioactive compound. Binary SVM was then performed between the subset and a corresponding control group of equal size (n).

To ensure that the number of hvCTSs in the subset had no effect, the calculations were performed with the sample size, n, equal to 6–10, which matched the range of numbers of strips used in each drug study of the 12 tested compounds. As expected, the SVM accuracy, regardless of the size of n, was approximately 50% for all serial additions (Figure 2C). These results indicated that there were no consistent and distinguishable trends within each serial addition, and therefore a reference of non-cardioactivity was created.

To validate this reference of non-cardioactivity, drug screens of aspirin from the database were used as negative controls. Aspirin is known to have no cardioactive effects on hPSC-CMs (Lu et al., 2015; Maddah et al., 2015; Scott







(A) hvCTSs were formed by compaction of hPSC-CMS, fibroblasts, and extracellular matrix solution around two PDMS posts. Force was derived from the deflection of the posts. Scale bar, 10 mm.

(B) hvCTSs of the vehicle study exhibited a drift in contractile behavior after serial additions of water (n = 28).

(C) To normalize for shift, data from each drug condition were compared with its respective vehicle condition (matching serial addition number). With this, a benchmark of non-cardioactivity was created with subsets of vehicle-treated strips being modeled as strips exposed to non-cardioactive compounds and compared with other vehicle-treated strips with binary SVM. The number of tissue strips in the subset (n = 6-10) had no effect as SVM accuracies were approximately 50% for all conditions.

(D) To assess the non-cardioactivity benchmark, the data of aspirin-treated hvCTSs indicated no statistical difference from the vehicle study over the tested range (n = 6). n refers to independent biological replicates.

All results are presented as means \pm SD.

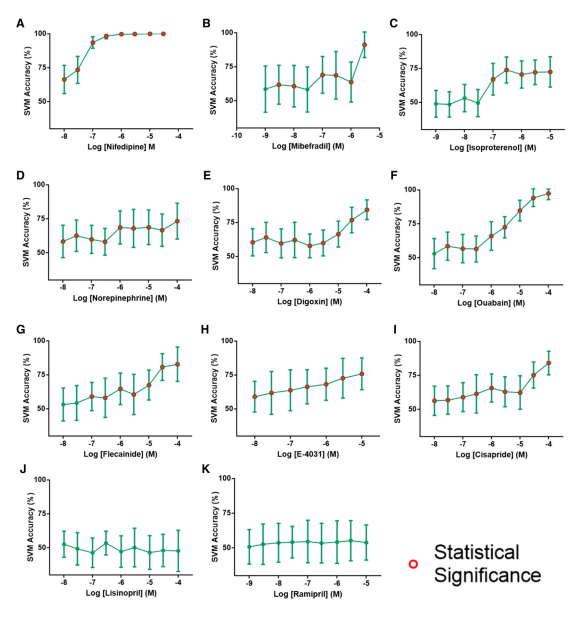
et al., 2014). The SVM accuracies of the aspirin drug screens (n = 6) had an average of 52.85% ± 1.77% among all serial additions (10 nM to 100 μ M). None of the conditions were statistically different from vehicle counterparts, indicating non-cardioactivity by aspirin (Figure 2D).

Generalizability of the Drug Classification Model

In setting up the drug classification model, the 11 nonreference compounds were compared with vehicle-treated tissue strips with the aforementioned binary SVM approach. At one or more of the tested concentrations, all but two compounds, lisinopril and ramipril, had SVM accuracies that were significantly greater than those of the respective vehicle studies (Figure 3). The ACE inhibitors, lisinopril and ramipril, did not have detectable cardioactive effects on hvCTS contractility, consistent with the results of other platforms (Harmer et al., 2012; Scott et al., 2014). Therefore, the ACE inhibitor class was removed from the library resulting in a four-class system. Mibefradil, norepinephrine, ouabain, and cisapride were chosen to represent the unknown compound group on which the model would make *de novo* predictions. Mibefradil and cisapride were of particular interest because both compounds had received market approval and were subsequently withdrawn (Li et al., 2016). Nifedipine, isoproterenol, and digoxin were chosen to represent the Ca^{2+} channel blocker, adrenergic agonist, and cardiac glycoside classes, respectively. As both flecainide and E-4031 have known hERG K⁺ channel-blocking capabilities, the impact of having either compound represent the hERG K⁺ blocker family was evaluated by generating the multiclass model under three different conditions: (1) flecainide only, (2) E-4031 only, and (3) both flecainide and E-4031.

A subset of the data was always withheld from the machine prior to training in each of the runs. This withheld set quantified the generalizability in the models and ensured that overfitting had not occurred. Upon asking the machine to classify these test sets, the multiclass models demonstrated good generalizability by being able







Implementation of binary SVM to determine cardioactive effects of 11 compounds, including: (A) nifedipine ($p \le 0.0063$; n = 10), (B) mibefradil ($p \le 0.0063$; n = 6), (C) isoproterenol ($p \le 0.0055$; n = 10), (D) norepinephrine ($p \le 0.0055$; n = 8), (E) digoxin ($p \le 0.0055$; n = 9), (F) ouabain ($p \le 0.0055$; n = 10), (G) flecainide ($p \le 0.0055$; n = 8), (H) E-4031 ($p \le 0.0071$; n = 8), (I) cisapride ($p \le 0.0055$; n = 9), (J) lisinopril ($p \le 0.0055$; n = 8), and (K) ramipril ($p \le 0.0055$; n = 7). Red circles indicate statistical significance in comparison with vehicle-treated hvCTSs. p values are adjusted with a Bonferroni correction. *n* refers to independent biological replicates. All results are presented as means \pm SD.

to correctly classify itself at an average accuracy rate of $76.09\% \pm 6.43\%$, $78.29\% \pm 5.34\%$, and $73.61\% \pm 5.19\%$ for the flecainide only, E-4031 only, and flecainide and E-4031 conditions, respectively (Figure 4).

In all three conditions, the multiclass models behaved similarly in that both the nifedipine and isoproterenol classifiers performed the best by always achieving the highest F_1 score values, a metric that ranges from 0 to 1 with 1 rep-

resenting perfection in the model's classification. This performance indicates that the data points of the nifedipine and isoproterenol compounds occupied very distinct boundaries compared with the other two classes, allowing for the binary learners to more accurately separate these compounds from others. For perspective on the quality of model performance, if there were no discernable differences among the four compounds for the machine to use,

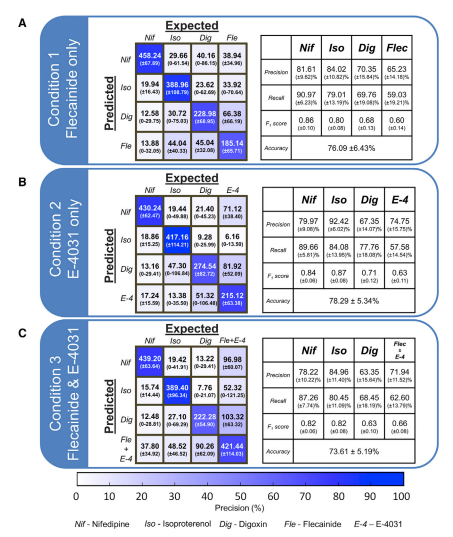


Figure 4. Generalizability of Drug Classification Models with Varying Representative Compound(s) for the hERG K⁺ Channel Blocker Family

(A) Condition 1: flecainide. Generalizability was evaluated by having the model classify a withheld test set composed of compounds used to define the library. (Left) The confusion matrix displays the average number of classified contractile events (individual twitches in acquired waveform) over 50 runs and is imposed with a color scale that indicates precision rate. (Right) Summary of the metrics evaluating performance. F_1 scores are above 0.6 for all classifiers, indicating good predictability.

(B) Condition 2: E-4031. (Left) Confusion matrix indicates that all compounds were being correctly classified as themselves as the diagonal of the matrix had the highest precision rate. (Right) Metrics indicate a similar performance between conditions 1 and 2.

(C) Condition 3: flecainide and E-4031. (Left) Confusion matrix indicates no major effect on generalizability by having two compounds define a class. (Right) Similar to conditions 1 and 2, F₁ scores are all above 0.6, indicating good generalizability of the model.

All results are presented as means \pm SD.

the expected values for precision (i.e., positive predictive value), recall (i.e., sensitivity), and accuracy would be a rate of 25% with an F_1 score of 0.25 (see Experimental Procedures). As all three multiclass models demonstrated good generalizability with average accuracy rates exceeding 70%, these results suggest the setup of the model was robust to the choice of compound representing the hERG K⁺ channel blocker family.

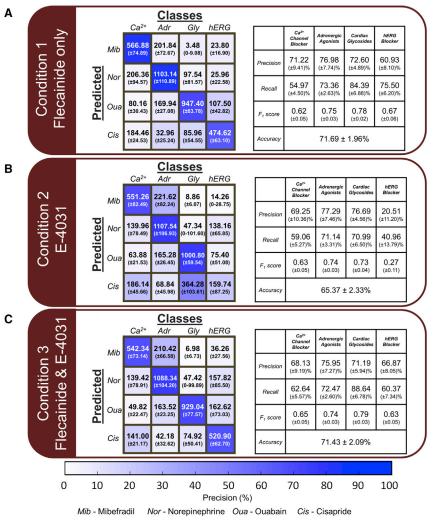
Prediction on Unknown Compounds

With each condition's model established and evaluated, the machine was then asked to predict the data from the unknown compounds group. In the first scenario with flecainide as the only hERG K⁺ channel blocker representative, the multiclass model was able to correctly assign the four unknown compounds to their corresponding counterparts with an average accuracy of 71.69% \pm 1.96% (Figure 5A). The four binary classifiers of this model demon-

strated overall predictability by all having an average F_1 score above 0.6.

When the second drug class model (only E-4031 defining the hERG K⁺ channel blocker class) was used to predict the unknown compounds, the average accuracy diminished to $65.37\% \pm 2.33\%$ (Figure 5B). This decrease was mainly the result of the contractile events from hvCTSs exposed to cisapride being miscategorized. Of the 779 contractile events from hvCTSs exposed to cisapride, on average, 46.76% ± 7.00% of the events were incorrectly labeled as hvCTSs affected by a cardiac glycoside. These misclassifications yielded a poor precision rate, $20.51\% \pm 11.20\%$, and subsequently, a low F_1 score, 0.27 \pm 0.11, for the hERG K⁺ channel blocker classifier. As for the classifiers of the other three families, they performed comparably with those of the first condition, and all had an average F₁ score over 0.6. These results indicate that using only data of hvCTSs exposed to E-4031 is not sufficient for defining the hERG K⁺ channel





Ca²⁺ - Ca²⁺ channel blockers Adr - adrenergic Agonists hERG - hERG K⁺ channel Gly - Cardiac glycoside blocker

Figure 5. Predictability of Drug Classification Libraries with Varying Representative Compound(s) for the hERG K⁺ Channel Blocker Family

(A) Condition 1: flecainide. Libraries of all three conditions were evaluated for capabilities to predict drug families of compounds previously unseen by the machine. (Left) Confusion matrix displays the average number of predicted contractile events (individual twitches in acquired waveform) from the unknown group over 50 runs. Unknown compounds were classified to the correct drug family as the precision rate was highest along the diagonal of the matrix. (Right) The model demonstrated good predictability as the macro-average of the F_1 score was 0.71. (B) Condition 2: E-4031. (Left) Cisapride was predominantly mislabeled as a member of the cardiac glycoside family, represented by digoxin. The other three compounds were classified correctly to their respective drug families. (Right) The classifier of the hERG K⁺ channel blocker class had an F₁ score of 0.27. As the other three classifiers maintained an F₁ score above 0.6, E-4031 did not define boundaries similar to that of cisapride.

(C) Condition 3: flecainide and E-4031. (Left) Precision rates are highest along the diagonal of the matrix. (Right) The model yielded good predictability and was not affected by the inclusion of E-4031 data to those of flecainide.

All results are presented as means \pm SD.

blocker family within a multiclass model and enabling that model to correctly predict cisapride's mechanistic action on cardiac tissue contractility.

In the last condition where the machine was trained with both flecainide and E-4031 representing the hERG K⁺ channel blocker family, the average accuracy was 71.43% \pm 2.09% (Figure 5C). All four classifiers had an average F₁ score higher than 0.6. Unlike the second condition, the hERG K⁺ channel blocker classifier had a precision rate higher than 25% at 66.87% \pm 8.05%. Similar to that of the first condition, the macro-average of the F₁ scores for this multiclass model was 0.71, demonstrating good predictability. These results suggest that the data points from the hvCTSs exposed to flecainide designated boundaries within the model that were similar to those of cisapridetreated hvCTSs. Finally, the inclusion of E-4031 data with the flecainide data did not negatively affect the multiclass model's ability to correctly predict cisapride's mechanistic action as neither the precision rate nor the recall rate fell below 41% as they did in the second condition.

Class Relationship Metrics

Once the drug classes were predicted, the concentrations of library compounds that induced the most similar cardioactive effects as the unknown compounds were computed (Figure S2; protocol in Supplemental Experimental Procedures). These relationship metrics are summarized in Table S1. For example, an estimated 5.35×10^{-5} M digoxin would be needed to evoke a level of cardioactivity that matches ouabain tested at 1.0×10^{-5} M. Such relationships could provide insights about drug potency. In the aforementioned example, ouabain would be considered the more potent compound as it requires approximately 5-fold lower concentration to achieve the same level of



cardioactivity. Ouabain's higher potency has been observed in other *in vitro* studies (Guo et al., 2011; Katz et al., 2010).

Decoupling Force-Frequency Relationship

While the concept of examining multiple parameters from waveforms has been pursued lately, some studies have suggested that only a few select parameters (e.g., peak count) are necessary in assessing a compound's cardioactivity as other parameters provide no further mechanistic insight (Lu et al., 2015; Pointon et al., 2016; Sirenko et al., 2013). This is primarily true when the hPSC-CMs are spontaneously beating, meaning the force generated is linked to beating frequency. This study's dataset affirmed the importance of decoupling this force-frequency relationship through the pacing of the tissues. By setting a fixed pacing frequency, any changes to the force waveform can be truly accredited to a compound's inotropic and lusitropic effects. For example, if the nifedipine-treated strips were allowed to spontaneously beat, a positive chronotropic effect would have most likely been observed (Guo et al., 2011; Harris et al., 2013; Pillekamp et al., 2012). As the hvCTSs displayed a negative force-frequency relationship (Figure S3A), a decrease in maximum developed force could not be directly linked to either chronotropic or inotropic effects. When 0.3 µM nifedipine-treated hvCTSs were paced at 1 Hz, the captured frequency of the tissue was 0.99 \pm 0.01 Hz and the maximum developed force decreased by $45.69\% \pm 10.42\%$ (Figure S3B). This paced data confidently confirmed nifedipine had a negative inotropic effect.

Examination of Cardioactive Effects

The data were further examined on an individual parameter basis to better comprehend and confirm the performance of the multiclass models and their ability to differentiate between different mechanistic actions (Figure S4). The adrenergic agonists and cardiac glycosides were expected to induce a positive inotropic response in the hvCTSs, while the Ca²⁺ and hERG K⁺ channel blockers would induce a negative inotropic response. The negative inotropic agents prompted distinct decreases in the maximum force generated among hvCTSs; however, the hvCTS sensitivity to positive inotropic agents was not very apparent. For example, hvCTSs exposed to 10 µM isoproterenol and paced at 0.5 Hz had a similar increase in maximum developed force to that of respective vehicle-treated strips (10.42% \pm 16.23% and 15.76% \pm 21.05%, respectively), suggesting the compound had negligible inotropic effects (Figures S5A and S5B) (Turnbull et al., 2014). Rather, isoproterenol's cardioactive effects manifested in other parameters. For example, the duration of the relaxation phase or time from maximum developed force to 95% cutoff decreased by 22.19% ± 19.35% for

the strips exposed to 10 μ M isoproterenol, while those of the vehicle-treated strips experienced essentially no change. Thus, through the various parameters, machine learning was able to leverage the positive lusitropic effects in both binary and multiclass SVM to distinguish the adrenergic agonist class.

Cardiac glycoside-treated hvCTSs also demonstrated the system's sensitivity to positive inotropes. Typically, these compounds increase the Ca²⁺ transient amplitude and amplitude of contractility transients; however, in this dataset, the hvCTSs decreased in maximum developed force as the concentration increased (Figure S5C) (Dempsey et al., 2016; Ravenscroft et al., 2016). Such results could be attributed to cardiac glycoside toxicity or immature phenotype. Studies have shown that above 3 μ M digoxin or ouabain, monolayers of hPSC-CMs stopped beating (Dempsey et al., 2016; Guo et al., 2013; Sirenko et al., 2013). In this dataset, the highest concentration applied for both cardiac glycosides was 100 μ M. At 100 μ M ouabain, 4 of the 10 treated strips stopped beating at all pacing frequencies.

Like those of isoproterenol, cardioactive effects of the cardiac glycosides at lower concentrations appeared in other parameters. When either the concentration of cisapride or digoxin increased, the maximum developed force decreased while the duration of the relaxation phase increased (Figures S5C and S5D). If limited to observation of only these two parameters, one might conclude that cisapride and digoxin are related. But upon visual inspection of the force traces, it is evident that the compounds have distinct effects on hvCTSs and mechanistic actions (Figures S3E and S3F). Changes in these parameters were clearly unique to the cardiac glycoside family as the ouabaintreated strips were correctly predicted from the other three classes.

DISCUSSION

In recognition of the need for better detection of druginduced cardiotoxicity, numerous methodologies have emerged to capture and quantify the attributes of hPSC-CMs when exposed to cardioactive compounds, ranging from calcium transients to contractile force. The nature of the output data becomes multidimensional when multiple experimental conditions are present or a multiplex system is used (Dempsey et al., 2016). In this study, we present the use of supervised machine learning to exploit multidimensional data and provide relevant information in an automated manner. Besides indicating if a compound was cardioactive, the machine constructed a multiclass drug model that accurately classified cardioactive compounds that it had never previously encountered. This comprehensive



approach can be readily applied to other screening platforms to more fully utilize generated datasets and enhance evidence-based decision-making for drug development.

With multiclass SVM, drug classification libraries were established under various conditions to examine the effects on predictive performance. The conditions that yielded the best performance in predicting mechanistic action were the two libraries that included flecainide as a representative of the hERG K⁺ channel blocker family. In both libraries, the macro-averages of F₁ scores were 0.71 (the macro-average of F₁ scores would be 0.25 if random classifiers were used). While this clear difference in F₁ scores indicates that the models have the capability to predict a compound's mechanistic action, there are opportunities to further improve model performance and obtain F₁ scores closer to 1, indicating reduction in errors.

One method to improve model performance is to define each drug family with multiple compounds. By having only one compound define a class, there is a risk of only defining a partial region of space that the drug class truly encompasses. The data of E-4031 exemplified this when it was tasked with defining the hERG K⁺ channel blocker family. E-4031's defined boundaries did not match or include those of cisapride, another hERG K⁺ channel blocker, causing classification of cisapride to be closer to that of the cardiac glycoside family. The inclusion of flecainide, a mixed hERG K⁺ channel blocker, with E-4031 in the definition of the class allowed for the correct prediction of cisapride without adversely affecting the predictive capability of the remaining classes. Although the addition of E-4031 to the hERG K⁺ channel blocker definition does not necessarily improve the predictive capability with respect to cisapride classification, establishing a more expansive region of space to define the hERG K⁺ channel blocker class may improve prediction of other unknown hERG K⁺ channel blockers that have effects more similar to E-4031 than flecainide. These results also suggest the potential of having subgroups within classes of the model, which can be achieved through a series of multiclass classifications. For instance, a compound can be predicted as a Ca²⁺ channel blocker in the first classification; within this family, the compound can be subsequently categorized into a subgroup (e.g., defined by frequency-dependent cardioactivity). As machine learning does not define drug classes with a priori knowledge (e.g., guidelines on how parameters are expected to change), the number of drug families and subclasses that can be defined within a model are not limited. The unbiased and automated nature of machine learning is also advantageous when a new drug family needs to be added, because no rubric needs to be manually amended and re-evaluated.

This study demonstrates the potential of machine learning for providing insights in the detection of cardioac-

tivity using hPSC-CMs. The basis of this study's libraries was an error-correcting output codes approach with binary learners being SVM. Different binary learners, such as decision trees, should be explored alongside completely different approaches (e.g., neural networks). The ideal machine learning technique should balance predictive capabilities and use of computational resources. In this study, all models were generated with a standard desktop. Each calculated instance of a model took approximately 4 hr. However, once all models were formed, the predictions made on unknown compounds were on the timescale of seconds.

Improvements of the multiclass drug libraries can also be achieved from enhancements of the hvCTSs and acquisition system. In particular, the sensitivity of this system to positive inotropic compounds can be increased by addressing two issues: the maturity of stem cell-derived cardiomyocytes and the drifting baseline of vehicle-treated strips. Studies have shown that hPSC-CMs elicit a minimal to non-existent response to certain positive inotropic compounds, such as β-adrenergic agonists, because of immature intracellular structures (Lundy et al., 2013; Pillekamp et al., 2012). When these diminished responses are paired with a baseline that has increasing contractility over time, positive inotropic effects of a compound can get masked and become harder to detect as seen in the aforementioned isoproterenol drug screen. While hvCTSs were arranged in an aligned manner and co-cultured with fibroblasts, they can be further matured through additional techniques, such as conditioning by electrical stimulation, a cellular tri-culture including endothelial cells, or forced expression of selected proteins (Eng et al., 2016; Liu et al., 2009; Ravenscroft et al., 2016). As for stabilization of the baseline, different components of the setup, ranging from pH to CO₂ levels in an ambient environment, should be re-evaluated to minimize overall drift during serial additions. Increasing the system's sensitivity to positive inotropic agents would yield even more distinct boundaries and subsequently better predictability in the drug classification libraries.

While we demonstrated the importance of pacing hvCTSs, a limitation of this study is the absence of responses of spontaneously beating hvCTSs to cardioactive compounds. These data would provide understanding of a compound's chronotropic effects. Even though the model has already demonstrated good predictability on inotropic- and lusitropic-related data alone, the model would benefit from chronotropic-related data as it would provide more dimensions in which compounds can further distinguish themselves and yield better predictability.

In summary, we present the implementation of supervised machine learning on multidimensional data of



hvCTSs exposed to drugs while paced at various frequencies. In an automated fashion, this machine learning approach is able to not only determine if a compound is cardioactive but it can predict the mechanistic action along with other metrics. Furthermore, this approach can be adapted to state-of-the-art tissue-engineered cardiac models, including different forms of signals (e.g., calcium transients, micro-electrode array, and optical recordings), and has the potential to integrate diverse output data of multiplex systems or even those across platforms. Along with analyses of compounds with acute cardioactive effects, machine learning can be readily applied with non-invasive techniques to longitudinal studies to inspect a compound's chronic effects. Moreover, in the future, machine learning may be utilized on a grander scale by incorporating past clinical data to determine the optimal combination of *in vitro* and *in silico* data for the prediction of drug-induced cardiotoxicity in patients.

EXPERIMENTAL PROCEDURES

hvCTS Formation

Human ventricular cardiomyocytes were differentiated from an hES2 stem cell line with a Wnt inhibitor-based protocol as described previously (Weng et al., 2014). Human ventricular cardiac tissue strips (hvCTS) were then formed by mixing cardiomyocytes (100,000 cells per strip) at 14-16 days post differentiation with a solution of bovine collagen I (2 mg/mL), Matrigel (0.9 mg/mL), and human foreskin fibroblasts (100,000 cells per strip) as previously described (Turnbull et al., 2014). The cell-matrix solution (100 µL per tissue strip) was injected into a custom polydimethylsiloxane (PDMS) force-sensing bioreactor device and placed in an incubator (37°C and 5% CO₂). Formed hvCTSs were fed DMEM with 10% newborn calf serum, 1% penicillinstreptomycin, and 0.1% amphotericin B. The PDMS device contains two flexible vertical end posts to which the tissue anchors, causing the posts to deflect as the tissue beats. Contractile force measurements were captured with a high-speed (100 fps) CCD camera while custom LabVIEW software tracked the centroid movement of the flexible post tips. Force was converted from the deflection of the PDMS posts by an elastic beam-bending equation (Serrao et al., 2012). A custom MATLAB script was used to calculate 17 parameters that described the overall shape of the force traces for each contractile event (Figure S1). Each contraction was regarded as an individual data point for the machine learning analysis.

Drug Treatment

Seven to eight days post tissue formation, hvCTS were exposed to drugs for pharmacodynamic analysis. Flecainide, lisinopril, norepinephrine, and ramipril were provided by Pfizer, while all other compounds were purchased form Sigma-Aldrich. Compounds were initially resuspended in DMSO and subsequently diluted in water for final concentrations of less than 0.1% (vol/vol) DMSO. The PDMS device containing the hvCTS was placed onto a heated stage $(37^{\circ}C)$ under a dissecting microscope. Before either vehicle or drug addition, the medium was replaced with DMEM containing high glucose (4.5 g/L) and HEPES without phenol red. Drug doses were added to a tissue in a consecutively increasing manner up to 10 concentrations with 3 min between measurements. Vehicle doses containing only water were applied similarly. A pulse stimulator (AMPI Master-9) connected to platinum wires electrically paced the hvCTSs with a monophasic electric field of 5 V/cm and a 10 ms pulse duration.

Machine Learning

To establish the drug class model, we identified individual compounds that respectively represented our defined classes. The compounds and the corresponding tested concentrations are listed in Table 1. To determine which concentration of a chosen compound to add to the model, we first gauged each compound's level of cardioactivity by utilizing binary SVM (Figure S6) (Lee et al., 2015). To normalize for the increasing variation seen in hvCTS contractile behavior during the later serial additions of the vehicle studies, the SVM was performed between each concentration of a compound and the vehicle data from the corresponding serial addition number in which the concentration was achieved (Figure 1A). Specific details and rationale regarding the optimization of SVM classifiers and other implemented machine learning approaches are in the Supplemental Experimental Procedures.

For multiclass classification, we then selected the compound concentration that met two criteria: (1) a binary SVM accuracy closest to 85% and (2) at least six of all screened tissue strips were still responsive to electrical stimulation (see Supplemental Experimental Procedures). As seen in Figure 1B, we then divided the compounds into two groups: one used to train the multiclass model and one to represent unknown compounds for predictions. During this separation, it was ensured that each class was at least represented by one compound within the two groups. For training of the model, we used an error-correcting output codes approach with the binary learners being SVM (Dietterich, 1995). To confirm generalizability of the generated models, we randomly pre-allocated a third of the data as a test set prior to the training. Finally, we evaluated the multiclass model with this test set and then asked the machine to predict the classes of the contractile beats derived from the unknown compounds group.

Statistics

SVM accuracies of strips exposed to a drug condition were compared with those of the non-cardioactive benchmark by using Student's t test (desired α value of 0.05) with a Bonferroni correction (*m*, number of tests or hypotheses, was dependent on the number of drug additions in a screen). If the adjusted p value was statistically significant, the drug condition was considered to have incited irregular behavior in hvCTSs and was labeled as cardioactive. The Bonferroni correction was also applied when examining changes in specific parameters.

To analyze the performance of the multiclass models, confusion matrices were generated for each of the 50 runs. In a confusion matrix, *M*, the precision and recall rates were defined as follows:

$$Precision_i = \frac{M_{ii}}{\sum_{j=1}^n M_{ij}}, \ Recall_j = \frac{M_{ii}}{\sum_{i=1}^n M_{ij}} \ .$$

The precision and recall rates were calculated for each of the classifiers. To further summarize these metrics, the F₁ score, the harmonic mean of precision and recall, was computed and defined as follows:

$$F_1 \ score_i = \frac{2 \times Precision_i \times Recall_i}{Precision_i + Recall_i}$$

A model that is perfect would achieve an F_1 score of 1. If a model were composed of *s* number of classes and had random classifiers, the expected F_1 score would be 1/s. To assess the model as a whole, accuracy, defined as

$$Accuracy_{model} = \frac{\sum_{i=1}^{n} M_{ii}}{\sum_{i=1}^{n} \sum_{j=1}^{n} M_{ij}}.$$

was calculated. In summarizing the 50 runs of each model, all calculated metrics were averaged and a confusion matrix containing the average number of contractile events over all runs was provided. Contractile events were defined as the individual twitches in the force readouts acquired from the tissue strips. All reported sample sizes (n) refer to independent tissue strips (biological replicates). All descriptive statistics are in the format of mean \pm SD.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, six figures, one table, and one data file and can be found with this article online at https://doi.org/10.1016/j.stemcr. 2017.09.008.

AUTHOR CONTRIBUTIONS

E.K.L. and D.D.T. initiated and conceived project. G.W., C.W.C., K.D.C., R.A.L., and M.K. supervised the project. W.K. and P.C. collected biological data. E.K.L. implemented the machine learning concepts. E.K.L., D.D.T., W.K., K.D.C., and M.K. analyzed data. E.K.L. and D.D.T. wrote the manuscript. W.K., K.D.C., R.A.L., and M.K. contributed to the writing of the manuscript.

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