Selecting anti-HIV therapies based on a variety of genomic and clinical factors

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ABSTRACT

Motivation: Optimizing HIV therapies is crucial since the virus rapidly develops mutations to evade drug pressure. Recent studies have shown that genotypic information might not be sufficient for the design of therapies and that other clinical and demographic factors may play a role in therapy failure. This study is designed to assess the improvement in prediction achieved when such information is taken into account. We use these factors to generate a prediction engine using a variety of machine learning methods and to determine which clinical conditions are most misleading in terms of predicting the outcome of a therapy.

Results: Three different machine learning techniques were used: generative-discriminative method, regression with derived evolutionary features, and mixture with regression and a mixture of weights. All three methods had similar performances with an area under the receiver operating characteristic curve (AUC) of 0.77. A set of three similar engines limited to genotypic information only achieved an AUC of 0.75. A straightforward combination of the three engines consistently improves the prediction, with significantly better performance than when the full set of features is employed. The combined engine improves on predictions obtained from an online state-of-the-art resistance interpretation system. Moreover, engines tend to disagree more on the outcome of failure therapies than regarding successful ones. Careful analysis of the differences between the engines revealed those mutations and drugs most closely associated with uncertainty of the therapy outcome.

Availability: The combined prediction engine will be available from July 2008, see http://engine.euresist.org

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1 INTRODUCTION

1.1 Treating HIV patients

Most of the antiretroviral compounds used to treat HIV patients belong to one of three classes: protease inhibitors (PI), nucleotide reverse transcriptase inhibitors (NRTI) and non-nucleotide reverse transcriptase inhibitors (NNRTI). Mutations in the target protein confer resistance to the drug targeting that protein. Although various HIV subtypes exist, our study focuses on subtype B, which is prevalent in Europe. Today, highly active antiretroviral therapy, known as HAART, is commonly used for treating HIV patients. A typical HAART comprises three to four carefully selected drugs from at least two different drug classes. This approach is designed to minimize the probability that the virus will develop escape mutations.

1.2 The EuResist dataset

The EuResist integrated database (IDB) combines the ARCA database (Italy),\textsuperscript{4} AREVIR database (Germany; Roopp et al., 2006) data coming from the Karolinska Infectious Diseases and Clinical Virology Department (Sweden), and a smaller dataset from Luxembourg. The EuResist database comprises records of 18K different patients and 65 K different therapies, of which 3K therapies contain genotype information. The information for the 18K different patients includes demographic data records such as gender, race and age, with the anonymization of patients data carried out before the data is stored in the IDB. Clinical measures of the disease state, such as viral load (VL) and CD4+ cell counts, are collected for each patient and for some of the therapies around a treatment switch. Sequence information of HIV protease (PRO) and reverse transcriptase (RT) obtained from genotypic resistance tests are also provided. A standard datum (SD) format was designed to define the characteristics of successful and failing therapies, respectively (minimal treatment length, handling of overlapping therapies, etc.). The SD format is also used to define which genotype and VL tests serve as baseline measures and are therefore associated with a therapy switch, and which follow-up VL is associated with a therapy outcome (8 weeks from the start of a therapy). Each therapy with a baseline and a follow-up VL is labeled a success if there is a drop of 2 log in VL or a drop below detection level (500 copies per milliliter); otherwise, the therapy is labeled as a failure. This process of generating a standard datum was motivated by the need to develop an online engine that performs as a decision support system and can recommend a drug combination for a patient. To this end, a minimal requirement for a prediction engine is defined containing the genotypic sequence for the two viral target proteins; the remaining information is treated as optional or maximal set of features.

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\textsuperscript{4}1http://www.hivarca.net/
1.3 Mutations
The HIV virus has a high mutation rate. In principle, all (about 10,000) positions in the viral genome are candidates for mutations. In particular, the related 99 amino acid positions of the PRO and the 440 amino acid positions of the RT have high mutation rates. This leads to an exceedingly high dimension. A prediction engine taking the whole relevant protein sequence into account is likely to suffer from the curse of dimensionality. One possibility of circumventing this problem is to make use of a predefined list of mutations expected to have the most significant effect on the therapy’s outcome. The International Aids Society (IAS) maintains a constantly updated reference mutation list, known as IAS mutations, of the mutations of PRO and RT sequence (and additional new viral target proteins) that are known to play a role in drug resistance (Johnson et al., 2007). Indeed, if we count the number of such mutations in our data, separating failure therapies from successful ones, the baseline genotype of failing therapies has on the average 3.1 ± 1.4 IAS mutations, while successful therapies have 2.2 ± 1.2 (Fig. 1). Finally, we find a highly significant correlation between the number of IAS mutations and successful therapies (using an indicator variable with 1 for successful therapy and 0 for failure), correlation of −0.32 (with P-value < 10−50). In our prediction engines described below, the list of IAS mutation was a natural candidate feature and in some of the engines this is the set of features picked for representing the protein sequences.

1.4 Related work
As soon as resistance to antiretroviral drugs was discovered, the need for tools to detect drug resistance arose. The majority of available interpretation systems (IS) work with hand-crafted tables (such as the IAS list) for every drug compiled by experts. These tables are used to classify the virus as susceptible or resistant to a single drug. These rules are built using expert knowledge from clinical experience and recently they incorporate experimentally generated-genotype-phenotype information. Some IS are completely data-driven rather than based on expert knowledge. For example, geno2pheno (Beeenwinkel et al., 2003) and VirtualPhenotype (Vermeiren et al., 2007) apply machine learning methods to predict in vitro drug resistance from the viral genotype. Recently published (data-driven) tools are able to predict in vivo response to a combination of antiretroviral drugs rather than single compounds. In Larder et al. (2007), committees of Artificial Neural Networks are used to predict the change in VL after treatment start, apart from information about the viral sequences and the intended regimen, CD4+ counts, baseline VL and four treatment history indicators that are used as input features. This approach was trained and validated on 1150 and 100 treatment change episodes respectively. Geno2pheno–THEO by Altmann et al. (2007), another online tool, predicts the success probability of a putative treatment based on sequence information and the estimated genetic barrier of the viral variant to resistance against every drug in the combination therapy. However, geno2pheno–THEO does not make use of information about the patient or other available clinical markers. In comparison to these two systems, our approach is able to predict the success probability of a regimen as well as the change in VL. Most important, our prediction system makes use of the fact that it employs three different highly optimized individual systems to provide a more robust prediction, i.e. one with a smaller variance in prediction accuracy and AUC scores, allowing us to gain further insight in the treatment–outcome relationship as this study demonstrates.

2 MATERIALS AND METHODS
We trained three separate engines. Each can be viewed as being composed of two layers, with a layer of feature generation and selection embedded within a discriminative method such as logistic regression, support vector machines (SVMs) or random forests. As logistic regression, compared with the other methods, requires much less parameter tuning efforts and performs similarly to the other two in the tests we carried out, we use logistic regression for the top layer and the engines differ in the type of features derived. We use the following three approaches:

1. Evolutionary models
2. Generative models
3. Mixture of effects

Given a core set of features selected by one of the methods above, we devoted extensive studies to figure out what other features, and in what format, should be added to the core set of extracted features to achieve a separate good performing engine. The result is a set of three engines—evolutionary engine (EV), a generative discriminative (GD) engine and a mixture of effects engine (ME).

Each of the engines is trained twice, once with the restriction that only drugs in the regimen and sequence information about the two viral targets are available (minimal feature set) and once with no such restriction (maximal feature set). In addition to these six engines aimed at binary classification, the prediction of the label of a therapy (success/failure), we train three engines with a target of predicting the drop in VL, i.e. regression (the minimal feature set is less adequate for this target of predicting change of VL as it requires

![Fig. 1. Number of IAS mutations in HIV genotypes submitted to drug therapies.](image-url)
All features defined in the SD, except for current and past drugs and genotype, are shown in Table 1. The table shows the minimal and maximal values for each of the features, the correlation with the therapy label, and the related \( P \)-value. We also apply the Kolmogorov–Smirnov (KS) test to check which features are distributed differently in the failure therapies and in the successful ones (second \( P \)-value in the parenthesis). In cases where the feature is categorical we use the \( \chi^2 \)-test instead. All \( P \)-values are adjusted with Bonferroni correction. The three measures—correlation, its related \( P \)-values and a \( P \)-value for rejecting the null hypothesis that there is no association between the values of the feature and success/failure—are derived for all therapies in the training set/incomplete labeled set (third/fourth column, respectively). The features are sorted by correlation in ascending order. The first item is the number of past treatment lines; the more therapy switches in the patient’s history, the more likely the current therapy will fail. The second, ‘RISKID’ stands for the infection risk group; this seems to be an indicator of whether a therapy will succeed or not. Although medically there is no reason that the manner of infection will impact response to treatment per se, it is known that intravenous drug abusers can experience less effective therapy due to lower adherence. Gender IDs stand for male/female/undifferentiated/unknown; it does not seem to be directly related to the success/failure of a therapy.

2.1 Raw features in the IDB

All features defined in the SD, except for current and past drugs and genotype, are shown in Table 1. The table shows the minimal and maximal values for each of the features, the correlation with the therapy label, and the related \( P \)-value. We also apply the Kolmogorov–Smirnov (KS) test to check which features are distributed differently in the failure therapies and in the successful ones (second \( P \)-value in the parenthesis). In cases where the feature is categorical we use the \( \chi^2 \)-test instead. All \( P \)-values are adjusted with Bonferroni correction. The three measures—correlation, its related \( P \)-values and a \( P \)-value for rejecting the null hypothesis that there is no association between the values of the feature and success/failure—are derived for all therapies in the training set/incomplete labeled set (third/fourth column, respectively). The features are sorted by correlation in ascending order. The first item is the number of past treatment lines; the more therapy switches in the patient’s history, the more likely the current therapy will fail. The second, ‘RISKID’ stands for the infection risk group; this seems to be an indicator of whether a therapy will succeed or not. Although medically there is no reason that the manner of infection will impact response to treatment per se, it is known that intravenous drug abusers can experience less effective therapy due to lower adherence. Gender IDs stand for male/female/undifferentiated/unknown; it does not seem to be directly related to the success/failure of a therapy.

Note that the two bottom rows in the table with the highest correlation defined as optional features in the SD. One row stands for the incomplete labeled set in which the drug occurs. We derive the success rate as obtained from the training set (and as obtained from the incomplete labeled set) for cases where that drug is administered ('SR with', second column) and for cases where the therapy does not contain the specific drug ('SR without', third column). The consistent lower success rate of the training set compared with the incomplete labeled set is due to the bias by which the training set is selected—therapies that present a greater challenge than usual and hence genotypic sequence is tested. We also provide \( P \)-values of \( \chi^2 \)-tests of whether the success/failure rate of a therapy is similar when the drug is part of the HAART and when it is missing. \( P \)-values lower than 0.05 are provided after adjustment of Bonferroni correction.

2.2 Evolutionary engine

This engine focuses on the development and use of evolutionary features. As stated before, one major obstacle in HIV-1 treatment is the virus’s escape from drug therapy by developing resistance mutations. In order to accurately predict the outcome of an antiretroviral therapy information about viral evolution has to be presented to the underlying statistical learning method. Our representation of the viral evolution is based on mutagenetic trees. Briefly, the mutagenetic tree is reconstructed from all pairwise probabilities of defined events. Here, these events are occurrences of drug resistance mutations in the viral genome. Hence, a mixture of reconstructed mutagenetic trees represents possible resistance pathways along with probabilities for the development of the participating mutations. Using these trees we can compute the so-called genetic barrier to drug resistance by calculating the probability of the virus not to develop further mutations leading to a complete phenotypic resistance to that drug. A complete resistance pattern is observed if on average viruses with that mutational pattern exceed a certain level of phenotypic drug resistance. Here the cut-offs used by geno2pheno (Beerenwinkel et al., 2003) were applied. In previous work Altmann et al. (2007) could show that using the genetic barrier significantly improves the

Table 1. IDB data fields

| Field                        | [min max] | Corr (training data) | Corr (full labeled set) |
|------------------------------|-----------|-----------------------|-------------------------|
| NUMBER OF PAST TREATMENT LINES | [0, 28]  | -0.26 (8.0 × 10^{-42}, 1.1 × 10^{-27}) | -0.15 (0, 0) |
| RISKID                       | [1, 6]    | -0.04 (--, 4.3 × 10^{-4}) | -0.03 (1.3 × 10^{-3}, 0) |
| GENDERID                     | [1, 4]    | -0.02 (--, --) | 0.00 (0, --) |
| AGE                          | [10, 102] | 0.01 (--, --) | 0.04 (1.0 × 10^{-7}, 5.6 × 10^{-15}) |
| BASELINE VL                  | [34, 7656900] | 0.01 (--, 6.8 × 10^{-15}) | 0.01 (--, 0) |
| BASELINE CD4                 | [0, 4441] | 0.06 (--, --) | 0.15 (0, 0) |
| BASELINE CD4 PERCENT         | [0, 70]   | 0.06 (--, 9.0 × 10^{-3}) | 0.17 (0, 0) |
| ALL TREATMENTS RECORDED       | [0, 1]    | 0.10 (9.3 × 10^{-5}, 5.4 × 10^{-3}) | 0.12 (7.2 × 10^{-4}, 0) |
| DATABASEID                   | [1, 4]    | 0.12 (7.4 × 10^{-5}, 1.5 × 10^{-3}) | 0.17 (0, 0) |

\( P \)-values are provided in parenthesis only if they are below 0.05; -- stands for higher values.

\( a \)-Categorical data, the \( \chi^2 \)-test, second in parenthesis, is more informative.
Table 2. Drugs and the associated success rate

| Drug   | Class | Occ  | SR with | SR without | \( \chi^2 \) -test |
|--------|-------|------|---------|-------------|-------------------|
| DDC    | NRTI  | 28 (343) | 0.36 (0.43) | 0.67 (0.69) | – (0) |
| APV    | PI    | 77 (344)  | 0.36 (0.40) | 0.68 (0.69) | 2.8 \times 10^{-6} (0) |
| SQV    | PI    | 195 (1560) | 0.50 (0.49) | 0.68 (0.70) | 4.4 \times 10^{-3} (0) |
| RTV    | PI    | 130 (1035) | 0.56 (0.57) | 0.67 (0.69) | – (1.1 \times 10^{-3}) |
| D4T    | NRTI  | 715 (5949) | 0.57 (0.60) | 0.70 (0.72) | 4.6 \times 10^{-6} (0) |
| NFV    | PI    | 235 (2329) | 0.57 (0.65) | 0.68 (0.69) | – (–) |
| IDV    | PI    | 193 (2322) | 0.58 (0.67) | 0.68 (0.69) | – (–) |
| DDI    | NRTI  | 864 (4943) | 0.59 (0.60) | 0.70 (0.71) | 1.2 \times 10^{-3} (0) |
| NVP    | NNRTI | 308 (2221) | 0.63 (0.68) | 0.67 (0.69) | – (–) |
| ABC    | NRTI  | 484 (3755) | 0.67 (0.75) | 0.67 (0.67) | – (0) |
| TDF    | NRTI  | 1053 (4711) | 0.70 (0.73) | 0.63 (0.64) | – (9.8 \times 10^{-3}) |
| TDF    | NRTI  | 1676 (13168) | 0.70 (0.71) | 0.63 (0.64) | 2.8 \times 10^{-3} (0) |
| RTV    | PI    | 1600 (6686) | 0.72 (0.74) | 0.61 (0.66) | 6.6 \times 10^{-7} (0) |
| LPV    | PI    | 1108 (4136) | 0.73 (0.73) | 0.64 (0.67) | 1.3 \times 10^{-4} (2.9 \times 10^{-10}) |
| ATV    | PI    | 271 (1643)  | 0.73 (0.81) | 0.66 (0.68) | – (0) |
| EFV    | NNRTI | 526 (3524)  | 0.73 (0.78) | 0.66 (0.67) | – (0) |
| FPV    | PI    | 118 (449)   | 0.74 (0.73) | 0.67 (0.69) | – (–) |
| AZT    | NRTI  | 970 (7536)  | 0.76 (0.74) | 0.63 (0.65) | 1.0 \times 10^{-13} (0) |
| FTC    | NRTI  | 242 (1194)  | 0.80 (0.88) | 0.66 (0.67) | 1.6 \times 10^{-3} (0) |

 zoals stands for \( P \)-value higher than 0.05, 0 stands for \( P \)-value lower than 10^{-3}, and SR stands for the success rate with and without the drug.

prediction of therapy response. Thus, the genetic barrier to drug resistance is computed for every single drug in the putative regimen given the viral sequences, and is together with other features, like indicators for single drugs in the treatment, the viral genotype as represented by indicators for IAS mutations, the treatment history, second-order interactions between these features and the baseline VL input to a logistic regression (linear regression for the regression task). To select the features used in the final engine, a simple feature selection approach based on SVMs was applied. The approach works in three steps: first, the cost parameter of a linear SVM is optimized by maximizing the AUC (correlation for regression) in a 10-fold cross validation setting; second, 25 different linear SVMs are generated by five repetitions of 5-fold cross-validation using the optimized cost parameter; third, all features having a mean \( z \)-score larger than two were kept for use in the final model.

2.3 GD engine

This engine develops and applies generative models of the interactions between current and history antiretroviral drugs. It is well known that subsequent use of previously employed and closely related drugs may be inefficient (see e.g. Pillero, 2003; Siliciano, 2004). Generative models are powerful since they add expert knowledge for guidance of the design of the network is available (Pearl, 1988). In this case we have 20 drugs naturally divided into three classes, PI, NRTI and NNRTI. A number of candidate networks containing nodes representing the drugs individually and/or the drug classes are tested using 10-fold cross-validation with 20 divisions on a large training set. The data available for training is the incomplete labeled set along with the training dataset, a total of about 20000 therapies. The network selected is shown in Figure 2. Root node and all leaves are binary nodes standing for success/failure, existence/not existence of a drug in current therapy, respectively. The three nodes in the middle are discrete nodes that stand for the count of number of history drugs adhered by the patient, a separate count per drug class. Each of these three nodes is a parent to the drugs from the related class. This Bayesian network, with no other features like genotype, yields an AUC of 0.716 \pm 0.001, it outperforms a Naive Bayes network trained with 20 leaves that stand for the drugs (AUC of 0.654 \pm 0.001) and a logistic regression engine trained on the same features (AUC of 0.662 \pm 0.001). The top layer of logistic regression engine trained on clinical genomic data with the 20 drugs, 20/25 mutations selected based on correlation with VL drop of PRO/RT in the baseline genotype and five mutations appearing in history genotype, see the Appendix for more details. The final list of features used in the generative discriminative engine contains, in addition to the above features, the 20 compounds, baseline VL and number of past treatment lines.

In a similar process a generative discriminative engine is developed for the case of the minimal feature set. The best performing Bayesian network is similar to the one in Figure 2, except that the middle layer with history drugs is replaced by an indicator that is 0/1 if the drug belong/does not belong to a drug class. This Bayesian network alone results with AUC of 0.672 \pm 0.001, it outperforms a naive Bayes network trained with 20 leaves that stand for the drugs (AUC of 0.654 \pm 0.001) and a logistic regression engine trained on the same features (AUC of 0.662 \pm 0.001). The top layer of logistic regression engine trained on clinical genomic data with the 20 drugs, 20/25 mutations selected based on correlation with VL drop of PRO/RT results in AUC of 0.744 \pm 0.003. This performance improves when to that the single Bayesian network trained feature is added—AUC of 0.749 \pm 0.003.

2.4 Mixture of effects engine

This engine takes second- and third-order variable interactions into account. The mixed-effects engine explores logistic regression for classification of virological success and multiple linear regression for regression of actual VL change.

Second- and third-order variable interactions were set up taking into account the following mixed effects:

1. drug \times mutations
2. mutations \times mutations
3. drug \times drug
4. drug \times drug \times mutations
5. drug \times drug \times drug
The following are additional features included in the engine: information of past treatments, using either a continuous exponentially decreasing function or a binary indicator for each drug class; epidemiological information (such as risk factor, country of infection, age, sex, …); clinical markers (viral RNA load and CD4+ cell counts) and viral subtypes [assessed through BLAST (Altschul et al., 1990) scores on a reference data base of viral subtypes].

The engine was fed with the clinical, epidemiological and all genomic data (more than 500 mutations), with the mixed effects added (thousands of interaction variables). The modeling required a strong effort in feature selection, since the input attribute space ranged from hundreds to thousands of variables. The feature selection techniques used were: (i) filters, using univariable analysis ($\chi^2$-rank-sum tests) and correlation-based feature selection (CFS; Hall, 1998); (ii) embedded methods, using AIC selection (Akaike, 1974) and ridge shrinkage (Hoerl, 1962; Le Cessie et al., 1992).

For both classification and regression, 10-fold cross-validation was executed multiple times, in order to obtain a Gaussian distribution that could be compared with a $t$-statistic (adjusted for sample overlap and multiple testing), useful in model comparison or model selection (Nadeau et al., 2000). The models were tested using different feature spaces and different loss functions (accuracy, AUC).

2.5 Results

Prediction results of the regression engines are provided in Table 3. Averaged correlation between actual and predicted change in log(VL) as obtained from the 10-fold cross-validation test (and SD) are provided for the training and the test datasets (first two columns); the same partition of 10-folds is used with each of the engines. Means squared error of the predictions are also provided.

Prediction results of the six classification engines are provided in Table 4. AUC and accuracy, i.e. 1-misclassification error, on the training dataset are derived from 10-fold cross-validation tests, SD is provided in parentheses. Results on simple combinations of engines are also provided. Note the consistent decrease in SD when AUC and accuracy measures of combinations of engines are derived, this means that an engine that is a resultant of a combination of the three engines provides a more robust prediction. In Figure 3 we compare the failures and successes of the three engines (training and test data altogether) separately for the case where the therapy failed and for the case where the therapy succeeded. When a single engine is referred as correct (wrong) in the figure it means that the other two are wrong (correct). We find two interesting effects: (a) all engines tend to provide the same wrong answer on failure therapies more than on successful ones and (b) inconsistency among engines on failure therapies is much larger than the cases of successful therapies. We discuss these observations in the next section.

A combined engine is derived by taking the average probability of successful therapy obtained from each of the engines. It seems to have the best performance as measured by accuracy and AUC (Table 3). The accuracy of this combined engine when a maximal feature set used is significantly higher than when a minimal set feature set is used. A paired $t$-test results in a $P$-value of 0.00013/0.048 on training/test dataset, respectively. The $P$-value on the test set is reduced because fewer samples are available.

Furthermore, we compare the performance of the three engines and the combined engine with Stanford’s online available (http://hivdb.stanford.edu) Interpretation System (IS) version 4.2 (De Luca et al., 2006). This system is widely used to classify a virus to be resistant against single drugs. Unlike in other expert algorithms individual scores are assigned to mutations in the genotype. The scores are derived from publications and from genotype–phenotype data. In this way the stanford system can be seen as an expert-derived linear regression model for resistance against single drugs. Comparison to a system that predicts therapy response to a combination of drugs was not possible, since the system by Lander et al. (2007), is not freely available. We extracted the nucleotide sequences for the test data from Table 3.

### Table 3. Log VL prediction results

| Model | Correlation | Mean squared error |
|-------|-------------|--------------------|
|       | Train | Test | Train | Test |
| GD    | 0.658 (0.023) | 0.602 | 2.519 |
| EV    | 0.679 (0.020) | 0.678 | 2.545 |
| ME    | 0.664 (0.023) | 0.642 | 1.723 |
The comparison between the engines performance has lead to the two observations mentioned above.

Regarding the first observation, it should be noted that noisy examples are probably more prevalent in the failure cases than in the successful therapies. To assess this, we carried out the following test. Recall that a label is defined by the SD via a follow-up VL measure. In particular if multiple VL tests are provided, the one closest to 8 weeks from the therapy switch determines the label of the therapy where the test needs to be in the time window 4–12 weeks in order to be considered as a follow-up VL. In the training (test) set 350 (35) failing therapies are predicted to be successful by all three engines. Once during the course of therapy 145 (16) of these achieve a VL measure below 500 copies per milliliter Of the remaining 550 (64) failing cases in the training (test) set 100 (13) have a VL measure below 500 copies per milliliter once during the course of the therapy. A Fisher’s exact test results in a P-value of 4.8 × 10−14 (0.011) on the training (test) set. Hence, the difference is highly significant. Our suspicion is that this behaviour is related to low adherence; it is known that law adherence disrupts the effectivity of HAART treatments, (see, e.g. Conway, 2007).

Regarding the second observation, we were interested in finding what are the features most related to disagreement between engines. Thus, we divide the 3K train/test therapies into the 592 cases where the engines provide identical prediction. We tested the correlation between individual drugs and individual mutations to an indicator of agreement/disagreement of the three engines. This is carried out to identify drugs and mutations that are most confusing, i.e. influence engines in different ways. We also use the χ²-test to check which features are distributed in the identical prediction case differently than in the different prediction case. In Table 5 we provide results with the lowest P-value after adjustment with Bonferroni correction.
Table 5. Drugs and mutations most related to inconsistency among engines

| Feature          | Agreement/disagreement | Success/failure |
|------------------|------------------------|-----------------|
| RT T215F        | -0.10 (2 \times 10^{-4}, 1) | -0.10 (1 \times 10^{-4}, 1) |
| RT K219Q        | -0.12 (2 \times 10^{-4}, 2 \times 10^{-5}) | -0.10 (1 \times 10^{-4}, 1) |
| RT L10I         | -0.10 (1 \times 10^{-4}, 1) | -0.10 (1 \times 10^{-4}, 1) |
| RT M41L         | -0.15 (5 \times 10^{-4}, 1 \times 10^{-5}) | -0.20 (0, 0) |
| RT T215Y        | -0.18 (0, 0) | -0.23 (0, 0) |
| RT M41I         | -0.15 (5 \times 10^{-4}, 1 \times 10^{-5}) | -0.20 (0, 0) |
| RT M184V        | -0.14 (2 \times 10^{-4}, 7 \times 10^{-5}) | -0.13 (3 \times 10^{-4}, 4 \times 10^{-5}) |
| RT D67N         | -0.14 (2 \times 10^{-4}, 6 \times 10^{-4}) | -0.16 (7 \times 10^{-4}, 8 \times 10^{-7}) |
| RT K219Q        | -0.12 (2 \times 10^{-4}, 3) | -0.12 (2 \times 10^{-4}, 3) |
| RT K70R         | -0.11 (6 \times 10^{-4}, 10^{-5}) | -0.08 (0, 0) |
| RT L211W        | -0.10 (1 \times 10^{-4}, 3) | -0.20 (0, 9.4 \times 10^{-11}) |
| RT T215F        | -0.10 (2 \times 10^{-4}, 2 \times 10^{-5}) | -0.10 (3 \times 10^{-4}, 4) |
| RT V118I        | -0.09 (4 \times 10^{-4}, 2 \times 10^{-5}) | -0.17 (2 \times 10^{-4}, 4) |

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**APPENDIX: MUTATIONS SELECTION BASED ON CORRELATIONS**

The original feature space of the mutations is the product of the four drugs, D4T, DDI, SQV and NFV, found as most correlated with disagreement among engines, are those drugs whose success rate is unbiased. In other words, their success rate in the training data is around 0.5, Table 2. Interestingly, all mutations found are from the list of EAS mutations. Moreover, the three PRO mutations appearing in the table are among those most involved in cross-resistance within the PI class and the RT mutations listed in the same table comprise the thymidine analog mutations well known to mediate cross-resistance to all the NRTIs, e.g. Gallant et al., 2003. These mutations seem to be related to therapies for which selecting antiretroviral compounds is most challenging.

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L33F, G73S, Q58E, L63P, I54I, L10F, K20R, G73T, V82T, I54M, K43T, V82F, I47V.

The 25 RT mutations with highest (absolute value of) correlations, sorted from the highest correlation (0.22) to lowest correlation (0.07): T215Y, M41L, L210W, V118I, D67N, H208Y, M184V, E44D, T69D, T215F, T39A, K101Q, M184M, V75M, K219Q, K70R, K103N, L74I, L228H, S162D, T215N, V181C, V75V, F77L, F116Y.

Top five mutations appearing in history genotype, followed by the value of correlation with therapy outcome: RT M41L, (0.19) PRO L10I, (0.19) PRO L90M, (0.19) PRO I54V, (0.18) PRO L63P, (0.17).