Heinz 95-845: Project Report

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Abstract

Opioid overdose deaths spiked in 2017 and continue to be a major concern for the US healthcare system. Major efforts have been dedicated to understand the demographic impacted and how the healthcare system can improve outcomes for individuals at risk. This paper explores the application of machine learning and statistical analyses to identify which could best predict (and prevent) overdose deaths using county-provided demographic, program activity and opiate prescription fills data for Medicaid beneficiaries from 2009-2017. The models with superior performance, Multivariate Logistic Regression and Random Forest, yield an AUROC of approximately 0.83, correctly identifying 57% of individuals who overdose. We found that the number of Tramadol prescribed, the number of drug and alcohol services received, whether an individual is White, and the number of lower court drug-related criminal cases were the most informative variables towards predicting opioid overdose deaths. We also note survival probabilities varying by gender, age and race. This paper showcases how county-provided data can be used to predict and prevent overdoses (translatable to other counties), identifies the variables indicative of risk, and highlights modeling techniques that can be tuned to improve predictive performance, thereby moving the field towards overdose prevention.

1. Introduction

Opioid abuse has been an emerging public health issue, and was declared a public health emergency in 2017¹. In the past two decades, national drug overdose deaths have increased from 16,849 in 1999 to 70,237 in 2017². This trend is especially affected by the onset of the opioid epidemic in the late 1990s³. Approximately 67% of the overdose deaths in 2017 were due to the involvement of any opioids (including illegal drugs like heroin and fetanyl) and 24% are specifically due to legally prescribed opioids, like oxycodone, hydrocodone, codeine, morphine, etc.⁴⁵. It has become critical to better understand the risk factors leading to overdoses and to determine the best way to prevent overdose deaths.

Crosier et al. predicted overdose frequency using random forests in order to uncover important features related to the frequency and development of overdose events⁶. Lobo et al. identified subgroups of Pennsylvania patients at greater risk for opioid abuse in a k-means clustering algorithm⁷. Our work aims to contribute to this existing body of work by identifying features that are most indicative of risk, which is crucial to preventing addiction and overdoses. There exists an Opioid Risk Tool, developed in 2005, to flag patients at risk for opioid abuse and overdose⁸. However, due to the subjective nature of this tool and the spike of deaths in 2017, machine learning has been sought to provide a more objective and quantitative approach to estimate risk of opioid abuse and overdose. A super learning approach was developed by Acion et al. to predict the successful treatment of patients with substance use disorders⁹. Artificial intelligence was also applied in the sphere of predicting opioid abuse by Haller et al., who implemented natural language processing on

electronic health record data to assess risk and predict opioid abuse¹⁰.

This paper strives to expand upon previous contributions to the field by identifying the machine learning algorithms that yield the highest predictive performance of overdose events. Specifically, those models that correctly identify the greatest number of at-risk individuals. This project uses datasets containing information concerning demographics, county-provided program usage, and opioid prescriptions to predict overdose deaths (opioid and non-opioid). Specifically, we analyze the data of Medicaid beneficiaries from 2009 to 2017 provided by the Allegheny County Department of Human Services (DHS) to predict the risk of death due to an non-opioid or opioid-related overdose. Here, we apply machine learning to uncover features and methods that successfully predict overdose death, and thereby enhance the space of addiction and overdose death prevention.

2. Methods

2.1 Original Data Description

We accessed 3 datasets from Allegheny County DHS: demographic, program activity, and opiate prescription fills. Information within contained data for 120,650 individuals who have utilized DHS services between 2009 and 2017. The demographic dataset (summarized in figure 6) contained variables for person ID, race, and gender. Of note, there was missingness in terms of race data for 19,531 individuals and gender information for 210 individuals. Additionally, there were demographics that were not heavily represented in the rest of the dataset: one transgender female and 19 Native Hawaiian / Pacific Islanders.

The program dataset (summarized in figure 7) contains 2,402,479 rows where each row denotes the activity (or activities) for an individual at any time since they entered the system. The program dataset contains variables for person ID, year and month of activity, overdose details (if any), and the DHS program-related relevant activity information. Program-related activity information included whether there was documentation for Child, Youth and Family (CYF) services used as child or parent (binary); the number of criminal court cases (drug or not) filed; mental health, drug and alcohol abuse, or prescription services (binary); and whether the individual was jailed (binary). If an individual experienced an overdose event, there were additional data fields for overdose date and a binary value for whether the overdose was opioid-related. The only missingness in the program activity dataset is in the overdose-related fields for individuals who did not overdose.

The opiate prescription fills dataset (summarized in figure 8) contains 1,161,650 rows, where each row represents a prescription fill for an individual. The prescription information variables are the claim number (unique for each row), person ID, age at prescription, dispensed quantity, days supply, fill date, and information specific to the drug (drug strength, name variations, package description, and dosage form). We did not include generic tier description and claim rank of prescriptions in our analyses for they are not informative. There were missing, extreme or incomprehensible values in the prescription dataset, such as an age of -7990, dispensed quantity of 17936 and days supply of 907. In addition, the prescription dataset contained multiple versions of drug names for the same drug. There were two columns pertaining to the drug dosage form, a condensed version and a descriptive version.

2.2 Data Cleaning and Feature Extraction

2.2.1 Demographic and Program Datasets

The missing race and gender information in the demographic dataset is likely missing not at random (MNAR) as it may be directly related to the unreported value itself. We dealt with this missingness

through assigning missing values their own category, "No Data". Furthermore, with respect to the race column, given that the "Native Hawaiian / Pacific Islander" demographic was underrepresented with only 19 individuals, we merged these individuals with the "No Data" race category to create a variable called "No Data and Other". This was replicated for the gender variable where we merged the "Transgendered male to female" and the "No Data" categories into "No Data and Other." With respect to the program activity dataset, we cleaned the outcome variable by converting the 0, 1 and NA to "Non-Opiate Overdose", "Opiate Overdose" and "No Overdose".

2.2.2 Opiate Prescription Fills Dataset

The age at which an individual got their first prescription was extracted. If the minimum age was less than 0, the maximum age was used. If both values were less than 0, then they were considered missing. There were 26 individuals with missing age values; these were replaced with the median age of individuals in the entire dataset. Drug strength values of oral solutions were updated to 5-325/5ML and of Naloxone were updated to 50MG-0.5MG. Days supply and dispensed quantity extreme values were censored. The top extreme values were replaced with the value at the 0.5% percentile and bottom extreme with the value at the 95.5% percentile. These percentile values were determined for each generic name and dosage form combination. Opioid strength values were pulled from the drug's label name, normalized to be per 1 ML, MCG or MG (if applicable). Missing values were replaced with those from provided drug strength column. Drug name (generic) and dosage forms were stripped to common terminology. The above-mentioned opioid strength values were converted into Morphine Milligram Equivalents (MME) using Opioid Morphine Equivalent Conversion Factors¹¹. The formula used was: opioid strength * (dispensed quantity/days supply) * conversion factor

2.2.3 Final Dataset

There were 30 meaningful features that were extracted at the person ID level. Demographic characteristic features included race, gender and age. DHS program usage features entailed cohort (the enrollment year in DHS services), usage of CYF program as child (total_cyfchild) and as parent (total_cyfparent), use of mental health, drug and alcohol abuse and prescription services (total_mh, total_da, total_rx), number of months spent in jail (total_acj), and number of criminal cases (total_cr_cases) and drug-related cases filed in court (total_cr_drug_cases).

Prescription activity features extracted included the number of prescriptions (num_presc) , name of the most prescribed drug $(most_presc_drug)$ and most prescribed dosage form $(most_dose_form)$, count of top three drugs $(oxy_count, tram_count, hydrobit_count)$, count of top three dose forms $(pill_count, patch_count, liquid_count)$, average, median and mode MME $(avg_mme, median_mme, mode_mme)$, as well as average days supply and dispensed quantity $(avg_supply$ and $avg_dispensed)$. Finally, overdose information $(od_type, od_month, od_year, od_date)$ was added.

2.2.4 Feature Choices

Some additional feature choices were made to enhance the final dataset and prepare for the prediction tasks. Features such as <code>most_presc_drug</code> and <code>most_dose_form</code> had categories that were comprised of less than 0.2% of the values. To prevent errors during the prediction task, these low occurrence categories were combined to create a separate Other category. We plotted the distribution of the average, median, and mode of morphine milligram equivalents (MME) variables to determine which mode of central tendency for MME was normally distributed. This led us to retain only median MME in the final dataset to avoid multicollinearity.

We examined the outcome variable under a binary value of 1 for an "opiate overdose" or "non-opiate overdose" and 0 for there being "no overdose". We include non-opiate overdose events in our

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outcome variable because these individuals were receiving opioid prescriptions and we are willing to increase the number of potential false positives to flag as many potentially at-risk individuals as possible. Furthermore, we argue that individuals who overdose (opiate or non-opiate) share similar characteristics and so the inclusion of both types of overdose in our outcome variable is informative in the prediction task. Columns pertaining to the date of overdose were removed to avoid leakage in the machine learning models, as they are proxies for the target variable. All categorical variables were converted to dummy variables and all variables were log transformed.

2.3 Oversampling, Machine Learning and Statistical Models, and Evaluation Metrics

Since there is a severe under-representation of outcome variable events (1,222 out of 120,650 individuals), the first model (multivariate logistic regression) was unable to capture a single case of overdose. As such, this paper utilized an oversampling method, Synthetic Minority Oversampling Technique (SMOTE) for the outcome variable. To do so, we divided the whole dataset into train and test with a 50-50% split. We then ran SMOTE on the train set to generate synthetic data to train our models and then calculated performance on the original test set. Furthermore, to increase the performance of our machine learning models, we trained the data only on the top 20 critical features identified using the random forest algorithm.

Six machine learning models were run and their performance in predicting overdose rates compared. These models include: multivariate logistic regression, ridge regression, random forest, AdaBoost, gradient boosting, and neural nets. We used 5-fold cross-validation for multivariate logistic regression, and 10-fold cross-validation for ridge regression, random forest, and gradient boosting. We also implemented lasso regularization. Note that for this use case in particular, it is more critical to identify the individuals who may overdose (true positives) than to eliminate individuals not at risk from consideration (true negatives). Furthermore, given the limited resources available to tackle such a pressing issue, it is valuable to minimize (to the extent possible) the number of individuals incorrectly identified to be at risk (false positives). Therefore, we evaluated and compared machine learning models through the area under the curve (AUC) of the Receiver Operating Characteristic (ROC) curves, the number of true positives, the number of false positives, sensitivity, specificity, and precision. The important or significant features used in these models will also be an additional point of comparison, specifically for the multivariate logistic regression and random forest models.

This paper also includes a survival analysis, conducted using Kaplan-Meier estimation and Cox Proportional Hazards models. For the survival analysis, the time variable was the months in system and the event variable was the status of the individual at their last known month in the system. The maximum month in the system is 108 months (December, 2017). The event was 1 for overdose death and 0 otherwise. The Kaplan-Meier estimation was evaluated visually (with the survival curve) along with the model output. Both normal and regularized versions of the Cox Model were produced. The coefficients, exponent coefficients and their corresponding p-values were used to understand and evaluate both models. Cross-validation was used to determine the best lambda to use for the regularized Cox Model. Code is available at https://github.com/nikitascmu/S19-aamlp-project-group1.

3. Results

Table 1 provides an overview of the 120,650 individuals in the Allegheny County DHS cohort between 2009 and 2017. No individuals were excluded from the cohort. We addressed all missingness to the extent possible.

Table 1. Demographics of Allegheny County DHS Cohort (2009-2017).

Characteristic	N	Percentage (%)
Race		
White	56,754	47.04
Black/African-American	40,573	33.63
Biracial/Multiracial	2,118	1.75
Asian	1,216	1.00
American Indian/Alaskan Native	439	0.36
No Data and Other	19,550	16.20
Total	120,650	100
Gender		
Female	73,582	60.99
Male	46,857	38.84
No Data and Other	211	0.17
Total	120,650	100
Age(years)		
0-19	27,128	22.48
20-39	51,883	43.00
40-59	35,941	29.79
60 & Over	5,698	4.72
Total	120,650	100

3.1 Evaluation of Machine Learning Models

The ROC curves of all 6 machine learning models run in this paper are showcased in figure 1 below.

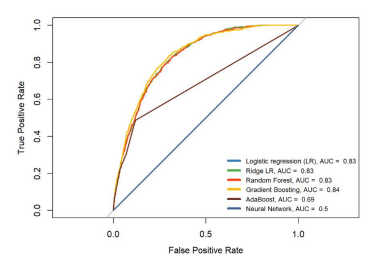


Figure 1. Receiver Operating Characteristic (ROC) curves with corresponding AUCs for the 6 ML models

These curves, along with their reported AUCs, indicate that Gradient Boosting performs the best with Logistic Regression, Ridge Regression, Random Forest close behind. AdaBoost does not perform well and Neural Networks perform only as good as chance. As such, AdaBoost and Neural Networks can be dropped from consideration for the time being. However, it would be premature to select Gradient Boosting as the best model. As stated earlier, it is vital to identify the individuals who are at risk of overdose (true positives) while keeping the number of false identifications to a minimum. As such, while Gradient Boosting may appear to perform the best, whether it is truly the best model in this case depends on its performance on additional evaluation metrics.

Table 2 summarizes the performance of all six machine learning models in predicting overdose deaths. Please note that there are 633 overdoses in the test dataset. As seen in this table, despite having the highest AUC, Gradient Boosting correctly identifies only 22% of overdose deaths (few relative to the other high performing models). Neural nets identifies all overdose deaths simply

Table 2. Evaluation Metrics of the Machine Learning Models

Model	True Positives	False Positives	Sensitivity (%)	Specificity (%)	Precision (%)	AUC
Multivariate Logistic Regression	362	8654	57	86	4	0.83
Ridge Regression	163	2422	26	96	6	0.83
Random Forest	362	8654	57	86	4	0.83
Gradient Boosting	138	1982	22	97	7	0.84
Ada Boost	147	2389	23	96	6	0.69
Neural Net	633	59692	100	0	0	0.5

because it classifies every individual at risk of overdose, so it can be excluded from consideration. Of the remaining four models, Multivariate Logistic Regression and Random Forest perform the best, correctly identifying 57% of the at-risk individuals.

3.2 Understanding the Survival Analysis

Survival analysis using the Kaplan-Meier estimation yields a survival curve seen in figure 2 in the appendix. The plot shows us that the probability of survival gradually declines from 100% at the beginning to 96.6% at the final month in the system (108 month mark). Individuals have dropped out of the system i.e. stopped utilizing DHS services and filling prescriptions at various points throughout the time-frame. The earliest time at which we start to see a decline in survival probability is 8 months.

Next, we explored if and how this probability of survival changes among various gender and race subgroups. Figure 3 shows us that females have a higher probability of survival than males in this dataset. Furthermore, individuals who are in the 'no data and other' category have a relatively lower probability of survival. Females have a 97.5% probability of surviving, males have a 95.3% probability of surviving, and individuals who have either not reported their gender or are transgender female have a 72.7% probability of surviving an overdose event by the time they conclude their time in the system.

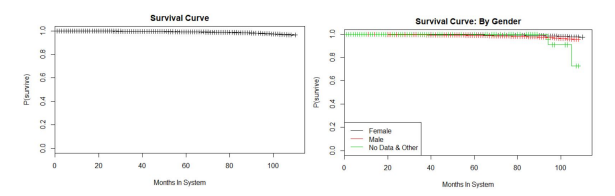


Figure 2. Kaplan Meier survival curve (all).

Figure 3. Kaplan Meier curve by gender.

Table 3 below display the survival probabilities at the conclusion of time in the system by race. Since no American Indian or Native Alaskan individuals experience an overdose event, no survival probability was calculated for them. There appear to be subtle differences in survival probabilities among the remaining racial categories. For example, we see that White individuals have the lowest relative survival probability (95%) and Asian individuals have the highest survival probability (99.6%). None of the probabilities, however, are lower than 95% and the differences are not as huge as those seen by gender alone.

Table 3. Probability of survival by Race

Race	Probability of Survival (%)
American Indian or Native Alaskan	Not Applicable
Asian	99.6
Biracial or Multiracial	96.6
Black or African American	98.6
No Data or Other	97.4
White	95.0

Next, we use a Cox Proportional Hazards model to analyze the survival probability of individuals by including age, gender and race as predictor variables. Figure 4 below shows the output of the Cox model which indicates that age, being male, being in the "No Data and Other" gender category, and being White are statistically significant predictors of survival. The base variables are being female and being Asian. Given that there are no overdose events for the American Indian or Native Alaskan individuals, their data was merged into the "No Data and Other" race category so as to not skew the results. The results show that, **holding all other things constant**: a one year increase in age is associated with an increase in hazard by a factor of 1.028 or 2.8%; compared to being female, being male increases hazard by a factor of 1.87 or 87% and being an individual with no reported gender or other increases hazard by a factor of 7.9 or 690%; and compared to being Asian, being White increases hazard by a factor of 8.2 or 720%. These findings are aligned with our findings using the Kaplan Meier estimation.

	coef	exp(coef)	se(coef)	Z	p
age	0.02736	1.02774	0.00208	13.19	<2e-16
genderMale	0.62619	1.87047	0.05815	10.77	<2e-16
genderNo Data and Other	2.06827	7.91115	0.71324	2.90	0.0037
raceBiracial/Multiracial	1.55887	4.75345	1.03539	1.51	0.1322
raceBlack/African-American	0.81961	2.26961	1.00269	0.82	0.4137
raceNo Data and Other	0.91127	2.48747	1.01580	0.90	0.3697
raceWhite	2.10949	8.24405	1.00066	2.11	0.0350

Figure 4. Cox Model Output with Age, Race and Gender as Predictors

3.3 Variable Importance

Figure 5 below showcases the top 15 variables from the Random Forest model. The variables most important in predicting overdose was the total number of Tramadol prescribed $(tram_count)$, the total number of drug and alcohol services received $(total_da)$, whether an individual is white $(race_White)$, and the total number of lower court drug-related criminal cases $(total_cr_drug_cases)$. These four variables were also statistically significant in the multivariate logistic regression. Most of the remaining variables were statistically significant except the average number of dispensed drugs $(avg_dispensed)$, whether an individual was male $(gender_Male)$, and the total number of mental health services received $(total_mh)$.

4. Discussion

Due to the under-representation of overdose events in our dataset, in spite of oversampling, even the best machine learning models yielded a relatively low sensitivity (57%), and fairly high number of false positives. While one could argue that it is beneficial to be extra cautious by overestimating the number of at-risk individuals as it would good a long way towards preventing addiction and overdose deaths. However, the high false positives may overburden counties' substance abuse prevention and treatment efforts. One way to address this could be gathering additional data, perhaps by pooling data from other counties or for additional years, or both. Another way to enhance the existing

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Top 15 Variables Determined via Random Forest

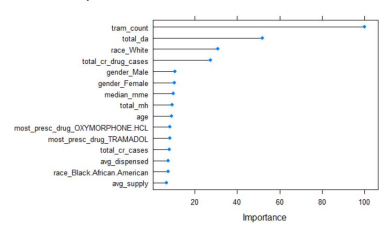


Figure 5. The top 15 variables of importance elucidated from the random forest model.

algorithms (especially boosting) would be to tune the hyperparameters. Moreover, the neural network had very poor performance because we utilized four dense layers and ran the algorithm for 30 epochs. While a deep neural network was computationally expensive for us, we would expect it to yield a much higher performance than the one seen in this paper.

While our analysis showcased gender- and race-related differences in survival, we cannot be certain that it is not merely a reflection of the Allegheny County population proportions or if it is reflective of some association between these factors and drug abuse. Our work may also be limited by the fact that it may not be generalizable to other counties, especially if they do not collect similar data in the same way. Furthermore, certain features were created specifically for this dataset, such as the count of the top three drugs (one of which was very predictive of overdose events) and dosage forms. Our analysis is also limited by the data collected by the county's DHS; we did not have access to potentially relevant variables, such as socioeconomic status, medical history, diagnoses, and insurance claims. Additionally, some feature creation could benefit from medical expertise, such as MME.

5. Conclusion

This paper applies the analytical pipeline on Allegheny County DHS demographic, prescription and program activity data for 120,650 individuals from 2009 to 2017 to predict the rate of opiate and non-opiate overdose. The best models, Logistic Regression and Random Forest, yielded an AUROC of 0.83 and correctly identified 57% of the at-risk population. We believe this analytical pipeline is a foundation that can be further improved using a bigger, enriched dataset that has a relatively more balanced outcome variable, better tuned models, and additional computational power, thereby moving the field one step closer towards preventing addiction and overdose deaths.

Appendix

PERSON_ID	F	RACE		GENDER
Min. : 1	American Indian/Alaskan Native	: 439	Female	:73582
1st Qu.: 30163	Asian	: 1216	Male	:46857
Median : 60326	Biracial/Multiracial	: 2118	No Data	: 210
Mean : 60326	Black/African-American	:40573	Transgendered male to	female: 1
3rd Qu.: 90488	Native Hawaiian/Pacific Islande	er: 19		
Max. :120650	No Data	:19531		
	White	: 56754		

Figure 6. Summary of the original demographic dataset.

PERSON_ID	YEAR	MONTH	CYFCHILD	CYFPARENT	MDJS_CR_CASES	MDJS_CR_DRUG_CASES
Min. : 1	Min. :2009	Min. : 1.00	Min. :0.00000	Min. :0.00000	Min. : 0.00000	Min. :0.00000
1st Qu.: 30364	1st Qu.:2011	1st Qu.: 3.00	1st Qu.: 0.00000	1st Qu.: 0.00000	1st Qu.: 0.00000	1st Qu.: 0.00000
Median : 60557	Median :2013	Median: 6.00	Median :0.00000	Median :0.00000	Median : 0.00000	Median : 0.00000
Mean : 60425	Mean :2013	Mean : 6.48	Mean :0.02653	Mean :0.07775	Mean : 0.04274	Mean :0.01301
3rd Qu.: 90391	3rd Qu.: 2015	3rd Qu.: 9.00	3rd Qu.: 0.00000	3rd Qu.: 0.00000	3rd Qu.: 0.00000	3rd Qu.: 0.00000
Max. :120650	Max. :2017	Max. :12.00	Max. :1.00000	Max. :1.00000	Max. :18.00000	Max. :6.00000
MH	DA	RX	ACJ	OVERDOSE_DATE	OPIATE_OVERDOSE	
Min. :0.0000	Min. :0.0000	Min. :0.000	0 Min. :0.0000	00 Min. :200901	Min. :0.0	
1st Qu.:0.0000	1st Qu.: 0.0000	1st Qu.: 0.000	0 1st Qu.:0.0000	00 1st Qu.:201411	1st Qu.:1.0	
Median : 0.0000	Median : 0.0000	Median : 0.000	0 Median: 0.0000	00 Median : 201607	Median :1.0	
Mean :0.4921	Mean :0.1509	Mean : 0.345	3 Mean : 0.0759	2 Mean :201539	Mean :0.9	
3rd Qu.:1.0000	3rd Qu.: 0.0000	3rd Qu.:1.000	0 3rd Qu.: 0.0000	00 3rd Qu.:201703	3rd Qu.:1.0	
Max. :1.0000	Max. :1.0000	Max. :1.000	0 Max. :1.0000	00 Max. :201802	Max. :1.0	
				NA's :236273	9 NA's :2362739	1

Figure 7. Summary of the original program activity dataset.

CLAIM_NR	PERSON_ID AG	E DISPENSED_QTY	DRUG_STRE	NGTH DAYS_SUPPLY
000000064742: 1	Min. : 1 Min.	:-7990.00 Min. : ().00 50 MG :1	.23829 Min. : 0.00
000000064849: 1	1st Qu.: 29949 1st Qu.	: 32.00 1st Qu.: 18	3.00 5MG-325MG:	89846 1st Qu.: 4.00
000000065039: 1	Median : 59718 Median	: 45.00 Median: 40).00 5 MG :	87096 Median: 11.00
000000066034: 1	Mean : 59960 Mean	: 42.69 Mean : 58	3.73 5 MG-325MG:	71169 Mean : 15.57
000000070555: 1	3rd Qu.: 89653 3rd Qu.	: 53.00 3rd Qu.: 90	7.5-750MG:	65156 3rd Qu.: 30.00
000000115549: 1	Max. :120650 Max.	: 354.00 Max. :17936	5.00 5MG-325M :	56623 Max. :907.00
(Other) :1161644	NA's	:126	(Other) :6	67931
FILL_DATE	L	ABEL_NAME	BRAND_NAME	GENERIC_NAME
2013-02-01: 868	TRAMADOL HCL 50 MG TABLET	:148444 HYDROCODONE	-ACETAMI: 495685	HYDROCODONE BITARTRATE: 498280
2013-07-01: 863	OXYCODONE-ACETAMINOPHEN 5-	325 :147962 OXYCODONE-A	ACETAMINO: 190069	OXYCODONE HCL/ACETAMIN: 218559
2014-08-01: 859	HYDROCODON-ACETAMINOPHEN 5	-500:132225 OXYCODONE F	ICL : 176250	OXYCODONE HCL :184272
2012-06-01: 855	HYDROCODON-ACETAMINOPH 7.5	-750:120385 TRAMADOL HO	: 148444	TRAMADOL HCL :148635
2013-08-01: 841	HYDROCODON-ACETAMINOPHEN 5	-325:112172 MORPHINE SU	JLFATE ER: 26912	MORPHINE SULFATE : 31889
2013-04-01: 837	OXYCODONE HCL 5 MG TABLET	: 93667 FENTANYL	: 21559	OXYMORPHONE HCL : 28456
(Other) :1156527	(Other)	:406795 (Other)	:102731	(Other) : 51559
PACKAGE_DESC	DOSAGE_FORM_CD	DOSAG	GE_FORM_DESC GENE	RIC_TIER_CLASS_DESC CLAIM_RANK
BOTTLE :1138114	TA :1059465 TABLET		:1059465 ANAL	GESICS:1161650 Min. :1
BOX : 22198		EXTENDED RELEASE	: 26805	1st Qu.:1
BLIST PACK: 843		RANSDERMAL 72 HOURS		Median :1
VIAL : 213	TM : 19707 TABLET,	EXTENDED RELEASE 12 HR	: 19707	Mean :1
SQUEEZ BTL: 198	SJ : 14668 SOLUTION	, ORAL	: 14668	3rd Qu.:1
AMPUL : 45		RAL ONLY, EXTENDED RELEASE		Max. :1
(Other) : 39	(Other): 5023 (Other)		: 5023	

Figure 8. Summary of the original opiate prescription fills dataset.

Table 4. Regularized Cox proportional Coefficients

Race	coefficient	exp(coefficient)	Change in hazard as %
age	0.027	1.027	2.7
gender_female	-0.617	0.540	-46
gender_male			
gender_No Data and Other	1.285	3.615	261.5
race_white	1.151	3.161	216.1
race_Black/African-American	-0.119	0.888	-12.2
race_Biracial/Multiracial	0.543	1.721	72.1
race_No Data and Other			
race_Asian	-0.459	0.632	-36.8
race_American Indian/Alaskan Native	-1.102	0.332	-66.8

Endnotes

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