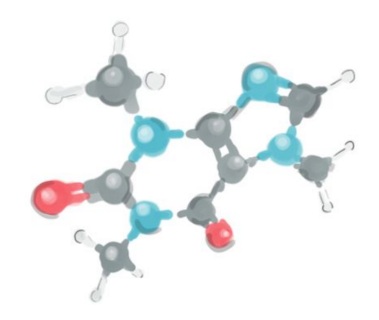
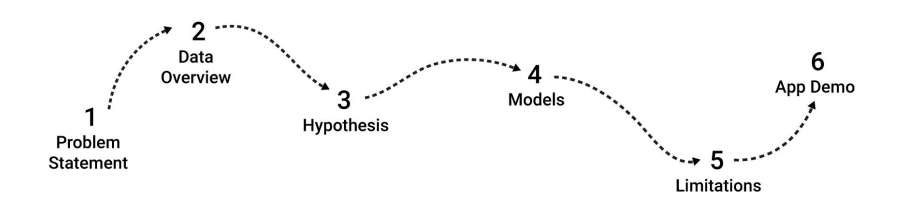
### Multi Class Drug Classification Using Molecular Structures

**Vivian Peng** 



https://github.com/veeps/molecular classification

### **Overview**





#### **Problem Statement**

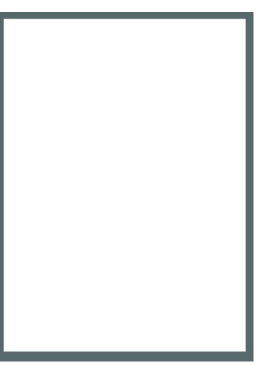
Current drug R&D is long and costly...
And it doesn't incentivize for research into treatments for neglected diseases.

What if we could improve the R&D process

by reducing redundancy, and screen for

multiple therapeutic uses in parallel?

#### **Neural Network**

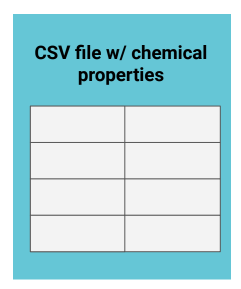


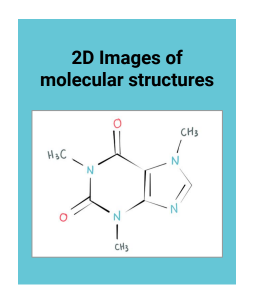
Antineoplastic

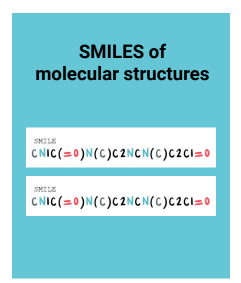
CNS

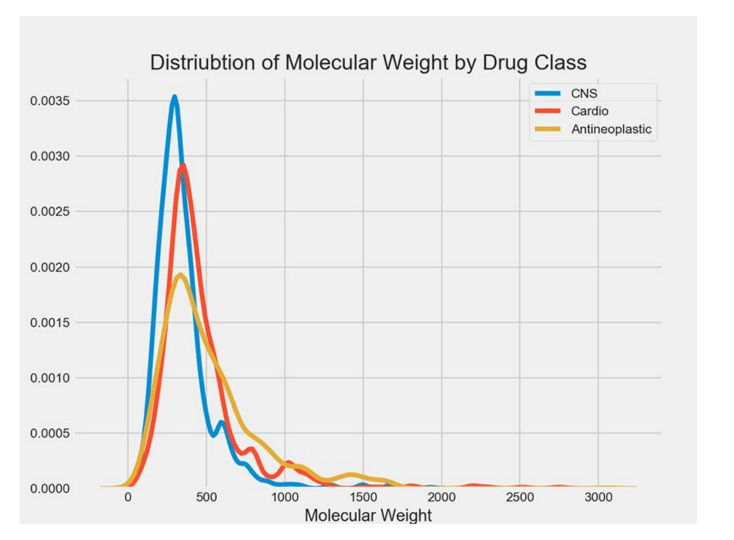
Cardio

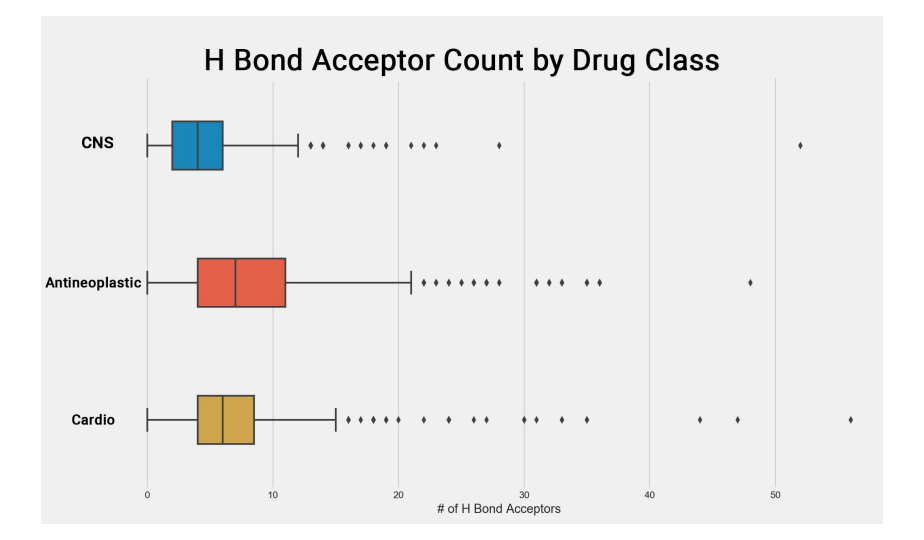
#### Here's what my data looked like:



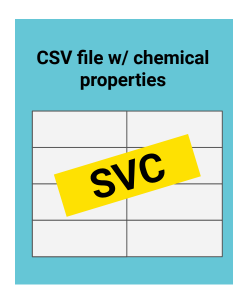


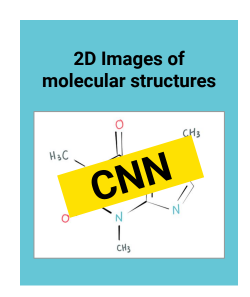


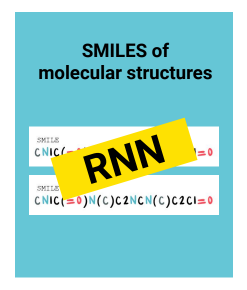




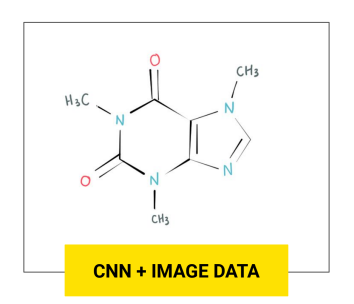
#### Ran 3 types of models:

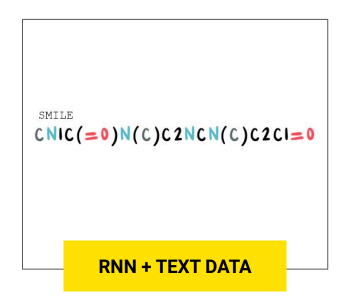


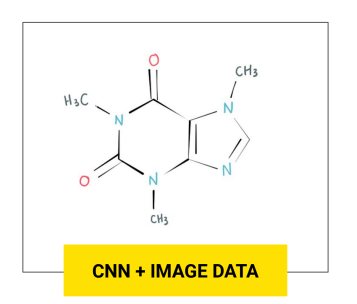




## How do these neural networks compare in classifying drug classes using molecular structure data?







My hypothesis was that a CNN model using image data would be better at classifying than an RNN model using SMILES.

### Models

## SVC Support Vector

Classification

#### **FEATURES:**

- Molecular Weight
- Hydrogen Bond Acceptors
- Hydrogen Bond Donors
- XlogP (Solubility)

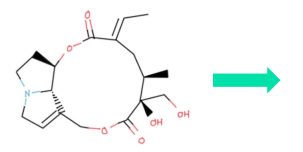
#### **HYPERPARAMETERS:**

- Ridge penalty
- Linear kernel
- Gamma = "scale"

### **CNN**

Convolutional Neural Network

#### **DATA PROCESSING:**

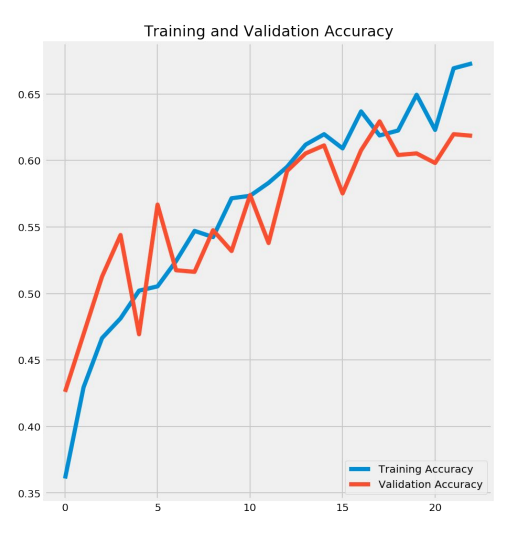


#### **MODELS:**

- Custom convolutional neural net
- Pre-trained VGG16 model
  - 2 hidden layers (256 and 128 neurons)
  - Adam optimizer

## CONVOLUTIONAL

**Neural Network** 



#### **DATA PROCESSING:**

Brc1c(NC2=NCCN2)ccc2nccnc12

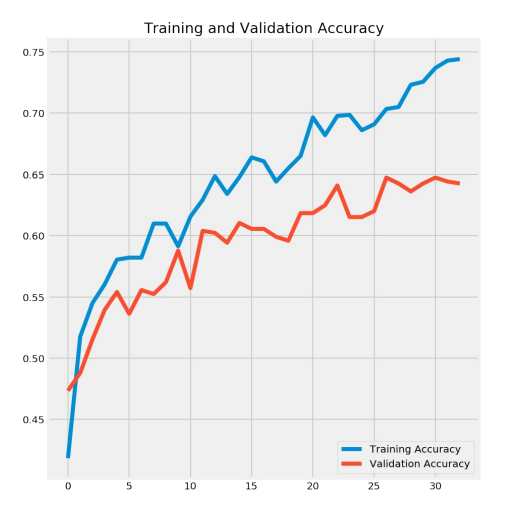


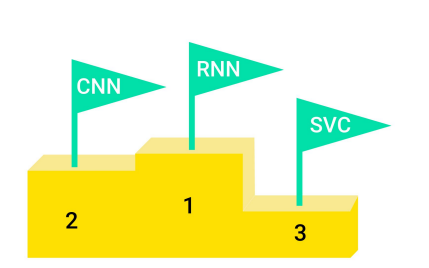
#### **MODEL:**

- Simple RNN
- Pre-trained VGG16 model
  - 5 hidden layers
  - Adam optimizer

# RECURRENT NEURAL Network

# RECURRENT Neural Network

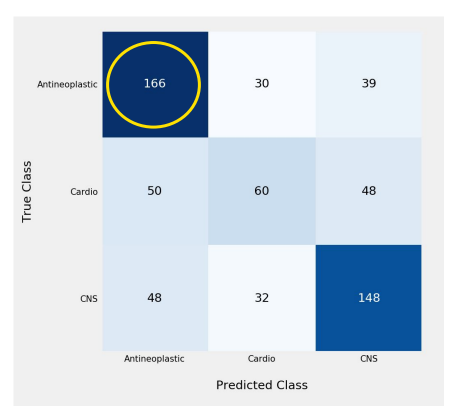




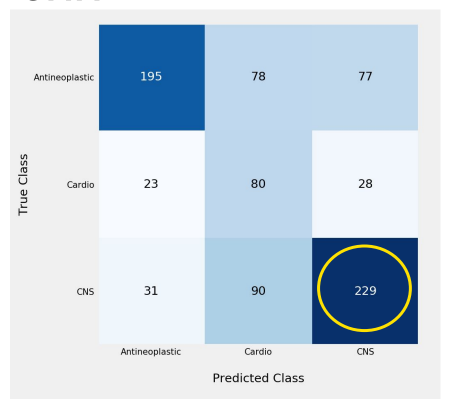
Model	Score
RNN	0.64
CNN	0.62
SVC	0.53
Baseline	0.37

Both types of neural networks performed **relatively the same** in predicting drug classes. **This is really interesting** because running an RNN with text data is **computationally much lighter** than running a CNN with image data.

#### **RNN**



#### **CNN**

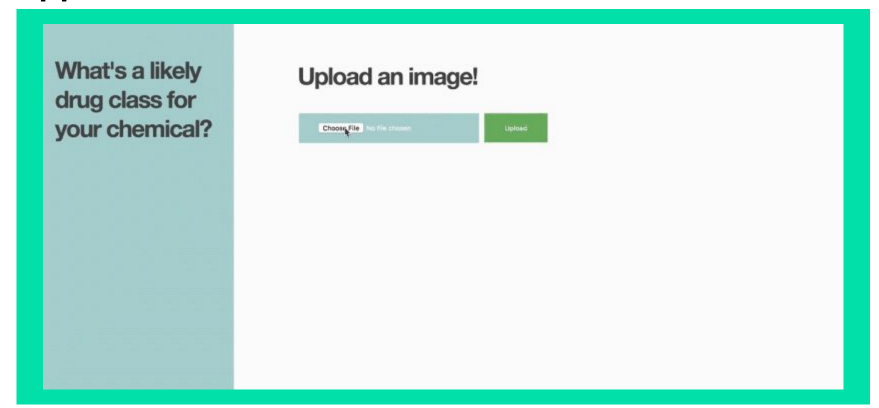


Models are good at predicting different types of drug classes.

#### Limitations

- Unable to tokenize SMILES to keep two-letter elements as one unit
- Lack of chemical expertise
- Unable to deploy app to Heroku

### App Demo



Thank you!