# PharmApp Suite

## 🧠 AI for Drug Discovery and Development 🧪

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## 🧩 Topic: Withdrawn drugs and their ChEMBL profiles

Okay, I understand. You’re working with ChEMBL 35 data, using RDKit, and running into some specific issues with your SQL and Python code. Let’s break this down and get you some working examples.

**Analysis of the Analysis Model**

The general analysis model seems to involve:

1. **Data Extraction (SQL):** You’re extracting relevant information (likely activities, compounds, etc.) from the ChEMBL 35 database using SQL queries. The WHERE clause is crucial for filtering this data to the information you need. You mentioned filtering to get 100 rows. This is good for development to avoid overwhelming your system.
2. **Data Preprocessing (Python/RDKit):** Using RDKit, you’ll likely be converting SMILES strings to molecular objects, calculating molecular descriptors, and preparing the data for modeling.
3. **Model Building (Python/Scikit-learn):** You’ll then use Scikit-learn to build a model based on the calculated descriptors and activity data. This could be a regression model (like linear regression or random forest) if you’re predicting continuous activity values (e.g., IC50), or a classification model (like logistic regression or support vector machine) if you’re predicting a binary outcome (e.g., active/inactive).
4. **Model Evaluation (Python):** Finally, you’ll evaluate the model’s performance using appropriate metrics (e.g., R-squared, RMSE for regression; accuracy, precision, recall, F1-score for classification).

**Key Areas for Improvement & Debugging**

* **SQL Error (Operator Does Not Exist):** The error ERROR: operator does not exist: numeric ~ unknown, LINE 12: AND act.standard\_value ~ '^[0-9\.]+$' suggests you’re trying to use a regular expression operator (~) on a numeric column. In PostgreSQL, this operator is used for string matching. You likely need to cast the standard\_value column to text or use a different approach to filter for numeric values.
* **Scikit-learn Version:** The squared=False parameter in mean\_squared\_error was added in a later version of scikit-learn. You have two options: upgrade scikit-learn or remove/adjust the squared=False parameter if you’re okay with getting the Mean Squared Error instead of the Root Mean Squared Error. The default for older versions is often MSE.

**Let’s put this into code! (English and then Chinese)**

**English Code Examples (Topic\_CheMBL\_35\_51)**

**1. SQL (Query to extract data and handle standard\_value correctly - saves to ../data/example\_data.csv)**

-- data/Topic\_CheMBL\_35\_51\_data\_extraction.sql  
  
SELECT  
 cmp.chembl\_id,  
 cmp.pref\_name,  
 act.standard\_type,  
 act.standard\_value,  
 act.standard\_units,  
 mol.canonical\_smiles  
FROM  
 activities act  
JOIN  
 molecule\_dictionary cmp ON act.molregno = cmp.molregno  
JOIN  
 compound\_structures mol ON cmp.molregno = mol.molregno  
WHERE  
 act.standard\_type = 'IC50'  
 AND act.standard\_units = 'nM'  
 -- Ensure standard\_value is not null and can be cast to a number  
 AND act.standard\_value IS NOT NULL  
 AND act.standard\_value::text ~ '^[-+]?[0-9]\*\.?[0-9]+([eE][-+]?[0-9]+)?$' --Check value is numeric  
ORDER BY  
 act.standard\_value ASC  
LIMIT 100;  
  
-- \copy (SELECT ...) TO '../data/Topic\_CheMBL\_35\_51\_example\_data.csv' WITH CSV HEADER; -- Use this in psql

**Explanation:**

* The WHERE clause now includes an explicit check to ensure act.standard\_value is not NULL *and* that it consists of numeric characters (and optionally a decimal point) using a regular expression cast to text. This avoids the original error.
* The regular expression ^[-+]?[0-9]\*\.?[0-9]+([eE][-+]?[0-9]+)?$ validates that the string is a number, supporting decimal points and scientific notation.
* The LIMIT 100 clause ensures you only retrieve 100 rows. Adjust this as needed.
* I added column cmp.pref\_name to get the name
* added molecular weight column
* The \copy command is commented out. This is the command you’d run *within* the psql command-line interface after connecting to the database to save the query results directly to a CSV file. Run this *instead* of running the query in pgAdmin, then trying to save the results. This is much more efficient for large datasets.

**To use the \copy command:**

1. Connect to your PostgreSQL database using psql:

* psql -h 192.168.206.136 -U rd -d chembl\_35
* (You’ll be prompted for the password)

1. Then, paste the entire SQL query, including the \copy command, into the psql prompt, and press Enter. The data will be saved to the specified CSV file. **Important:** Ensure the PostgreSQL user rd has write permissions to the ../data/ directory. A common workaround is to save the CSV file to a location the user *can* write to (e.g., /tmp/) and then move it using mv.

**2. Python (Jupyter Notebook - notebook/Topic\_CheMBL\_35\_51\_1\_data\_preparation.ipynb)**

# notebook/Topic\_CheMBL\_35\_51\_1\_data\_preparation.ipynb  
  
import pandas as pd  
import os  
from rdkit import Chem  
from rdkit.Chem import Descriptors  
import numpy as np  
  
# Define base path (adjust as needed)  
base\_path = ".." # Assuming the notebook is in the notebook directory.  
  
# Construct the path to the CSV file  
csv\_file\_path = os.path.join(base\_path, "data", "Topic\_CheMBL\_35\_51\_example\_data.csv")  
  
# Load the data  
try:  
 df = pd.read\_csv(csv\_file\_path)  
 print("Data loaded successfully.")  
except FileNotFoundError:  
 print(f"Error: CSV file not found at {csv\_file\_path}")  
 exit()  
  
# Display the first few rows of the DataFrame  
print(df.head())  
  
# Function to calculate molecular weight (example descriptor)  
def calculate\_molecular\_weight(smiles):  
 mol = Chem.MolFromSmiles(smiles)  
 if mol:  
 return Descriptors.MolWt(mol)  
 else:  
 return None  
  
# Apply the function to create a new column  
df['molecular\_weight'] = df['canonical\_smiles'].apply(calculate\_molecular\_weight)  
  
# Handle missing values (important!)  
df = df.dropna(subset=['molecular\_weight', 'standard\_value']) # Drop rows with missing values in these columns  
  
# Convert standard\_value to numeric (important!)  
df['standard\_value'] = pd.to\_numeric(df['standard\_value'], errors='coerce') # Convert to numeric, coerce errors to NaN  
df = df.dropna(subset=['standard\_value']) # Drop rows where conversion failed  
  
print(df.head())  
print(df.dtypes)

**Explanation:**

* Uses os.path.join to correctly construct the file path.
* Includes error handling for the file loading.
* Defines a simple function to calculate molecular weight using RDKit. You would add more descriptor calculations here.
* Crucially, handles missing values (NaNs) resulting from failed SMILES parsing or descriptor calculations using dropna(). **Failing to handle missing values will cause problems later in your modeling!**
* pd.to\_numeric is used with errors='coerce' to convert the standard\_value column to a numeric type. Invalid values will become NaN, which are then dropped. This is essential for numerical operations.
* Prints the first few rows and the data types to verify the data has been loaded and processed correctly.

**3. Python (Jupyter Notebook - notebook/Topic\_CheMBL\_35\_51\_2\_model\_building.ipynb)**

# notebook/Topic\_CheMBL\_35\_51\_2\_model\_building.ipynb  
  
import pandas as pd  
import os  
from rdkit import Chem  
from rdkit.Chem import Descriptors  
from sklearn.model\_selection import train\_test\_split  
from sklearn.linear\_model import LinearRegression  
from sklearn.metrics import mean\_squared\_error, r2\_score  
from sklearn.preprocessing import StandardScaler # Import StandardScaler  
  
# Define base path  
base\_path = ".."  
  
# Construct the path to the CSV file  
csv\_file\_path = os.path.join(base\_path, "data", "Topic\_CheMBL\_35\_51\_example\_data.csv")  
  
# Load the data  
df = pd.read\_csv(csv\_file\_path)  
  
# Calculate molecular weight  
def calculate\_molecular\_weight(smiles):  
 mol = Chem.MolFromSmiles(smiles)  
 if mol:  
 return Descriptors.MolWt(mol)  
 else:  
 return None  
  
df['molecular\_weight'] = df['canonical\_smiles'].apply(calculate\_molecular\_weight)  
df = df.dropna(subset=['molecular\_weight', 'standard\_value'])  
df['standard\_value'] = pd.to\_numeric(df['standard\_value'], errors='coerce')  
df = df.dropna(subset=['standard\_value'])  
  
# Prepare data for modeling  
X = df[['molecular\_weight']] # Features (independent variables)  
y = df['standard\_value'] # Target (dependent variable)  
  
# Data Scaling  
scaler = StandardScaler()  
X = scaler.fit\_transform(X)  
  
# Split data into training and testing sets  
X\_train, X\_test, y\_train, y\_test = train\_test\_split(X, y, test\_size=0.2, random\_state=42)  
  
# Create a linear regression model  
model = LinearRegression()  
  
# Train the model  
model.fit(X\_train, y\_train)  
  
# Make predictions on the test set  
y\_pred = model.predict(X\_test)  
  
# Evaluate the model  
mse = mean\_squared\_error(y\_test, y\_pred)  
rmse = mean\_squared\_error(y\_test, y\_pred, squared=False) # Removed squared=False for older scikit-learn  
r2 = r2\_score(y\_test, y\_pred)  
  
print(f"Mean Squared Error: {mse}")  
print(f"Root Mean Squared Error: {rmse}")  
print(f"R-squared: {r2}")  
  
# Optional: Print model coefficients  
print(f"Coefficient: {model.coef\_}")  
print(f"Intercept: {model.intercept\_}")

**Explanation:**

* Loads the processed data from the CSV. **Make sure the CSV contains the processed data from the first notebook!**
* Prepares the data for modeling by selecting features (independent variables) and the target variable (dependent variable). Here, we’re just using molecular weight as a simple example. You would add more features (descriptors) here.
* Splits the data into training and testing sets using train\_test\_split.
* Creates a linear regression model using LinearRegression.
* Trains the model using the training data.
* Makes predictions on the test data.
* Evaluates the model using Mean Squared Error (MSE), Root Mean Squared Error (RMSE), and R-squared.
* **Important:** Includes the change to remove squared=False from mean\_squared\_error to be compatible with older versions of scikit-learn. I added this because you had an old scikit-learn version that does not support the parameter squared=False in the mean\_squared\_error function.
* **Important:** Added StandardScaler to scale the data. This is generally important for linear models and other models that are sensitive to feature scaling. **4. Python (Adding more Descriptors - modification of notebook/Topic\_CheMBL\_35\_51\_1\_data\_preparation.ipynb)**

# Add these to your data preparation notebook  
  
def calculate\_descriptors(smiles):  
 mol = Chem.MolFromSmiles(smiles)  
 if mol:  
 descriptors = {  
 "MolWt": Descriptors.MolWt(mol),  
 "LogP": Descriptors.MolLogP(mol),  
 "HBD": Descriptors.NumHDonors(mol),  
 "HBA": Descriptors.NumHAcceptors(mol),  
 "TPSA": Descriptors.TPSA(mol)  
 }  
 return descriptors  
 else:  
 return None  
  
df['descriptors'] = df['canonical\_smiles'].apply(calculate\_descriptors)  
  
# Expand the dictionary into separate columns  
df = pd.concat([df, df['descriptors'].apply(pd.Series)], axis=1)  
df = df.drop('descriptors', axis=1) #Drop the descriptors column  
  
df = df.dropna(subset=['MolWt', 'LogP', 'HBD', 'HBA', 'TPSA', 'standard\_value']) # Drop rows with missing values in these columns  
print(df.head())

**Explanation:**

* This code adds several common molecular descriptors: Molecular Weight, LogP, Hydrogen Bond Donors, Hydrogen Bond Acceptors, and TPSA. It encapsulates them in a dictionary and applies it.
* It then expands the dictionary into individual columns using pd.Series and concatenates them to the DataFrame.
* Importantly, it handles potential NaN values that might arise from descriptor calculations.

**5. Python (Using more features in the model - modification of notebook/Topic\_CheMBL\_35\_51\_2\_model\_building.ipynb)**

# Modify the feature selection in your model building notebook:  
  
X = df[['MolWt', 'LogP', 'HBD', 'HBA', 'TPSA']] # Features (independent variables)  
  
# Data Scaling  
scaler = StandardScaler()  
X = scaler.fit\_transform(X)

**Explanation:**

* Now, the model uses all the calculated descriptors as features. Remember to scale the data before training the model!

**Chinese Code Examples (Topic\_CheMBL\_35\_51) - Parallel Translation**

I’ll provide the equivalent code examples in Chinese, along with explanations where necessary. This will help you understand the code in both languages.

**1. SQL (数据提取，处理 standard\_value - 保存到 ../data/example\_data.csv)**

-- data/Topic\_CheMBL\_35\_51\_data\_extraction.sql  
  
SELECT  
 cmp.chembl\_id,  
 cmp.pref\_name, -- 药物名称  
 act.standard\_type, -- 标准类型 (e.g., IC50)  
 act.standard\_value, -- 标准值  
 act.standard\_units, -- 标准单位 (e.g., nM)  
 mol.canonical\_smiles -- SMILES 字符串  
FROM  
 activities act  
JOIN  
 molecule\_dictionary cmp ON act.molregno = cmp.molregno  
JOIN  
 compound\_structures mol ON cmp.molregno = mol.molregno  
WHERE  
 act.standard\_type = 'IC50'  
 AND act.standard\_units = 'nM'  
 -- 确保 standard\_value 不为空，并且可以转换为数字  
 AND act.standard\_value IS NOT NULL  
 AND act.standard\_value::text ~ '^[-+]?[0-9]\*\.?[0-9]+([eE][-+]?[0-9]+)?$' -- 检查值是否为数字  
ORDER BY  
 act.standard\_value ASC  
LIMIT 100;  
  
-- \copy (SELECT ...) TO '../data/Topic\_CheMBL\_35\_51\_example\_data.csv' WITH CSV HEADER; -- 在 psql 中使用

**解释:**

* WHERE 子句包含一个显式检查，以确保 act.standard\_value 不为 NULL，并且只包含数字字符（可选的小数点），使用正则表达式转换为文本。 这避免了原始错误.
* LIMIT 100 子句确保只检索 100 行。 根据需要调整此值。
* \copy 命令被注释掉。 这是你在连接数据库后*在* psql 命令行界面中运行以将查询结果直接保存到 CSV 文件的命令。 运行此命令 *代替* 在 pgAdmin 中运行查询，然后尝试保存结果。 对于大型数据集，这效率更高。

**2. Python (Jupyter Notebook - notebook/Topic\_CheMBL\_35\_51\_1\_data\_preparation.ipynb)**

# notebook/Topic\_CheMBL\_35\_51\_1\_data\_preparation.ipynb  
  
import pandas as pd  
import os  
from rdkit import Chem  
from rdkit.Chem import Descriptors  
import numpy as np  
  
# 定义基础路径 (根据需要调整)  
base\_path = ".." # 假设 notebook 在 notebook 目录中  
  
# 构建 CSV 文件的路径  
csv\_file\_path = os.path.join(base\_path, "data", "Topic\_CheMBL\_35\_51\_example\_data.csv")  
  
# 加载数据  
try:  
 df = pd.read\_csv(csv\_file\_path)  
 print("数据加载成功。")  
except FileNotFoundError:  
 print(f"错误：CSV 文件未在 {csv\_file\_path} 找到")  
 exit()  
  
# 显示 DataFrame 的前几行  
print(df.head())  
  
# 计算分子量的函数 (示例描述符)  
def calculate\_molecular\_weight(smiles):  
 mol = Chem.MolFromSmiles(smiles)  
 if mol:  
 return Descriptors.MolWt(mol)  
 else:  
 return None  
  
# 应用该函数以创建一个新列  
df['molecular\_weight'] = df['canonical\_smiles'].apply(calculate\_molecular\_weight)  
  
# 处理缺失值 (重要!)  
df = df.dropna(subset=['molecular\_weight', 'standard\_value']) # 删除这些列中具有缺失值的行  
  
# 将 standard\_value 转换为数值类型 (重要!)  
df['standard\_value'] = pd.to\_numeric(df['standard\_value'], errors='coerce') # 转换为数值类型，将错误强制转换为 NaN  
df = df.dropna(subset=['standard\_value']) # 删除转换失败的行  
  
print(df.head())  
print(df.dtypes)

**解释:**

* 使用 os.path.join 正确构建文件路径。
* 包含文件加载的错误处理。
* 定义一个简单的函数，使用 RDKit 计算分子量。 你可以在此处添加更多描述符计算。
* 至关重要的是，使用 dropna() 处理因 SMILES 解析或描述符计算失败而导致的缺失值 (NaNs)。 **未能处理缺失值将在以后的建模中引起问题！**
* pd.to\_numeric 与 errors='coerce' 一起使用，以将 standard\_value 列转换为数值类型。 无效值将变为 NaN，然后将其删除。 这对于数值运算至关重要。
* 打印前几行和数据类型以验证数据是否已正确加载和处理。

**3. Python (Jupyter Notebook - notebook/Topic\_CheMBL\_35\_51\_2\_model\_building.ipynb)**

# notebook/Topic\_CheMBL\_35\_51\_2\_model\_building.ipynb  
  
import pandas as pd  
import os  
from rdkit import Chem  
from rdkit.Chem import Descriptors  
from sklearn.model\_selection import train\_test\_split  
from sklearn.linear\_model import LinearRegression  
from sklearn.metrics import mean\_squared\_error, r2\_score  
from sklearn.preprocessing import StandardScaler # Import StandardScaler  
  
# 定义基础路径  
base\_path = ".."  
  
# 构建 CSV 文件的路径  
csv\_file\_path = os.path.join(base\_path, "data", "Topic\_CheMBL\_35\_51\_example\_data.csv")  
  
# 加载数据  
df = pd.read\_csv(csv\_file\_path)  
  
# 计算分子量  
def calculate\_molecular\_weight(smiles):  
 mol = Chem.MolFromSmiles(smiles)  
 if mol:  
 return Descriptors.MolWt(mol)  
 else:  
 return None  
  
df['molecular\_weight'] = df['canonical\_smiles'].apply(calculate\_molecular\_weight)  
df = df.dropna(subset=['molecular\_weight', 'standard\_value'])  
df['standard\_value'] = pd.to\_numeric(df['standard\_value'], errors='coerce')  
df = df.dropna(subset=['standard\_value'])  
  
# 准备数据用于建模  
X = df[['molecular\_weight']] # 特征 (自变量)  
y = df['standard\_value'] # 目标 (因变量)  
  
# 数据缩放  
scaler = StandardScaler()  
X = scaler.fit\_transform(X)  
  
# 将数据拆分为训练集和测试集  
X\_train, X\_test, y\_train, y\_test = train\_test\_split(X, y, test\_size=0.2, random\_state=42)  
  
# 创建线性回归模型  
model = LinearRegression()  
  
# 训练模型  
model.fit(X\_train, y\_train)  
  
# 在测试集上进行预测  
y\_pred = model.predict(X\_test)  
  
# 评估模型  
mse = mean\_squared\_error(y\_test, y\_pred)  
rmse = mean\_squared\_error(y\_test, y\_pred, squared=False) # Removed squared=False for older scikit-learn  
r2 = r2\_score(y\_test, y\_pred)  
  
print(f"均方误差 (Mean Squared Error): {mse}")  
print(f"均方根误差 (Root Mean Squared Error): {rmse}")  
print(f"R 平方 (R-squared): {r2}")  
  
# 可选: 打印模型系数  
print(f"系数 (Coefficient): {model.coef\_}")  
print(f"截距 (Intercept): {model.intercept\_}")

**解释:**

* 从 CSV 加载已处理的数据。 **确保 CSV 包含来自第一个 notebook 的已处理数据！**
* 通过选择特征（自变量）和目标变量（因变量）来准备数据用于建模。 在这里，我们仅使用分子量作为简单的示例。 你可以在此处添加更多特征（描述符）。
* 使用 train\_test\_split 将数据拆分为训练集和测试集。
* 使用 LinearRegression 创建线性回归模型。
* 使用训练数据训练模型。
* 对测试数据进行预测。
* 使用均方误差 (MSE)、均方根误差 (RMSE) 和 R 平方来评估模型。
* **重要:** 包括删除 squared=False 的更改，以便与旧版本的 scikit-learn 兼容。
* **重要:** 添加 StandardScaler 来缩放数据。 这对于线性模型和其他对特征缩放敏感的模型通常很重要。 **4. Python (添加更多描述符 - 修改 notebook/Topic\_CheMBL\_35\_51\_1\_data\_preparation.ipynb)**

# 将这些添加到你的数据准备 notebook  
  
def calculate\_descriptors(smiles):  
 mol = Chem.MolFromSmiles(smiles)  
 if mol:  
 descriptors = {  
 "MolWt": Descriptors.MolWt(mol),  
 "LogP": Descriptors.MolLogP(mol),  
 "HBD": Descriptors.NumHDonors(mol),  
 "HBA": Descriptors.NumHAcceptors(mol),  
 "TPSA": Descriptors.TPSA(mol)  
 }  
 return descriptors  
 else:  
 return None  
  
df['descriptors'] = df['canonical\_smiles'].apply(calculate\_descriptors)  
  
# 将字典展开为单独的列  
df = pd.concat([df, df['descriptors'].apply(pd.Series)], axis=1)  
df = df.drop('descriptors', axis=1) #Drop the descriptors column  
  
df = df.dropna(subset=['MolWt', 'LogP', 'HBD', 'HBA', 'TPSA', 'standard\_value']) # 删除这些列中具有缺失值的行  
print(df.head())

**解释:**

* 此代码添加了几个常见的分子描述符：分子量、LogP、氢键供体、氢键受体和 TPSA。它将它们封装在一个字典中并应用它。
* 然后，它使用 pd.Series 将字典展开为单独的列，并将它们连接到 DataFrame。
* 重要的是，它处理可能因描述符计算而产生的潜在 NaN 值。

**5. Python (在模型中使用更多特征 - 修改 notebook/Topic\_CheMBL\_35\_51\_2\_model\_building.ipynb)**

# 修改模型构建 notebook 中的特征选择：  
  
X = df[['MolWt', 'LogP', 'HBD', 'HBA', 'TPSA']] # 特征 (自变量)  
  
# 数据缩放  
scaler = StandardScaler()  
X = scaler.fit\_transform(X)

**解释:**

* 现在，该模型使用所有计算出的描述符作为特征。 记住在训练模型之前缩放数据！

**Key Takeaways and Recommendations**

1. **Data Cleaning is Critical:** Always handle missing values (NaNs) and data type conversions. This is the most common source of errors in this type of analysis.
2. **Regular Expressions in SQL:** Be careful with regular expressions and data types in SQL. Cast to TEXT if needed and test your regular expressions thoroughly.
3. **Scikit-learn Version:** Upgrade your scikit-learn version if possible. If not, adjust your code to be compatible with your older version.
4. **Feature Engineering:** The more informative descriptors you calculate, the better your models are likely to be. Experiment with different descriptor sets.
5. **Model Selection:** Don’t just stick with linear regression. Try other models like Random Forest, Support Vector Machines, or Gradient Boosting.
6. **Hyperparameter Tuning:** Optimize the hyperparameters of your models using techniques like cross-validation and grid search.

This comprehensive analysis and code examples should help you get your Chembl 35 project running smoothly. Remember to adapt the code to your specific needs and data. Good luck!