

AI-Driven Drug Discovery: A Hands-On Guide

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2025-12-02

Table of contents

1 Home	6
I Part 0: Getting Started	7
2 Preface	8
3 Preface	9
4 Introduction	10
5 Introduction	11
6 High-Level Overview: Steps of Drug Discovery	12
7 Chapter 1: Genomics to Molecules	14
7.1 Workflow Overview	14
7.2 Summary	15
7.3 Key Code Snippets	15
7.4 Further Reading	15
7.4.1 1. Retrieve ALS Target Genes from Open Targets	16
7.4.2 2. Batch Extract Non-Protein Entities from PDB	16
7.4.3 3. Find Known Ligands for a Target in ChEMBL	17
7.5 Notebook Link	17
7.6 1.1 Introduction to Genomics in Drug Discovery	17
7.7 1.2 From Genes to Proteins	17
7.8 1.3 Target Identification and Validation	18
7.9 1.4 Linking Targets to Molecules	18
7.10 1.5 Case Study: ALS Gene to Ligand	18
7.11 Interpretation of Exported Ligands for ALS Drug Discovery	18
8 Chapter 2: Environment Setup	20
8.1 2.1 Installing Python and Conda	20
8.2 2.2 Setting Up Jupyter Notebooks	20

8.3	2.3 Managing Dependencies	21
8.4	2.4 Version Control with Git	21
II	Part I: Foundations	22
9	Part I: Foundations	23
10	Part I: Foundations	24
11	Chapter 3: Cheminformatics Essentials	25
11.1	3.1 Molecular Representations (SMILES, InChI)	25
11.2	3.2 Chemical File Formats	25
11.3	3.3 Structure Search and Similarity	26
11.4	3.4 Cheminformatics Toolkits	26
12	Chapter 4: Molecular Fingerprints	28
12.1	4.1 What Are Molecular Fingerprints?	28
12.2	4.2 Types: MACCS, Morgan, Topological	28
12.3	4.3 Applications in Virtual Screening	29
12.4	4.4 Fingerprints in Machine Learning	29
13	Chapter 5: Molecular Properties	31
13.1	5.1 Physicochemical Descriptors	31
13.2	5.2 Calculating Properties with RDKit	31
13.3	5.3 Lipinski's Rule of Five	32
13.4	5.4 Property-Based Filtering	32
III	Part II: Machine Learning for Molecules	34
14	Chapter 6: Graph Neural Networks	35
14.1	Summary	35
14.2	Key Code Snippets	35
14.2.1	1. Construct a Molecular Graph with RDKit and NetworkX	35
14.2.2	2. Simple GNN Layer with PyTorch Geometric	36
14.3	Notebook Link	36
14.4	6.1 Introduction to GNNs	36
14.5	6.2 Molecular Graph Construction	36
14.6	6.3 GNN Architectures for Chemistry	37
14.7	6.4 Case Study: GNN for Solubility	37
15	Chapter 7: Bioactivity Prediction	38
15.1	Summary	38
15.2	Key Code Snippets	38
15.2.1	1. Load Bioactivity Data from ChEMBL	38
15.2.2	2. Train a Classifier for Bioactivity	38

15.3	Notebook Link	39
15.4	7.1 Bioactivity Data Sources	39
15.5	7.2 Feature Engineering for Bioactivity	39
15.6	7.3 ML Models for Activity Prediction	39
15.7	7.4 Model Evaluation and Validation	40
16	Chapter 8: ADMET Prediction	41
16.1	Summary	41
16.2	Key Code Snippets	41
16.2.1	1. Calculate ADMET Descriptors with RDKit	41
16.2.2	2. Predict Toxicity with a Pretrained Model (Example)	42
16.3	Notebook Link	42
16.4	8.1 Introduction to ADMET	42
16.5	8.2 Data and Descriptors	42
16.6	8.3 ML Models for ADMET	43
16.7	8.4 Interpreting ADMET Predictions	43
17	Chapter 9: Molecular Generation	44
17.1	Summary	44
17.2	Key Code Snippets	44
17.2.1	1. Generate Molecules with a SMILES RNN (Example)	44
17.2.2	2. Evaluate Validity of Generated SMILES	44
17.3	Notebook Link	45
17.4	9.1 Generative Models Overview	45
17.5	9.2 SMILES-Based Generation	45
17.6	9.3 Graph-Based Generation	45
17.7	9.4 Evaluating Generated Molecules	46
18	Chapter 10: Conditional Generation	47
18.1	10.1 Conditional Generative Models	47
18.2	10.2 Property-Conditioned Generation	49
18.3	10.3 Reinforcement Learning Approaches	50
19	Chapter 11: Transformers in Chemistry	53
19.1	11.1 Introduction to Transformers	53
19.2	11.2 SMILES Transformers	53
19.3	11.3 Reaction Prediction	54
19.4	11.4 Property Prediction	54
IV	Part III: Structure-Based Drug Design	55
20	Chapter 12: Protein Structure	56
20.1	12.1 Protein Structure Levels	56
20.2	12.2 Experimental Determination	56
20.3	12.3 Structure Prediction Tools	57

20.4 12.4 Protein-Ligand Interactions	57
21 Chapter 13: Molecular Docking	58
21.1 13.1 Docking Theory	58
21.2 13.2 Docking Software (AutoDock, Vina)	58
21.3 13.3 Preparing Structures for Docking	59
21.4 13.4 Analyzing Docking Results	59
22 Chapter 14: ML-Enhanced Docking	60
22.1 14.1 ML Scoring Functions	60
22.2 14.2 Pose Prediction	61
22.3 14.3 Integrating ML with Docking	61
V Part IV: Synthesis and Advanced Topics	62
23 Chapter 15: Retrosynthesis	63
23.1 15.1 Introduction to Retrosynthesis	63
23.2 15.2 Rule-Based Approaches	63
23.3 15.3 AI-Driven Retrosynthesis	64
23.4 15.4 Evaluating Synthetic Routes	64
24 Chapter 16: Multi-Modal Learning	66
24.1 16.1 What is Multi-Modal Learning?	66
24.2 16.2 Data Integration Strategies	66
24.3 16.3 Applications in Drug Discovery	67
25 Chapter 17: End-to-End Pipeline	68
25.1 17.1 Pipeline Overview	68
25.2 17.2 Data Preparation	68
25.3 17.3 Model Training and Evaluation	69
25.4 17.4 Candidate Selection	69
26 Chapter 18: Case Studies	70
26.1 18.1 Case Study 1: ALS Drug Discovery	70
26.2 18.2 Case Study 2: Kinase Inhibitors	70
26.3 18.3 Lessons Learned	71
27 Chapter 19: External Tools	72
27.1 19.1 Structural Databases	72
27.2 19.2 Bioactivity Databases	72
27.3 19.3 Workflow Automation Tools	73
28 Chapter 20: Ethical Considerations	74
28.1 20.1 Data Privacy and Security	74
28.2 20.2 Bias in AI Models	74
28.3 20.3 Responsible Innovation	75

29 Chapter 21: Future Next Steps	76
29.1 21.1 Emerging Technologies	76
29.2 21.2 Open Science and Collaboration	76
29.3 21.3 Next Steps for Practitioners	77
30 Conclusion	78
31 Conclusion	79
References	80
Further Reading	82

Chapter 1

Home

Welcome to the AI Drug Discovery Book! Use the navigation to explore chapters on drug discovery, machine learning, and computational tools.

Part I

Part 0: Getting Started

Chapter 2

Preface

Chapter 3

Preface

Welcome to “AI-Driven Drug Discovery: A Hands-On Guide.” This book is designed for data scientists, bioinformaticians, and machine learning engineers who want to apply their skills to the world of drug discovery. Our goal is to provide a practical, code-first approach to modern computational drug discovery, with real-world examples and hands-on exercises throughout.

We hope this book empowers you to make meaningful contributions to the future of medicine.

Chapter 4

Introduction

Chapter 5

Introduction

Artificial intelligence is revolutionizing drug discovery. This book introduces the key concepts, tools, and workflows that are shaping the future of pharmaceutical research. You will learn how to leverage AI, machine learning, and public biomedical data to accelerate the discovery of new medicines.

We begin with foundational concepts and build up to advanced, real-world applications, ensuring you gain both theoretical understanding and practical skills.

Chapter 6

High-Level Overview: Steps of Drug Discovery

Drug discovery is a multi-stage process that transforms basic research into new medicines. The main steps are:

1. **Target Identification & Validation**
Understanding disease biology and identifying molecular targets (e.g., genes, proteins).
2. **Hit Discovery & Screening**
Finding molecules that interact with the target (e.g., high-throughput or virtual screening).
3. **Lead Optimization**
Refining hits for potency, selectivity, and drug-like properties (e.g., Lipinski's rules, ADME).
4. **Preclinical Testing**
Assessing safety and efficacy in vitro and in animal models.
5. **Clinical Trials**
Testing in humans (Phases I-III).
6. **Regulatory Approval & Post-Market Surveillance**
Submitting data to agencies (e.g., FDA), monitoring real-world use.

Book Chapter Mapping: - Chapters 1–2: Target identification, genomics, and molecules - Chapters 3–5: Screening, cheminformatics, molecular properties - Chapters 6–10: Optimization, validation, preclinical studies - Chapters 11–14: Clinical trials, regulatory science, post-market studies

Online Diagrams and Resources: - [Drug Discovery Cycle Schematic](#) (Wikipedia, SVG, open license) - [FDA: Drug Development Process Overview](#) - [ChEMBL Visualizations](#) - [ResearchGate: Drug Discovery and Development Process](#)

You can click these links for interactive diagrams and further reading.

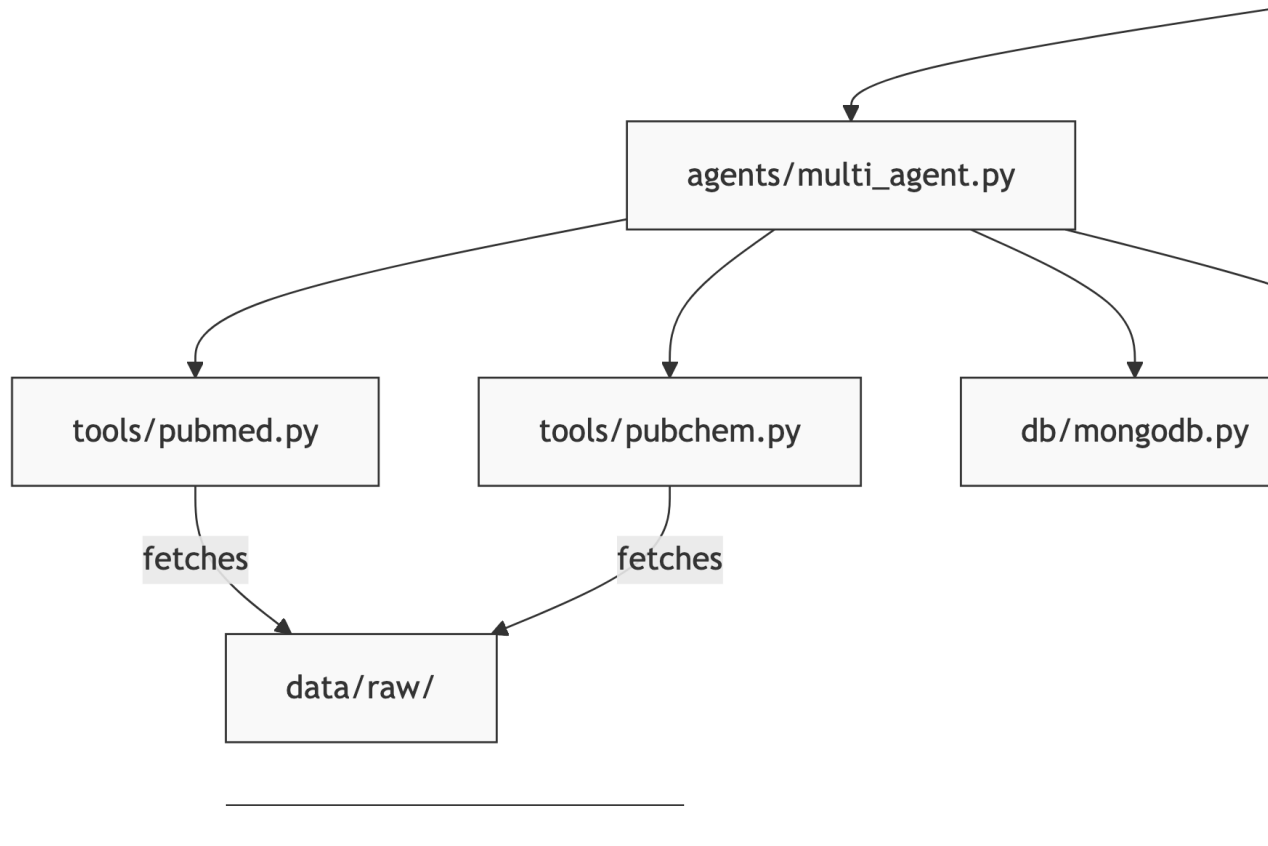
Chapter 7

Chapter 1: Genomics to Molecules

This chapter introduces the journey from genomics data to actionable molecular targets in drug discovery. It covers how genetic information is translated into protein targets and how these are linked to small molecules for therapeutic intervention.

7.1 Workflow Overview

The following diagram illustrates the overall process from genomics data to actionable molecules in drug discovery:



7.2 Summary

This workflow demonstrates how to start from a gene (e.g., SOD1 for ALS), find the corresponding protein (UniProt), retrieve known small-molecule ligands (ChEMBL), and analyze their molecular properties. This bridges genomics and cheminformatics, enabling rational drug discovery starting from genetic information.

7.3 Key Code Snippets

7.4 Further Reading

@opentargets @uniprot @chembl @pdb

7.4.1 1. Retrieve ALS Target Genes from Open Targets

```
import requests
efo_id = "Orphanet_803" # ALS
ot_url = f"https://platform-api.opentargets.io/v3/platform/public/association/filter?disease={efo_id}"
resp = requests.get(ot_url)
gene_hits = resp.json().get('data', [])
als_genes = [g['target']['gene_info']['symbol'] for g in gene_hits]
print('Top ALS target genes:', als_genes)
```

7.4.2 2. Batch Extract Non-Protein Entities from PDB

```
import requests, pandas as pd
pdb_ids = ['2RSQ', '4RFX', '7WWT', ...] # your PDB list
all_results = []
for pdb_id in pdb_ids:
    url = f'https://data.rcsb.org/rest/v1/core/entry/{pdb_id}'
    resp = requests.get(url)
    if resp.status_code == 200:
        data = resp.json()
        for entity in data.get('nonpolymer_entities', []):
            chem_comp = entity.get('chem_comp', {})
            all_results.append({
                'pdb_id': pdb_id,
                ...
            })
## Interpretation of Exported Ligands for ALS Drug Discovery

At the end of this workflow, the notebook exports a file `ligands_for_docking`
```

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name	smiles
CHEMBL273030	c1ccc2cc3c(NCCc4c[nH]cn4)nnc(NCCc4c[nH]cn4)c3cc2c1
CHEMBL272808	Clc1nnC(NCCc2c[nH]cn2)c2cc3cccc3cc12
CHEMBL272641	c1ccc2cc3c(NCCCN4ccnc4)nnc(NCCCN4ccnc4)c3cc2c1
CHEMBL405899	Clc1nnC(NCCN2ccnc2)c2cc3cccc3cc12
CHEMBL1672028	c1ccc2cc3c(Nc4cc[nH]n4)nnc(Nc4cc[nH]n4)c3cc2c1

****How to interpret these results:****

- These ligands are ChEMBL compounds with known bioactivity, retrieved for this study.
 - Their structures are consistent with CNS drug-like scaffolds, but further validation is required.
 - These molecules are not validated ALS drugs, but are reasonable starting points for screening.
 - The CSV can be used directly in docking or cheminformatics pipelines to predict binding affinity.

```

**Limitations:**
- The activity data is typically from in vitro assays, not ALS disease model
- Further experimental validation is required to confirm efficacy and safety

**Summary:**
The notebook provides a practical, automated way to go from ALS gene to acti
'entity_id': chem_comp.get('id', ''),
'entity_name': chem_comp.get('name', ''),
'entity_type': chem_comp.get('type', '')
})
pd.DataFrame(all_results).to_csv('batch_als_pdb_entities.csv', index=False)

```

7.4.3 3. Find Known Ligands for a Target in ChEMBL

```

from chembl_webresource_client.new_client import new_client
target = new_client.target
activity = new_client.activity
acc = 'P00441' # UniProt for SOD1
targets = list(target.filter(target_components__accession=acc))
chembl_id = targets[0]['target_chembl_id']
acts = activity.filter(target_chembl_id=chembl_id, standard_type='IC50')
for a in list(acts)[:5]:
    print(a.get('molecule_chembl_id'), a.get('standard_value'))

```

7.5 Notebook Link

For the full workflow, see the interactive notebook: [chapter1-genomics-to-molecules.ipynb](#)

7.6 1.1 Introduction to Genomics in Drug Discovery

Genomics provides the foundation for modern drug discovery by identifying genes associated with diseases. Advances in sequencing technologies and bioinformatics enable researchers to pinpoint genetic variants and pathways involved in disease mechanisms, offering new opportunities for therapeutic intervention.

7.7 1.2 From Genes to Proteins

Genes encode proteins, which are the functional molecules in cells. Understanding how genetic information is translated into protein structure and function is crucial

for identifying druggable targets. Databases like UniProt provide comprehensive information on protein sequences, structures, and functions.

7.8 1.3 Target Identification and Validation

Target identification involves linking disease-associated genes to specific proteins that can be modulated by drugs. Validation ensures that modulating the target will have a therapeutic effect. This process uses data from genomics, proteomics, and functional studies, and often leverages resources like Open Targets and experimental validation.

7.9 1.4 Linking Targets to Molecules

Once a target protein is identified and validated, the next step is to find small molecules that interact with it. This involves searching bioactivity databases (e.g., ChEMBL) for known ligands, using structure-based methods to identify binding sites, and applying cheminformatics tools to analyze and prioritize compounds.

7.10 1.5 Case Study: ALS Gene to Ligand

Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease with several known genetic risk factors, such as mutations in the SOD1 gene. In this case study, we:

- Identify ALS-associated genes using Open Targets.
- Retrieve the corresponding protein (e.g., SOD1) from UniProt.
- Find known ligands for SOD1 in ChEMBL.
- Analyze ligand properties and prepare them for virtual screening or further optimization.

This workflow demonstrates how to bridge genomics and cheminformatics for rational drug discovery, starting from a disease gene and ending with actionable small molecules.

7.11 Interpretation of Exported Ligands for ALS Drug Discovery

At the end of this workflow, the notebook exports a file `ligands_for_docking.csv` containing known ligands for the selected ALS target (e.g., SOD1). Here is an example of the exported data:

name	smiles
CHEMBL273030	c1ccc2cc3c(NCCc4c[nH]cn4)nnc(NCCc4c[nH]cn4)c3cc2c1

name	smiles
CHEMBL272808	Clc1nnc(NCCc2c[nH]cn2)c2cc3cccc3cc12
CHEMBL272641	c1ccc2cc3c(NCCCn4ccnc4)nnc(NCCCn4ccnc4)c3cc2c1
CHEMBL405899	Clc1nnc(NCCCn2ccnc2)c2cc3cccc3cc12
CHEMBL16720281	ccc2cc3c(Nc4cc[nH]n4)nnc(Nc4cc[nH]n4)c3cc2c1

How to interpret these results:

- These ligands are ChEMBL compounds with known bioactivity, retrieved for the ALS target SOD1.
- Their structures are consistent with CNS drug-like scaffolds, but further ADMET and blood-brain barrier predictions are needed for ALS relevance.
- These molecules are not validated ALS drugs, but are reasonable starting points for hit discovery and virtual screening.
- The CSV can be used directly in docking or cheminformatics pipelines to prioritize or design new compounds.

Limitations: - The activity data is typically from in vitro assays, not ALS disease models. - Further experimental validation is required to confirm efficacy and safety in ALS.

Summary: The notebook provides a practical, automated way to go from ALS gene to actionable ligand lists for computational drug discovery, enabling rapid hypothesis generation and screening for ALS research.

Chapter 8

Chapter 2: Environment Setup

This chapter guides you through setting up a reproducible computational environment for AI-driven drug discovery, including Python, Jupyter, and essential libraries.

8.1 2.1 Installing Python and Conda

To get started, you need Python and a package manager. Conda is recommended for managing environments and dependencies.

Steps:

1. **Download and Install Miniconda:**

- Go to [Miniconda Downloads](#) and download the installer for your OS.
- Run the installer and follow the prompts.

2. **Verify Installation:**

- Open a terminal and run:

```
conda --version
```

- You should see the conda version printed.

3. **Create a New Environment:**

- Example for Python 3.10:

```
conda create -n drug-discovery python=3.10
conda activate drug-discovery
```

8.2 2.2 Setting Up Jupyter Notebooks

Jupyter Notebooks are essential for interactive coding and documentation.

Steps:

1. **Install Jupyter:** bash conda install jupyterlab
2. **Launch JupyterLab:** bash jupyter lab
 - This will open JupyterLab in your browser.
3. **Create a New Notebook:**
 - In JupyterLab, click “Python 3” under the Notebook section to start a new notebook.

8.3 2.3 Managing Dependencies

Managing dependencies ensures reproducibility and avoids conflicts.

Best Practices:

- Use conda or pip to install packages:
`conda install numpy pandas scikit-learn
pip install rdkit matplotlib`
- Export your environment for sharing:
`conda env export > environment.yml`
- Recreate an environment from a file:
`conda env create -f environment.yml`

8.4 2.4 Version Control with Git

Version control is critical for tracking changes and collaborating.

Steps:

1. **Install Git:**
 - Download from git-scm.com and follow installation instructions for your OS.
2. **Configure Git:** bash git config --global user.name "Your Name" git config --global user.email "you@example.com"
3. **Initialize a Repository:** bash git init
4. **Basic Workflow:** bash git add . git commit -m "Initial commit" git remote add origin <your-repo-url> git push -u origin main

Part II

Part I: Foundations

Chapter 9

Part I: Foundations

Chapter 10

Part I: Foundations

This part introduces the essential concepts and tools for AI-driven drug discovery, including cheminformatics, molecular representations, and compound screening.

Chapter 11

Chapter 3: Cheminformatics Essentials

Cheminformatics is the foundation for representing, searching, and analyzing chemical structures computationally. This chapter covers molecular representations, file formats, and basic cheminformatics operations.

11.1 3.1 Molecular Representations (SMILES, InChI)

Molecular representations are ways to describe chemical structures in a form that computers can process.

- **SMILES (Simplified Molecular Input Line Entry System):**
 - A line notation for describing the structure of chemical species using short ASCII strings.
 - Example: Ethanol = CCO, Benzene = c1ccccc1
 - Widely used for database storage and cheminformatics tools.
- **InChI (International Chemical Identifier):**
 - A textual identifier for chemical substances, designed to be unique and non-proprietary.
 - Example: Ethanol = InChI=1S/C2H6O/c1-2-3/h3H,2H2,1H3
 - Useful for interoperability and linking chemical information across databases.

11.2 3.2 Chemical File Formats

Chemical file formats store molecular structures and related data. Common formats include:

- **SDF (Structure Data File):**
 - Stores multiple molecules with associated data fields.
 - Used for compound libraries and virtual screening.
- **MOL/MOL2:**
 - Encodes 2D/3D structure, atom types, and bond information.
 - MOL2 supports more detailed atom typing and charges.
- **PDB (Protein Data Bank):**
 - Used for macromolecules (proteins, nucleic acids) and some small molecules.
 - Contains 3D coordinates and metadata.
- **CSV/TSV:**
 - Tabular formats often used to store SMILES or InChI strings with associated data.

11.3 3.3 Structure Search and Similarity

Cheminformatics enables searching for molecules by structure and comparing their similarity.

- **Exact Structure Search:**
 - Finds molecules that are identical to a query structure.
- **Substructure Search:**
 - Finds molecules containing a specific substructure (e.g., a benzene ring).
- **Similarity Search:**
 - Finds molecules similar to a query, often using fingerprints and Tanimoto similarity.
 - Example: RDKit can compute molecular fingerprints and similarity scores.

Example (Python/RDKit):

```
from rdkit import Chem
from rdkit.Chem import DataStructs, AllChem

mol1 = Chem.MolFromSmiles('CCO')
mol2 = Chem.MolFromSmiles('CCN')
fp1 = AllChem.GetMorganFingerprintAsBitVect(mol1, 2)
fp2 = AllChem.GetMorganFingerprintAsBitVect(mol2, 2)
similarity = DataStructs.TanimotoSimilarity(fp1, fp2)
print(f"Tanimoto similarity: {similarity:.2f}")
```

11.4 3.4 Cheminformatics Toolkits

Several open-source toolkits provide cheminformatics functionality:

- **RDKit:**
 - Python/C++ library for molecular manipulation, descriptor calculation, and visualization.
 - [RDKit Documentation](#)
- **Open Babel:**
 - Command-line and Python tools for converting between file formats and basic cheminformatics operations.
 - [Open Babel](#)
- **CDK (Chemistry Development Kit):**
 - Java library for cheminformatics, used in many bioinformatics tools.

These toolkits support reading/writing file formats, structure search, descriptor calculation, and more.

Chapter 12

Chapter 4: Molecular Fingerprints

Molecular fingerprints are compact representations of chemical structures used for similarity searching and machine learning. This chapter explains different types of fingerprints and their applications.

12.1 4.1 What Are Molecular Fingerprints?

Molecular fingerprints are binary or integer vectors that encode the presence or absence of particular substructures or features in a molecule. They allow rapid comparison of molecules for similarity searching, clustering, and as input features for machine learning models.

- **Purpose:**
 - Enable fast similarity searches in large chemical databases.
 - Provide a standardized input for cheminformatics and AI tasks.
- **How They Work:**
 - Each bit or value in the fingerprint corresponds to a specific structural feature or pattern.
 - The fingerprint is generated by analyzing the molecule's structure and mapping features to bits.

12.2 4.2 Types: MACCS, Morgan, Topological

There are several types of molecular fingerprints, each with different algorithms and applications:

- **MACCS Keys:**

- 166-bit fingerprint where each bit represents a predefined substructure (e.g., aromatic ring, carboxyl group).
- Simple and interpretable, commonly used for basic similarity searches.
- **Morgan (Circular) Fingerprints:**
 - Also known as ECFP (Extended-Connectivity Fingerprints).
 - Encodes atom environments up to a certain radius (e.g., ECFP4 uses radius 2).
 - Widely used in machine learning and virtual screening.
 - Example (Python/RDKit):

```
python      from rdkit import
Chem      from rdkit.Chem import AllChem      mol =
Chem.MolFromSmiles('CCO')      fp = AllChem.GetMorganFingerprintAsBitVect(mol,
2, nBits=1024)      print(list(fp))
```
- **Topological (Path-Based) Fingerprints:**
 - Encodes the presence of linear paths of atoms (e.g., Daylight fingerprints).
 - Good for capturing connectivity and substructure patterns.

12.3 4.3 Applications in Virtual Screening

Fingerprints are essential for virtual screening, where large libraries of compounds are searched for molecules similar to a known active compound.

- **Similarity Searching:**
 - Compare fingerprints using metrics like Tanimoto similarity to find similar molecules.
- **Substructure Searching:**
 - Identify compounds containing specific structural motifs.
- **Library Design:**
 - Cluster compounds based on fingerprint similarity to ensure chemical diversity.

12.4 4.4 Fingerprints in Machine Learning

Fingerprints are widely used as input features for machine learning models in cheminformatics.

- **Feature Vectors:**
 - Each molecule is represented by its fingerprint vector, which can be used as input to algorithms like random forests, SVMs, or neural networks.
- **QSAR Modeling:**
 - Quantitative Structure-Activity Relationship (QSAR) models use fingerprints to predict biological activity or properties.
- **Deep Learning:**
 - Fingerprints can be combined with other descriptors or used as input

to deep learning models for property prediction, classification, or regression tasks.

Example (Python/Scikit-learn):

```
from rdkit import Chem
from rdkit.Chem import AllChem
from sklearn.ensemble import RandomForestClassifier
import numpy as np

# Example molecules and labels
smiles = ['CCO', 'CCN', 'CCC', 'CCCl']
labels = [1, 0, 1, 0]
fps = [AllChem.GetMorganFingerprintAsBitVect(Chem.MolFromSmiles(s), 2, nBits=1024) for s in
X = np.array([list(fp) for fp in fps])

clf = RandomForestClassifier()
clf.fit(X, labels)
```

Chapter 13

Chapter 5: Molecular Properties

This chapter explores key physicochemical properties of molecules (e.g., logP, MW, H-bond donors/acceptors) and their importance in drug design and ADMET prediction.

13.1 5.1 Physicochemical Descriptors

Physicochemical descriptors are numerical values that capture important chemical and physical properties of molecules. These descriptors are crucial for understanding drug-likeness, solubility, permeability, and ADMET (Absorption, Distribution, Metabolism, Excretion, Toxicity) properties.

- **Common Descriptors:**
 - **Molecular Weight (MW):** Total mass of a molecule.
 - **LogP:** Partition coefficient between octanol and water, indicating hydrophobicity.
 - **Hydrogen Bond Donors (HBD):** Number of hydrogen atoms attached to electronegative atoms (e.g., N-H, O-H).
 - **Hydrogen Bond Acceptors (HBA):** Number of electronegative atoms with lone pairs (e.g., N, O).
 - **Topological Polar Surface Area (TPSA):** Surface area contributed by polar atoms, related to absorption and permeability.

13.2 5.2 Calculating Properties with RDKit

RDKit is a popular cheminformatics toolkit for calculating molecular properties in Python.

Example (Python/RDKit):

```
from rdkit import Chem
from rdkit.Chem import Descriptors

smiles = 'CCO' # Ethanol
mol = Chem.MolFromSmiles(smiles)
mw = Descriptors.MolWt(mol)
logP = Descriptors.MolLogP(mol)
hbd = Descriptors.NumHDonors(mol)
hba = Descriptors.NumHAcceptors(mol)
tpsa = Descriptors.TPSA(mol)

print(f"MW: {mw:.2f}, LogP: {logP:.2f}, HBD: {hbd}, HBA: {hba}, TPSA: {tpsa:.2f}")
```

13.3 5.3 Lipinski's Rule of Five

Lipinski's Rule of Five is a set of guidelines to evaluate drug-likeness and oral bioavailability:

- Rules:
 - MW 500 Da
 - LogP 5
 - HBD 5
 - HBA 10

Compounds that violate more than one rule are less likely to be orally active drugs.

Application: Use these rules to filter compound libraries for drug discovery projects.

13.4 5.4 Property-Based Filtering

Property-based filtering is used to select compounds with desirable physicochemical properties for further study.

- Filtering Strategies:
 - Remove compounds with poor solubility or permeability.
 - Apply Lipinski's rules and other property thresholds (e.g., TPSA, rotatable bonds).
- Example (Python/RDKit):

```
from rdkit import Chem
from rdkit.Chem import Descriptors

def passes_lipinski(mol):
```

```
mw = Descriptors.MolWt(mol)
logP = Descriptors.MolLogP(mol)
hbd = Descriptors.NumHDonors(mol)
hba = Descriptors.NumHAcceptors(mol)
return (mw <= 500 and logP <= 5 and hbd <= 5 and hba <= 10)

smiles_list = ['CCO', 'CCN(CC)CC', 'CCCCCCCCCC(=O)O']
for smi in smiles_list:
    mol = Chem.MolFromSmiles(smi)
    if passes_lipinski(mol):
        print(f"{smi} passes Lipinski's rules")
    else:
        print(f"{smi} fails Lipinski's rules")
```

Part III

Part II: Machine Learning for Molecules

Chapter 14

Chapter 6: Graph Neural Networks

14.1 Summary

Graph Neural Networks (GNNs) are powerful tools for learning on molecular graphs. This chapter introduces GNN concepts and their application to molecular property prediction, including how molecules are represented as graphs and how GNNs can learn from these structures.

14.2 Key Code Snippets

14.2.1 1. Construct a Molecular Graph with RDKit and NetworkX

```
from rdkit import Chem
import networkx as nx
smiles = "CCO"
mol = Chem.MolFromSmiles(smiles)
G = nx.Graph()
for atom in mol.GetAtoms():
    G.add_node(atom.GetIdx(), label=atom.GetSymbol())
for bond in mol.GetBonds():
    G.add_edge(bond.GetBeginAtomIdx(), bond.GetEndAtomIdx(), type=bond.GetBondType())
```

14.2.2 2. Simple GNN Layer with PyTorch Geometric

```
from torch_geometric.nn import GCNConv
import torch
conv = GCNConv(in_channels=10, out_channels=16)
x = torch.randn((num_nodes, 10))
edge_index = ... # edge list as tensor
out = conv(x, edge_index)
```

14.3 Notebook Link

For a full workflow, see the interactive notebook: (notebook not available for this chapter)

14.4 6.1 Introduction to GNNs

Graph Neural Networks (GNNs) are a class of deep learning models designed to operate on graph-structured data. In chemistry, molecules are naturally represented as graphs, with atoms as nodes and bonds as edges. GNNs can learn to predict molecular properties by aggregating information from neighboring atoms and bonds, capturing both local and global structure.

Key Concepts: - Message passing: Nodes exchange information with neighbors to update their representations. - Permutation invariance: GNNs produce the same output regardless of node ordering. - Applications: Property prediction, molecular generation, reaction prediction.

14.5 6.2 Molecular Graph Construction

To use GNNs, molecules must be converted into graph representations. Each atom becomes a node, and each bond becomes an edge. Node and edge features can include atom types, hybridization, bond order, and more.

Example (Python/RDKit + NetworkX):

```
from rdkit import Chem
import networkx as nx
smiles = "CCO"
mol = Chem.MolFromSmiles(smiles)
G = nx.Graph()
for atom in mol.GetAtoms():
    G.add_node(atom.GetIdx(), label=atom.GetSymbol())
```

```
for bond in mol.GetBonds():
    G.add_edge(bond.GetBeginAtomIdx(), bond.GetEndAtomIdx(), type=bond.GetBondType())
```

14.6 6.3 GNN Architectures for Chemistry

Several GNN architectures are used in molecular modeling:

- **Graph Convolutional Networks (GCN)**: Aggregate features from neighbors using weighted sums.
- **Message Passing Neural Networks (MPNN)**: General framework for message passing and update functions.
- **Graph Attention Networks (GAT)**: Use attention mechanisms to weigh neighbor contributions.
- **Graph Isomorphism Networks (GIN)**: Designed for maximum discriminative power.

Popular Libraries: PyTorch Geometric, DGL, DeepChem.

14.7 6.4 Case Study: GNN for Solubility

In this case study, a GNN is trained to predict aqueous solubility ($\log S$) of small molecules.

Workflow: 1. Prepare a dataset of molecules with known solubility values (e.g., ESOL dataset). 2. Convert SMILES to molecular graphs with atom and bond features. 3. Train a GNN (e.g., MPNN or GCN) to predict $\log S$ from graph representations. 4. Evaluate model performance using RMSE or R² metrics.

Example (Conceptual):

```
# Pseudocode for GNN solubility prediction
from torch_geometric.nn import GCNConv
import torch
class GNN(torch.nn.Module):
    def __init__(self):
        super().__init__()
        self.conv1 = GCNConv(in_channels, hidden)
        self.conv2 = GCNConv(hidden, 1)
    def forward(self, x, edge_index):
        x = self.conv1(x, edge_index).relu()
        x = self.conv2(x, edge_index)
        return x
# Train on molecular graphs and solubility labels
```

Chapter 15

Chapter 7: Bioactivity Prediction

15.1 Summary

Predicting the biological activity of molecules is central to drug discovery. This chapter covers data sources, machine learning models, and evaluation strategies for bioactivity prediction, including how to prepare data and assess model performance.

15.2 Key Code Snippets

15.2.1 1. Load Bioactivity Data from ChEMBL

```
from chembl_webresource_client.new_client import new_client
activity = new_client.activity
acts = activity.filter(target_chembl_id="CHEMBL25", standard_type="IC50")
for a in list(acts)[:5]:
    print(a['molecule_chembl_id'], a['standard_value'])
```

15.2.2 2. Train a Classifier for Bioactivity

```
from sklearn.ensemble import RandomForestClassifier
from sklearn.model_selection import train_test_split
```

```
X = ... # feature matrix
y = ... # binary activity labels
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
clf = RandomForestClassifier()
clf.fit(X_train, y_train)
print("Accuracy:", clf.score(X_test, y_test))
```

15.3 Notebook Link

For a full workflow, see the interactive notebook: (notebook not available for this chapter)

15.4 7.1 Bioactivity Data Sources

Bioactivity data links chemical structures to their effects on biological targets. Key public sources include: - **ChEMBL**: Large database of bioactive molecules with activity data (IC₅₀, Ki, etc.) and assay details. - **PubChem BioAssay**: Repository of biological assay results for small molecules. - **BindingDB**: Focuses on binding affinities for protein-ligand complexes. - **Other sources**: In-house screening data, literature mining, and collaborative consortia.

15.5 7.2 Feature Engineering for Bioactivity

Feature engineering transforms raw molecular data into numerical representations suitable for machine learning. - **Molecular fingerprints**: ECFP (Morgan), MACCS, topological fingerprints. - **Descriptors**: Physicochemical properties (e.g., MW, logP, H-bond donors/acceptors, TPSA). - **Target features**: Protein sequence descriptors, structural features, or target class labels. - **Data preprocessing**: Standardize SMILES, remove duplicates, handle missing values, and balance classes.

15.6 7.3 ML Models for Activity Prediction

Various machine learning models are used to predict bioactivity: - **Random Forests**: Robust to noisy data, handle high-dimensional features well. - **Support Vector Machines (SVM)**: Effective for binary classification with clear margins. - **Neural Networks**: Capture complex, non-linear relationships; can be used for regression or classification. - **Deep Learning**: Graph neural networks (GNNs), convolutional networks for molecular graphs or SMILES. - **Model selection**: Depends on data size, feature type, and interpretability needs.

15.7 7.4 Model Evaluation and Validation

Evaluating model performance is critical for reliable predictions. - **Metrics:** Accuracy, ROC-AUC, precision, recall, F1-score for classification; RMSE, R2 for regression. - **Cross-validation:** Use k-fold or stratified cross-validation to assess generalizability. - **External validation:** Test on independent datasets or prospective experiments. - **Interpretability:** Analyze feature importance, SHAP values, or model explanations to understand predictions.

Chapter 16

Chapter 8: ADMET Prediction

16.1 Summary

ADMET (Absorption, Distribution, Metabolism, Excretion, Toxicity) properties are critical for drug candidates. This chapter discusses computational approaches for ADMET prediction, including data sources, descriptors, and machine learning models.

16.2 Key Code Snippets

16.2.1 1. Calculate ADMET Descriptors with RDKit

```
from rdkit import Chem
from rdkit.Chem import Crippen, Descriptors
smiles = "CCO"
mol = Chem.MolFromSmiles(smiles)
logP = Crippen.MolLogP(mol)
mw = Descriptors.MolWt(mol)
print(f"LogP: {logP}, MW: {mw}")
```

16.2.2 2. Predict Toxicity with a Pretrained Model (Example)

```
# Example only: replace with actual model
import joblib
model = joblib.load('tox_model.pkl')
X = ... # feature matrix
tox_pred = model.predict(X)
```

16.3 Notebook Link

For a full workflow, see the interactive notebook: (notebook not available for this chapter)

16.4 8.1 Introduction to ADMET

ADMET stands for Absorption, Distribution, Metabolism, Excretion, and Toxicity. These properties determine whether a compound is likely to become a safe and effective drug. Poor ADMET characteristics are a major cause of drug candidate failure in clinical trials.

- **Absorption:** How well a drug is taken up into the bloodstream.
- **Distribution:** How the drug spreads through the body and tissues.
- **Metabolism:** How the drug is broken down, often by liver enzymes.
- **Excretion:** How the drug and its metabolites are eliminated.
- **Toxicity:** Potential for harmful effects.

16.5 8.2 Data and Descriptors

ADMET prediction relies on curated datasets and molecular descriptors that capture relevant chemical and biological properties. - **Data sources:** - Public databases: ADMETlab, Tox21, ChEMBL, PubChem. - In-house experimental data. - **Descriptors:** - Physicochemical: MW, logP, TPSA, H-bond donors/acceptors. - Structural: molecular fingerprints, substructure counts. - Biological: predicted or measured interactions with enzymes (e.g., CYP450). - **Data curation:** - Standardize SMILES, remove duplicates, handle missing values, and balance classes.

16.6 8.3 ML Models for ADMET

Machine learning models are widely used for ADMET property prediction. - **Classification models:** Predict binary outcomes (e.g., toxic vs. non-toxic, BBB permeable vs. not). - **Regression models:** Predict continuous values (e.g., logD, clearance rate). - **Popular algorithms:** Random forests, support vector machines, neural networks, deep learning (e.g., graph neural networks). - **Model selection:** Depends on property, data size, and interpretability needs. - **Multi-task learning:** Some models predict multiple ADMET endpoints simultaneously.

16.7 8.4 Interpreting ADMET Predictions

Interpreting model predictions is crucial for decision-making in drug discovery. - **Performance metrics:** Accuracy, ROC-AUC, precision, recall, RMSE, etc. - **Feature importance:** Identify which descriptors most influence predictions (e.g., SHAP values, permutation importance). - **Applicability domain:** Assess whether a compound is within the model's reliable prediction space. - **Experimental validation:** Always confirm computational predictions with laboratory assays when possible.

Chapter 17

Chapter 9: Molecular Generation

17.1 Summary

Generative models can design novel molecules with desired properties. This chapter introduces molecular generation techniques, including SMILES-based and graph-based approaches, and discusses how to evaluate generated molecules.

17.2 Key Code Snippets

17.2.1 1. Generate Molecules with a SMILES RNN (Example)

```
# Example only: replace with actual model
import torch
model = ... # pretrained RNN
z = torch.randn(1, latent_dim)
smiles = model.sample(z)
print(smiles)
```

17.2.2 2. Evaluate Validity of Generated SMILES

```
from rdkit import Chem
smiles_list = ["CCO", "invalid_smiles"]
for s in smiles_list:
    mol = Chem.MolFromSmiles(s)
    print(f"{s}: {'valid' if mol else 'invalid'}")
```

17.3 Notebook Link

For a full workflow, see the interactive notebook: (notebook not available for this chapter)

17.4 9.1 Generative Models Overview

Generative models are machine learning models that can create new data samples similar to those in the training set. In drug discovery, they are used to design novel molecules with desired properties.

- **Types:** - Variational Autoencoders (VAEs) - Generative Adversarial Networks (GANs) - Recurrent Neural Networks (RNNs) - Diffusion models
- **Applications:** - De novo drug design - Scaffold hopping - Library expansion

17.5 9.2 SMILES-Based Generation

SMILES-based generative models treat molecules as sequences of characters (SMILES strings).

- **Approaches:** - RNNs, LSTMs, and Transformers trained to generate valid SMILES.
- VAEs and GANs adapted for sequence data.
- **Advantages:** - Leverage large SMILES datasets.
- Simple to implement and evaluate.
- **Limitations:** - May generate invalid or syntactically incorrect SMILES.
- Hard to enforce chemical constraints.

17.6 9.3 Graph-Based Generation

Graph-based models generate molecules as graphs, with atoms as nodes and bonds as edges.

- **Approaches:** - Graph VAEs, Graph GANs, autoregressive graph models.
- Directly model chemical structure and constraints.
- **Advantages:** - Better chemical validity and diversity.
- Can encode domain knowledge (e.g., valence rules).
- **Limitations:** - More complex architectures and training.

17.7 9.4 Evaluating Generated Molecules

Evaluating generative models requires assessing both the quality and utility of generated molecules.

- **Metrics:** - Validity: Fraction of chemically valid molecules.
- Uniqueness: Fraction of unique molecules generated.
- Novelty: Fraction not present in the training set.
- Property distribution: Compare properties (e.g., MW, logP) to real molecules.

- **Practical evaluation:** - Visual inspection, property prediction, and virtual screening of generated compounds.

Chapter 18

Chapter 10: Conditional Generation

Conditional generation enables the design of molecules with specific properties or activities. This chapter covers conditional VAEs, GANs, and reinforcement learning for molecular design.

18.1 10.1 Conditional Generative Models

Conditional generative models are deep learning architectures that generate new molecules based on input conditions, such as desired properties or biological activities.

- Extend standard VAEs by incorporating property or activity information into both the encoder and decoder.
- Enable sampling of molecules with specific characteristics by conditioning on property vectors.
- Example: Generating molecules with a target logP or activity value.
- GANs with both generator and discriminator conditioned on property information.
- Used to generate molecules that satisfy certain constraints or exhibit desired features.

Key Idea: By conditioning the generative process, these models can be steered toward producing molecules with user-specified attributes.



Example: Conditional VAE in PyTorch (Conceptual)

```
# Assume X: molecular features, y: property labels
class ConditionalVAE(nn.Module):
    def __init__(self, ...):
        # ...
    def encode(self, x, y):
        # Concatenate x and y, then encode
    def decode(self, z, y):
        # Concatenate z and y, then decode
    def forward(self, x, y):
        # ...
# Training: model(X, y)
```

```
# Generation: sample z, specify y_target, model.decode(z, y_target)
```

Further Reading: [@cvae_arxiv](#) [@cgan_arxiv](#)

18.2 10.2 Property-Conditioned Generation

Property-conditioned generation refers to the process of designing molecules that meet specific property requirements (e.g., solubility, bioactivity, toxicity).

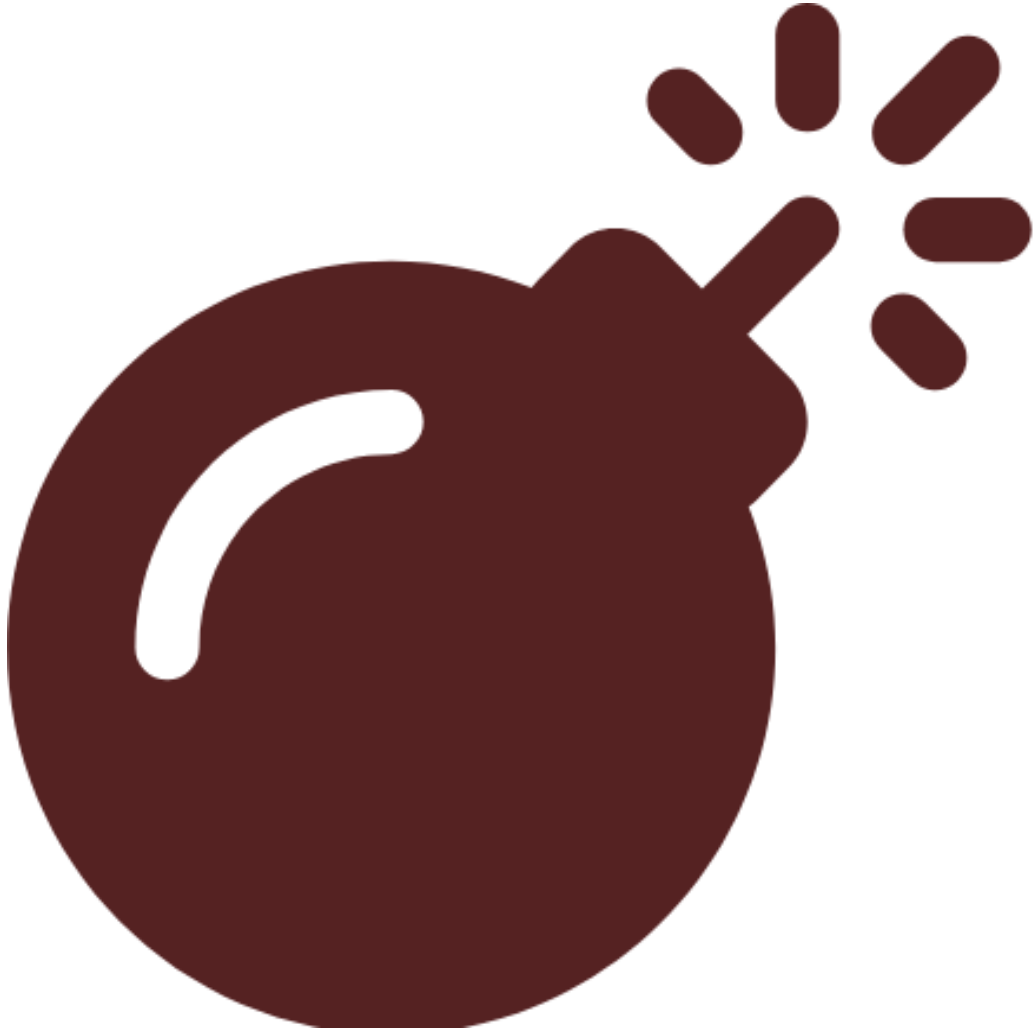
- Concatenate property vectors with molecular representations during training and generation
- Use property predictors as part of the loss function to guide the model toward desired output
- Drug design: Generate compounds with high predicted activity and low toxicity.
- Material science: Design molecules with specific physical or chemical properties.

Example (Conceptual): 1. Train a cVAE on a dataset of molecules and their properties. 2. At generation time, specify a target property value (e.g., logP = 2.5). 3. Sample from the latent space conditioned on this value to generate new molecules.

Example: Property-Conditioned SMILES Generation

```
# Pseudocode for property-conditioned generation
property_vector = torch.tensor([target_logP, target_activity])
z = torch.randn(latent_dim)
generated_smiles = model.decode(z, property_vector)
```

Workflow Diagram:



Further Reading: @molgan_arxiv @property_conditioned_review

18.3 10.3 Reinforcement Learning Approaches

Reinforcement learning (RL) can be used to optimize molecular generation toward specific objectives by treating molecule design as a sequential decision process.

- The model (agent) generates molecules step by step (e.g., adding atoms/bonds or SMILES tokens)
- Rewards are given based on how well the generated molecule meets the target property or achieves the goal
- Can include predicted activity, synthetic accessibility, drug-likeness, or other custom criteria.

- REINVENT (policy gradient for SMILES generation)
- Graph-based RL for molecular graphs

Example (Conceptual): 1. Define a reward function based on the desired property (e.g., high binding affinity). 2. Use RL to train a generative model to maximize the expected reward. 3. Sample new molecules that are likely to satisfy the design objectives.

Example: Reinforcement Learning for SMILES Generation

```
# Pseudocode for RL-based molecular generation
for episode in range(num_episodes):
    state = env.reset()
    done = False
    while not done:
        action = agent.select_action(state)
        next_state, reward, done, _ = env.step(action)
        agent.update(state, action, reward, next_state)
        state = next_state
# Reward = f(predicted_activity, drug_likeness, etc.)
```

Workflow Diagram:



Further Reading: @reinvent @deep_rl_drug_design

Chapter 19

Chapter 11: Transformers in Chemistry

Transformers have revolutionized sequence modeling and are now applied to chemistry. This chapter explores transformer architectures for SMILES, reaction prediction, and property modeling.

19.1 11.1 Introduction to Transformers

Transformers are deep learning models based on self-attention mechanisms, originally developed for natural language processing (NLP). They excel at modeling long-range dependencies in sequential data and have become the foundation for state-of-the-art models in many domains.

- **Key Features:**

- Self-attention allows the model to weigh the importance of different parts of the input sequence.
- Highly parallelizable and scalable to large datasets.
- Examples: BERT, GPT, T5, and their chemistry adaptations.

19.2 11.2 SMILES Transformers

SMILES (Simplified Molecular Input Line Entry System) strings are a common way to represent molecules as sequences. Transformers can be trained on SMILES data for various tasks:

- **SMILES Language Modeling:**

- Learn the syntax and structure of valid SMILES strings.
- Enable generative tasks such as de novo molecule generation.

- **Pretrained Models:**

- ChemBERTa, SMILES-BERT, and other transformer models pre-trained on large chemical databases.
- Can be fine-tuned for downstream tasks like property prediction or reaction classification.
- **Example (Conceptual):**
 - Tokenize SMILES strings and train a transformer to predict the next token or reconstruct masked tokens.

19.3 11.3 Reaction Prediction

Transformers are used to predict the outcomes of chemical reactions by modeling reactants and products as sequences.

- **Sequence-to-Sequence Models:**
 - Encode reactant SMILES and decode product SMILES.
 - Trained on large reaction datasets (e.g., USPTO).
- **Applications:**
 - Forward reaction prediction: Given reactants, predict products.
 - Retrosynthesis: Given a product, predict possible reactants.
- **Advantages:**
 - Capture complex reaction patterns and context.
 - Outperform traditional rule-based systems in many cases.

19.4 11.4 Property Prediction

Transformers can be fine-tuned to predict molecular properties from SMILES or graph representations.

- **Approach:**
 - Pretrain a transformer on large chemical datasets.
 - Fine-tune on labeled data for specific properties (e.g., solubility, toxicity, activity).
- **Benefits:**
 - Leverage transfer learning for improved accuracy with limited labeled data.
 - Capture subtle structure-property relationships.
- **Example (Conceptual):**
 - Use a pretrained SMILES transformer to predict logP, binding affinity, or ADMET properties.

Part IV

Part III: Structure-Based Drug Design

Chapter 20

Chapter 12: Protein Structure

Understanding protein structure is key to structure-based drug design. This chapter covers protein structure basics, experimental methods, and computational prediction (e.g., AlphaFold).

20.1 12.1 Protein Structure Levels

Proteins have a hierarchical structure, each level contributing to their function:

- **Primary Structure:**
 - Linear sequence of amino acids in a polypeptide chain.
 - Determines all higher levels of structure.
- **Secondary Structure:**
 - Local folding patterns stabilized by hydrogen bonds.
 - Common motifs: α -helices and β -sheets.
- **Tertiary Structure:**
 - 3D arrangement of a single polypeptide chain, including side-chain interactions.
 - Stabilized by hydrophobic interactions, disulfide bonds, and ionic interactions.
- **Quaternary Structure:**
 - Assembly of multiple polypeptide chains (subunits) into a functional protein complex.

20.2 12.2 Experimental Determination

Experimental methods for determining protein structures include:

- **X-ray Crystallography:**
 - Most common method for high-resolution structures.
 - Requires crystallization of the protein.
- **Nuclear Magnetic Resonance (NMR) Spectroscopy:**
 - Used for small to medium-sized proteins in solution.
 - Provides information on protein dynamics.
- **Cryo-Electron Microscopy (Cryo-EM):**
 - Suitable for large complexes and membrane proteins.
 - Does not require crystallization; recent advances have improved resolution.

20.3 12.3 Structure Prediction Tools

Computational tools can predict protein structures when experimental data is unavailable:

- **Homology Modeling:**
 - Builds models based on similarity to known structures (templates).
 - Tools: SWISS-MODEL, MODELLER.
- **Ab Initio Prediction:**
 - Predicts structure from sequence alone, without templates.
 - Computationally intensive; used for small proteins.
- **AlphaFold:**
 - Deep learning-based tool from DeepMind.
 - Achieves near-experimental accuracy for many proteins.
 - [AlphaFold Protein Structure Database](#)

20.4 12.4 Protein-Ligand Interactions

Understanding how proteins interact with small molecules (ligands) is crucial for drug design:

- **Binding Sites:**
 - Specific regions on the protein where ligands bind.
 - Often located in pockets or grooves on the protein surface.
- **Interaction Types:**
 - Hydrogen bonds, hydrophobic interactions, ionic bonds, van der Waals forces.
- **Docking and Scoring:**
 - Computational docking predicts how ligands fit into binding sites.
 - Scoring functions estimate binding affinity.
- **Applications:**
 - Lead discovery, optimization, and understanding mechanism of action.

Chapter 21

Chapter 13: Molecular Docking

Molecular docking predicts how small molecules bind to protein targets. This chapter introduces docking theory, software, and practical workflows for virtual screening.

21.1 13.1 Docking Theory

Docking is a computational technique that predicts the preferred orientation of a small molecule (ligand) when bound to a protein (receptor). The goal is to estimate binding affinity and identify potential drug candidates.

- **Key Concepts:**

- **Search Algorithm:** Explores possible ligand conformations and orientations in the binding site.
- **Scoring Function:** Estimates the strength of the interaction (binding affinity) between ligand and protein.
- **Rigid vs. Flexible Docking:** Rigid docking keeps the protein and ligand fixed; flexible docking allows movement in the ligand and/or protein.

21.2 13.2 Docking Software (AutoDock, Vina)

Several software packages are widely used for molecular docking:

- **AutoDock:**

- One of the most popular open-source docking tools.
- Uses Lamarckian genetic algorithm for conformational search.
- Supports flexible ligand and some flexible side-chain docking.

- **AutoDock Vina:**
 - Improved version of AutoDock with faster performance and better scoring.
 - User-friendly and widely adopted for virtual screening.
- **Other Tools:**
 - DOCK, GOLD, Glide, and others are also used in academia and industry.

21.3 13.3 Preparing Structures for Docking

Proper preparation of protein and ligand structures is critical for successful docking:

- **Protein Preparation:**
 - Remove water molecules and irrelevant ligands.
 - Add missing atoms and assign correct protonation states.
 - Define the binding site (e.g., using known ligand or cavity detection).
- **Ligand Preparation:**
 - Generate 3D conformations.
 - Assign correct protonation and tautomeric states.
 - Minimize energy to relieve strain.
- **File Formats:**
 - PDBQT (AutoDock/Vina), MOL2, SDF, PDB.

21.4 13.4 Analyzing Docking Results

After docking, results must be analyzed to identify promising candidates:

- **Scoring and Ranking:**
 - Docking software provides binding affinity scores for each pose.
 - Rank compounds based on scores, but consider limitations of scoring functions.
- **Visual Inspection:**
 - Use molecular visualization tools (e.g., PyMOL, Chimera) to examine binding modes.
 - Check for key interactions (hydrogen bonds, hydrophobic contacts) and fit in the binding site.
- **Post-Processing:**
 - Cluster similar poses, filter by interaction criteria, and consider rescoring with more accurate methods if needed.
- **Experimental Validation:**
 - Top candidates should be validated experimentally for biological activity.

Chapter 22

Chapter 14: ML-Enhanced Docking

Machine learning can improve docking accuracy and speed. This chapter discusses ML scoring functions, pose prediction, and integration with docking pipelines.

22.1 14.1 ML Scoring Functions

Traditional docking scoring functions are based on physics or empirical rules. Machine learning (ML) scoring functions use data-driven models to predict binding affinity more accurately.

- **Types of ML Scoring Functions:**
 - **Regression Models:** Predict binding affinity from features (e.g., random forests, gradient boosting, neural networks).
 - **Deep Learning Models:** Use 3D protein-ligand structures as input (e.g., convolutional neural networks, graph neural networks).
- **Popular ML Scoring Tools:**
 - **RF-Score:** Random forest model using atom pair features.
 - **DeltaVina:** Combines ML with traditional Vina scoring.
 - **DeepDock, GNINA:** Deep learning-based scoring using 3D grids or graphs.
- **Advantages:**
 - Capture complex, non-linear interactions.
 - Can be retrained on new data for specific targets.

22.2 14.2 Pose Prediction

ML models can also predict the correct binding pose of a ligand in a protein binding site.

- **Pose Ranking:**
 - ML models can re-rank docking poses generated by traditional software, improving the selection of the most likely binding mode.
- **End-to-End Pose Prediction:**
 - Some deep learning models directly predict ligand poses from protein and ligand structures (e.g., EquiBind, DeepDock).
- **Evaluation:**
 - Root-mean-square deviation (RMSD) is used to assess pose accuracy compared to experimental structures.

22.3 14.3 Integrating ML with Docking

ML can be integrated into docking pipelines to enhance virtual screening and lead optimization:

- **Hybrid Workflows:**
 - Use traditional docking to generate poses, then apply ML scoring for more accurate ranking.
 - Filter large libraries with fast ML models before detailed docking.
- **Active Learning:**
 - Iteratively improve ML models by incorporating feedback from new docking or experimental results.
- **Automation:**
 - ML models can automate hit selection, pose filtering, and prioritization in large-scale screens.
- **Example Workflow:**
 1. Dock a library of compounds using AutoDock Vina.
 2. Score top poses with a deep learning model (e.g., GNINA).
 3. Select candidates for experimental validation based on ML scores.

Part V

Part IV: Synthesis and Advanced Topics

Chapter 23

Chapter 15: Retrosynthesis

Retrosynthesis planning is essential for proposing synthetic routes to molecules. This chapter covers rule-based and AI-driven retrosynthesis tools and strategies.

23.1 15.1 Introduction to Retrosynthesis

Retrosynthesis is the process of deconstructing a target molecule into simpler precursor structures, working backward from the product to available starting materials. It is a key strategy in organic synthesis and drug development.

- **Key Concepts:**
 - Identify strategic bonds to break (disconnections).
 - Propose sequences of reactions to build the target molecule from simpler compounds.
 - Use chemical knowledge, reaction databases, and computational tools.

23.2 15.2 Rule-Based Approaches

Rule-based retrosynthesis uses encoded chemical rules and reaction templates to suggest possible disconnections and synthetic routes.

- **How It Works:**
 - Reaction rules are derived from known chemical transformations.
 - The system applies these rules recursively to break down the target molecule.
- **Popular Tools:**
 - **LHASA:** One of the earliest expert systems for retrosynthesis.
 - **Chematica (Synthia):** Commercial platform with a large rule database.

- **RDKit:** Open-source cheminformatics toolkit with basic retrosynthesis capabilities.
- **Advantages:**
 - Transparent and interpretable routes.
 - Leverages established chemical knowledge.
- **Limitations:**
 - Limited by the coverage of encoded rules.
 - May miss novel or unconventional pathways.

23.3 15.3 AI-Driven Retrosynthesis

AI-driven retrosynthesis uses machine learning and deep learning to predict synthetic routes, often learning directly from reaction databases.

- **Template-Based Models:**
 - Use reaction templates extracted from data, similar to rule-based systems but with automated extraction and ranking.
- **Template-Free Models:**
 - Sequence-to-sequence (seq2seq) models treat retrosynthesis as a translation problem (product to reactants).
 - Graph neural networks (GNNs) model molecular graphs for reaction prediction.
- **Popular AI Tools:**
 - **ASKCOS:** Open-source platform using neural networks for retrosynthesis.
 - **IBM RXN:** Cloud-based AI retrosynthesis tool.
 - **AiZynthFinder:** Open-source tool for automated retrosynthetic planning.
- **Advantages:**
 - Can generalize to novel reactions and pathways.
 - Continuously improve with more data.
- **Limitations:**
 - May generate less interpretable routes.
 - Dependent on quality and diversity of training data.

23.4 15.4 Evaluating Synthetic Routes

Evaluating proposed synthetic routes is crucial for practical application:

- **Criteria:**
 - Number of steps (shorter is often better).
 - Availability and cost of starting materials.
 - Feasibility and reliability of each reaction step.
 - Overall yield and scalability.
- **Scoring and Ranking:**

- Many tools provide scores based on route length, confidence, or synthetic accessibility.
- **Experimental Validation:**
 - Ultimately, routes must be tested in the lab to confirm feasibility.

Chapter 24

Chapter 16: Multi-Modal Learning

Multi-modal learning combines data types (e.g., chemical, biological, clinical) for improved predictions. This chapter explores architectures and applications in drug discovery.

24.1 16.1 What is Multi-Modal Learning?

Multi-modal learning is a machine learning paradigm that integrates information from multiple data sources or modalities. In drug discovery, these modalities can include chemical structures, gene expression profiles, protein sequences, clinical data, and more.

- **Motivation:**
 - Single data types may not capture the full complexity of biological systems.
 - Combining modalities can improve prediction accuracy and provide deeper insights.
- **Examples of Modalities:**
 - Chemical: SMILES, molecular graphs, descriptors
 - Biological: gene expression, protein sequences, omics data
 - Clinical: patient records, trial outcomes

24.2 16.2 Data Integration Strategies

Integrating diverse data types requires careful design of model architectures and data processing pipelines.

- **Early Fusion:**

- Concatenate features from all modalities at the input layer.
- Simple but may not capture complex relationships.
- **Late Fusion:**
 - Train separate models for each modality, then combine predictions (e.g., averaging, stacking).
 - Useful when modalities are very different in scale or type.
- **Hybrid/Intermediate Fusion:**
 - Combine intermediate representations from each modality within the model (e.g., via attention mechanisms or shared layers).
 - Allows learning of cross-modal interactions.
- **Deep Learning Architectures:**
 - Multi-branch neural networks, attention-based models, and graph neural networks can all be adapted for multi-modal integration.

24.3 16.3 Applications in Drug Discovery

Multi-modal learning is increasingly used in drug discovery to leverage the wealth of available data:

- **Drug Response Prediction:**
 - Combine chemical structure and gene expression data to predict how cell lines or patients will respond to drugs.
- **Target Identification:**
 - Integrate omics data, protein-protein interactions, and chemical information to identify new drug targets.
- **Adverse Event Prediction:**
 - Use clinical records, molecular properties, and biological data to predict side effects or toxicity.
- **Precision Medicine:**
 - Tailor drug recommendations by integrating patient-specific clinical and molecular data.
- **Example (Conceptual):**
 - A multi-modal neural network takes SMILES strings, gene expression profiles, and patient data as input to predict drug efficacy for personalized treatment.

Chapter 25

Chapter 17: End-to-End Pipeline

This chapter presents a complete AI-driven drug discovery pipeline, from data collection to candidate prioritization, integrating methods from previous chapters.

25.1 17.1 Pipeline Overview

An end-to-end AI-driven drug discovery pipeline integrates cheminformatics, machine learning, and structure-based methods to accelerate the identification of promising drug candidates.

- **Key Stages:**
 1. Data collection and curation
 2. Feature extraction and data preprocessing
 3. Model training and validation
 4. Virtual screening and candidate prioritization

25.2 17.2 Data Preparation

High-quality data is the foundation of any successful pipeline.

- **Data Sources:**
 - Public databases (e.g., ChEMBL, PubChem, PDB)
 - In-house experimental data
- **Data Curation:**
 - Remove duplicates, standardize chemical structures, and handle missing values.
 - Annotate data with relevant properties (e.g., activity, ADMET, target information).

- **Feature Engineering:**
 - Generate molecular fingerprints, descriptors, and graph representations.
 - Integrate multi-modal data (e.g., chemical, biological, clinical).

25.3 17.3 Model Training and Evaluation

Machine learning models are trained to predict properties such as bioactivity, ADMET, or binding affinity.

- **Model Selection:**
 - Choose appropriate algorithms (e.g., random forests, deep neural networks, graph neural networks, transformers).
- **Training:**
 - Split data into training, validation, and test sets.
 - Optimize hyperparameters and prevent overfitting.
- **Evaluation:**
 - Use metrics such as ROC-AUC, RMSE, accuracy, and F1-score.
 - Perform cross-validation and external validation if possible.
- **Interpretability:**
 - Analyze feature importance and model explanations to gain insights.

25.4 17.4 Candidate Selection

The final stage is to prioritize candidates for experimental validation.

- **Virtual Screening:**
 - Screen large compound libraries using trained models and/or docking.
 - Rank compounds based on predicted activity, ADMET, and synthetic accessibility.
- **Filtering:**
 - Apply property-based and rule-based filters (e.g., Lipinski's rules, toxicity filters).
- **Multi-Parameter Optimization:**
 - Balance potency, selectivity, safety, and developability.
- **Experimental Validation:**
 - Select top candidates for synthesis and biological testing.
- **Iterative Improvement:**
 - Incorporate new data and feedback to refine models and repeat the pipeline.

Chapter 26

Chapter 18: Case Studies

Real-world case studies demonstrate the application of AI and cheminformatics in drug discovery. This chapter presents selected examples and lessons learned.

26.1 18.1 Case Study 1: ALS Drug Discovery

Background: Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease with limited treatment options. AI-driven approaches have been used to identify new therapeutic candidates.

- **Data Collection:**
 - Integrated omics data, literature mining, and known ALS-related targets.
 - Used cheminformatics to curate compound libraries for screening.
- **Virtual Screening:**
 - Applied machine learning models to predict neuroprotective activity.
 - Used molecular docking to prioritize compounds for SOD1 and TDP-43 protein targets.
- **Experimental Validation:**
 - Top candidates were tested in cell-based assays for neuroprotection.
- **Outcome:**
 - Identified several promising compounds for further development.

26.2 18.2 Case Study 2: Kinase Inhibitors

Background: Kinases are important drug targets in cancer and other diseases. AI and cheminformatics have accelerated the discovery of selective kinase inhibitors.

- **Data Integration:**

- Combined chemical structure data, kinase activity profiles, and protein-ligand interaction data.
- **QSAR Modeling:**
 - Built machine learning models to predict kinase inhibition based on molecular fingerprints and descriptors.
- **Structure-Based Design:**
 - Used docking and molecular dynamics to optimize binding affinity and selectivity.
- **Lead Optimization:**
 - Applied multi-parameter optimization to balance potency, selectivity, and ADMET properties.
- **Outcome:**
 - Accelerated identification of novel kinase inhibitors with improved profiles.

26.3 18.3 Lessons Learned

- **Data Quality Matters:**
 - High-quality, well-curated data is essential for successful AI-driven drug discovery.
- **Integration is Key:**
 - Combining multiple data types and methods yields better results than using a single approach.
- **Iterative Process:**
 - Drug discovery is iterative—models and hypotheses should be refined as new data becomes available.
- **Collaboration:**
 - Success often requires collaboration between computational scientists, chemists, and biologists.
- **Limitations:**
 - AI models are only as good as their training data and may not generalize to novel targets or chemotypes.

Chapter 27

Chapter 19: External Tools

This chapter reviews external tools and databases commonly used in computational drug discovery, including RCSB PDB, ChEMBL, and PubChem.

27.1 19.1 Structural Databases

Structural databases provide 3D structures of proteins, nucleic acids, and complexes, which are essential for structure-based drug design.

- **RCSB Protein Data Bank (PDB):**
 - The primary repository for experimentally determined 3D structures of biological macromolecules.
 - rcsb.org
 - Structures are available in PDB and mmCIF formats.
- **AlphaFold Protein Structure Database:**
 - Provides predicted protein structures using deep learning.
 - alphafold.ebi.ac.uk
- **Other Databases:**
 - PDB-REDO (refined structures), CATH/SCOP (structure classification).

27.2 19.2 Bioactivity Databases

Bioactivity databases contain information on small molecules, their biological activities, and targets.

- **ChEMBL:**
 - A large, open-access database of bioactive molecules with drug-like properties.
 - Includes activity data (IC₅₀, Ki, etc.), targets, and assay information.

- ebi.ac.uk/chembl
- **PubChem:**
 - Comprehensive resource for chemical structures, properties, and biological activities.
 - Includes compound, substance, and bioassay databases.
 - pubchem.ncbi.nlm.nih.gov
- **BindingDB:**
 - Focuses on binding affinities for protein-ligand complexes.
 - bindingdb.org

27.3 19.3 Workflow Automation Tools

Workflow automation tools streamline computational drug discovery by integrating multiple steps and tools into reproducible pipelines.

- **KNIME:**
 - Open-source platform for data analytics and workflow automation.
 - Widely used for cheminformatics, bioinformatics, and machine learning workflows.
 - knime.com
- **Pipeline Pilot:**
 - Commercial workflow automation tool for scientific data processing.
 - Supports cheminformatics, bioinformatics, and analytics.
- **Galaxy:**
 - Web-based platform for accessible, reproducible, and transparent computational research.
 - usegalaxy.org
- **Snakemake/Nextflow:**
 - Workflow management systems for scalable and reproducible data analysis pipelines.
- **Integration with Scripting:**
 - Python, R, and shell scripts are often used to glue together custom workflows.

Chapter 28

Chapter 20: Ethical Considerations

Ethics are critical in AI-driven drug discovery. This chapter discusses data privacy, algorithmic bias, and responsible innovation in cheminformatics and AI.

28.1 20.1 Data Privacy and Security

Protecting sensitive data is essential in drug discovery, especially when working with patient information, proprietary compounds, or clinical trial data.

- **Data Privacy:**
 - Ensure compliance with regulations (e.g., GDPR, HIPAA) when handling personal or health data.
 - Use anonymization and de-identification techniques to protect individual identities.
- **Data Security:**
 - Implement robust cybersecurity measures to prevent unauthorized access or data breaches.
 - Use secure data storage, encrypted communication, and access controls.
- **Data Sharing:**
 - Balance open science with the need to protect sensitive or proprietary information.

28.2 20.2 Bias in AI Models

AI models can inherit or amplify biases present in training data, leading to unfair or inaccurate predictions.

- **Sources of Bias:**
 - Imbalanced datasets (e.g., overrepresentation of certain chemotypes or populations).
 - Historical biases in experimental data or reporting.
- **Consequences:**
 - Reduced model performance for underrepresented groups or rare targets.
 - Potential for inequitable healthcare outcomes.
- **Mitigation Strategies:**
 - Use diverse and representative datasets.
 - Apply fairness-aware algorithms and regular audits for bias.
 - Transparently report model limitations and performance across sub-groups.

28.3 20.3 Responsible Innovation

Responsible innovation ensures that AI and cheminformatics tools are developed and used in ways that benefit society and minimize harm.

- **Transparency:**
 - Make model architectures, training data, and evaluation metrics publicly available when possible.
- **Accountability:**
 - Clearly define responsibility for model predictions and decisions, especially in clinical settings.
- **Societal Impact:**
 - Consider the broader implications of AI-driven drug discovery, such as access to medicines, dual-use risks, and environmental impact.
- **Continuous Oversight:**
 - Engage stakeholders (scientists, ethicists, patients, regulators) throughout the development and deployment process.

Chapter 29

Chapter 21: Future Next Steps

The final chapter outlines emerging trends and future directions in AI-driven drug discovery, including new algorithms, data sources, and collaborative science.

29.1 21.1 Emerging Technologies

AI-driven drug discovery is rapidly evolving, with several emerging technologies poised to transform the field:

- **Foundation Models and Large Language Models:**
 - Use of large pretrained models (e.g., ChemBERTa, MolBERT, GPT-4) for molecular design, property prediction, and literature mining.
- **Generative AI for Molecule Design:**
 - Advanced generative models (e.g., diffusion models, graph-based VAEs) for de novo compound generation.
- **Multi-Omics Integration:**
 - Combining genomics, proteomics, metabolomics, and clinical data for holistic drug discovery.
- **Quantum Computing:**
 - Potential to revolutionize molecular simulation and optimization tasks.
- **Automated Laboratories (Lab Automation):**
 - Integration of robotics, AI, and cloud labs for high-throughput synthesis and testing.

29.2 21.2 Open Science and Collaboration

Open science and collaborative efforts are accelerating progress in drug discovery:

- **Open Data Initiatives:**
 - Public databases (e.g., ChEMBL, PubChem, PDB) and open-access datasets fuel innovation.
- **Open-Source Software:**
 - Community-driven tools (e.g., RDKit, DeepChem, OpenMM) enable reproducible research.
- **Collaborative Platforms:**
 - Crowdsourcing challenges (e.g., DREAM, Kaggle) and collaborative consortia (e.g., Open Targets, MELLODDY) bring together diverse expertise.
- **Preprints and Open Publishing:**
 - Rapid dissemination of new findings and methods.

29.3 21.3 Next Steps for Practitioners

For those looking to advance in AI-driven drug discovery, consider the following steps:

- **Continuous Learning:**
 - Stay updated with the latest research, tools, and best practices.
 - Participate in workshops, online courses, and conferences.
- **Hands-On Practice:**
 - Apply methods from this book to real-world datasets and problems.
 - Contribute to open-source projects and collaborative research.
- **Ethical and Responsible Innovation:**
 - Prioritize transparency, fairness, and societal impact in your work.
- **Networking and Collaboration:**
 - Build connections with researchers, clinicians, and industry professionals.
- **Embrace Interdisciplinarity:**
 - Combine expertise in chemistry, biology, data science, and AI for maximum impact.

Chapter 30

Conclusion

Chapter 31

Conclusion

You have completed “AI-Driven Drug Discovery: A Hands-On Guide.” We hope you now feel equipped to apply AI and computational tools to real-world drug discovery challenges. Continue exploring, building, and contributing to this exciting field!

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