# Appendix B: Dataset Catalogue

This appendix provides a comprehensive list of datasets relevant for neuroAl research, along with access information and example code for loading and working with these datasets.

# **B.1 Neuroscience Datasets**

# **Electrophysiology Data**

### Allen Brain Observatory

- Description: Single-cell recordings from visual cortex of mice during visual stimulation
- Contents: Neural activity from ~75,000 neurons across multiple cortical areas and layers
- Access: <a href="https://observatory.brain-map.org/">https://observatory.brain-map.org/</a>
- Documentation: https://allensdk.readthedocs.io/en/latest/brain\_observatory.html

```
# Example: Loading Allen Brain Observatory data
from allensdk.core.brain observatory_cache import BrainObservatoryCache
import matplotlib.pyplot as plt
# Initialize the cache
boc = BrainObservatoryCache(manifest_file='boc_manifest.json')
# Get experiment containers for specific targeted structures and imaging depths
experiments = boc.get_experiment_containers(targeted_structures=['VISp'],
                                       imaging depths=[175])
# Access the first experiment
container_id = experiments[0]['id']
# Get experiment with natural scenes stimulus
experiment id = boc.get ophys experiments(
    experiment container ids=[container id],
    stimuli=['natural scenes'])[0]['id']
# Get neural data
data_set = boc.get_ophys_experiment_data(experiment_id)
# Get fluorescence traces for all cells
timestamps, traces = data_set.get_fluorescence_traces()
print(f"Recorded {traces.shape[0]} neurons for {len(timestamps)} timepoints")
# Plot example trace from first neuron
plt.figure(figsize=(12, 4))
plt.plot(timestamps, traces[0])
plt.xlabel('Time (s)')
plt.ylabel('Fluorescence (a.u.)')
plt.title('Example Neuron Activity')
plt.show()
```

#### Collaborative Research in Computational Neuroscience (CRCNS)

- **Description**: Diverse electrophysiology datasets from various labs and species
- Contents: Single-unit and multi-unit recordings, EEG, MEG, and more
- Access: https://crcns.org/
- **Documentation**: Varies by dataset

```
# Example: Loading CRCNS data (HC1 - hippocampal recordings from rats)
import h5py
import numpy as np
import matplotlib.pyplot as plt
# Open the HDF5 file (you would need to download this)
with h5py.File('example_data.h5', 'r') as f:
    # List all groups
    print("Keys: %s" % list(f.keys()))
    # Get spike times for a specific cell
    spikes = np.array(f['spikes']['times'])
    # Plot raster
    plt.figure(figsize=(12, 4))
    plt.plot(spikes, np.ones_like(spikes), '|', markersize=10)
    plt.xlabel('Time (s)')
    plt.ylabel('Neuron')
    plt.title('Spike Raster')
    plt.grid(True)
    plt.show()
```

#### Neurodata Without Borders (NWB)

- **Description**: Standardized format for neurophysiology data
- Contents: Various datasets following the NWB format standard
- Access: https://www.nwb.org/example-datasets/
- **Documentation**: https://pynwb.readthedocs.io/

```
# Example: Reading NWB formatted data
from pvnwb import NWBHDF5I0
import matplotlib.pyplot as plt
# Open NWB file (you would need to download an example file)
with NWBHDF5IO('example.nwb', 'r') as io:
    nwbfile = io.read()
    # Print available acquisition data
    print(nwbfile.acquisition)
    # Access LFP data if available
    if 'LFP' in nwbfile.acquisition:
        lfp_data = nwbfile.acquisition['LFP'].data[:]
        lfp_timestamps = nwbfile.acquisition['LFP'].timestamps[:]
        # Plot LFP
        plt.figure(figsize=(12, 4))
        plt.plot(lfp_timestamps, lfp_data)
        plt.xlabel('Time (s)')
        plt.ylabel('Voltage (mV)')
        plt.title('LFP Recording')
        plt.show()
```

# **Neuroimaging Data**

#### Human Connectome Project (HCP)

- **Description**: High-quality neuroimaging data from 1,200+ healthy young adults
- Contents: MRI (structural, functional, diffusion), MEG, behavioral, and genetic data
- Access: <a href="https://www.humanconnectome.org/study/hcp-young-adult">https://www.humanconnectome.org/study/hcp-young-adult</a>
- **Documentation**: <a href="https://www.humanconnectome.org/study/hcp-young-adult/documentation">https://www.humanconnectome.org/study/hcp-young-adult/documentation</a>

```
# Example: Working with HCP data using nilearn
from nilearn import datasets, plotting
import matplotlib.pyplot as plt
# Download a preprocessed resting-state fMRI from HCP
# Note: You need to register and get credentials for actual HCP data
# This example uses a sample from nilearn, not the actual HCP dataset
hcp_dataset = datasets.fetch_development_fmri(n_subjects=1)
# Get the first subject's data
func_filename = hcp_dataset.func[0]
# Plot the mean image
mean_img = plotting.plot_epi(func_filename, title='Mean fMRI image')
plt.show()
# Plot the temporal variance
var_img = plotting.plot_stat_map(plotting.mean_img(func_filename),
                              title='Temporal variance')
plt.show()
```

#### **OpenNeuro**

- Description: Open platform for sharing BIDS-compatible neuroimaging data
- Contents: fMRI, EEG, MEG, iEEG datasets from various studies
- Access: <a href="https://openneuro.org/">https://openneuro.org/</a>
- **Documentation**: Varies by dataset

```
# Example: Using openneuro-py to download data
# !pip install openneuro-pv
import openneuro
import os
# Create a download instance
api = openneuro.OpenNeuroAPI()
# Specify dataset ID (ds002422 is an example, you can find many on the website)
dataset id = 'ds002422'
# Set download directory
download_dir = './openneuro_data'
os.makedirs(download dir, exist ok=True)
# Download a specific dataset
# This downloads the dataset metadata
downloader = api.download(dataset id, download dir)
# List all files (doesn't download them vet)
files = [f for f in downloader.files if not f.endswith('/')]
print(f"Total files: {len(files)}")
# Download a specific file
# downloader.download([files[0]]) # Uncomment to actually download
# For working with the data, you would typically use BIDS tools
# !pip install pybids
from bids import BIDSLayout
# Create a layout to work with the BIDS dataset (only metadata)
layout = BIDSLayout(download dir)
# Get all available sessions
sessions = layout.get sessions()
print(f"Available sessions: {sessions}")
# Get functional MRI runs
runs = layout.get(datatype='func', extension='.nii.gz')
print(f"Number of fMRI runs: {len(runs)}")
```

#### NeuroVault

- **Description**: Repository for statistical maps of the human brain
- Contents: fMRI and PET statistical maps
- Access: https://neurovault.org/
- Documentation: NeuroVault/NeuroVault

```
# Example: Using neurovault's API
# !pip install requests
import requests
import matplotlib.pyplot as plt
from nilearn import plotting
import nibabel as nib
import numpy as np
import os
# Get collection info (collection 1952 is an example)
base_url = 'https://neurovault.org/api/collections/1952/'
response = requests.get(base url)
collection_data = response.json()
# Get images in the collection
images url = base url + 'images/'
response = requests.get(images url)
images data = response.json()['results']
print(f"Found {len(images_data)} images")
# Download and display the first image
if len(images data) > 0:
    # Get download URL for the first image
    img url = images data[0]['file']
    img name = os.path.basename(img url)
    # Download the image
    img response = requests.get(img url)
    with open(img_name, 'wb') as f:
        f.write(img response.content)
    # Load and display the image
    img = nib.load(img name)
    plotting.plot stat map(img, title=images data[0]['name'])
    plt.show()
```

# Behavior and Cognition

#### International Brain Laboratory (IBL)

- **Description**: Standardized mouse behavioral and neural data
- Contents: Neural recordings during decision-making tasks
- Access: <a href="https://www.internationalbrainlab.com/public-resources">https://www.internationalbrainlab.com/public-resources</a>
- Documentation: <a href="https://docs.internationalbrainlab.org/">https://docs.internationalbrainlab.org/</a>

```
# Example: Loading IBL data
# !pip install ONE-api
from one.api import ONE
import numpy as no
import matplotlib.pyplot as plt
# Initialize ONE
one = ONE()
# Search for sessions with available data types
selection = one.search(dataset_types=['spikes.times', 'trials'])
print(f"Found {len(selection)} sessions")
# Select the first session
eid = selection[0]
# List all available data for this session
datasets = one.list datasets(eid)
print(f"Available datasets: {datasets}")
# Load spike data
spikes times, spikes clusters = one.load dataset(eid, 'spikes.times'), one.load d
# Load trial data
trials = one.load object(eid, 'trials')
# Plot raster for a few neurons
unique clusters = np.unique(spikes clusters)
print(f"Total neurons: {len(unique_clusters)}")
# Plot raster for 5 clusters
plt.figure(figsize=(15, 10))
for i, cluster id in enumerate(unique clusters[:5]):
    cluster spikes = spikes times[spikes clusters == cluster id]
    plt.plot(cluster spikes, np.ones like(cluster spikes) * i, '|', markersize=1)
plt.xlabel('Time (s)')
plt.vlabel('Neuron ID')
plt.title('Spike Raster')
plt.show()
```

#### Brain Imaging Data Structure (BIDS) Examples

- **Description**: Standardized datasets following BIDS format
- Contents: Various types of brain imaging and behavioral data
- Access: bids-standard/bids-examples
- **Documentation**: <a href="https://bids.neuroimaging.io/">https://bids.neuroimaging.io/</a>

```
# Example: Working with BIDS datasets
# !pip install pvbids
from bids import BIDSLayout
import os
# Path to the BIDS dataset
bids path = './bids dataset' # You would need to download or have a BIDS dataset
# Check if path exists
if os.path.exists(bids path):
    # Initialize layout
    layout = BIDSLayout(bids path)
    # Get participant information
    subjects = layout.get subjects()
    print(f"Subjects: {subjects}")
    # Get available modalities
    modalities = layout.get_modalities()
    print(f"Modalities: {modalities}")
    # Get all T1w images
    t1w_images = layout.get(suffix='T1w', extension='.nii.gz')
    print(f"Found {len(t1w_images)} T1w images")
    # Get task fMRI data
    func_runs = layout.get(datatype='func', suffix='bold')
    print(f"Found {len(func runs)} functional runs")
    # Access metadata for a specific file
    if func runs:
        metadata = layout.get_metadata(func_runs[0].path)
        print("\nSample metadata:")
        for key, value in list(metadata.items())[:5]:
            print(f"{key}: {value}")
else:
    print(f"BIDS dataset not found at {bids_path}")
```

# **B.2 Al Benchmark Datasets**

# **Computer Vision**

#### **ImageNet**

• **Description**: Large-scale image classification benchmark

- Contents: ~14 million images across 20,000+ categories
- Access: https://www.image-net.org/
- **Documentation**: https://www.image-net.org/challenges/LSVRC/

```
# Example: Using PyTorch's ImageNet loader
import torch
from torchvision import datasets, transforms
import matplotlib.pyplot as plt
import numpy as np
# Define transformations
transform = transforms.Compose([
    transforms.Resize(256),
    transforms.CenterCrop(224),
    transforms.ToTensor(),
    transforms.Normalize(mean=[0.485, 0.456, 0.406],
                         std=[0.229, 0.224, 0.225])
])
# Load ImageNet validation set (you need to have this downloaded)
# Replace 'path/to/imagenet/val' with your actual path
try:
    val_dataset = datasets.ImageNet('path/to/imagenet/val', split='val', transfor
    val loader = torch.utils.data.DataLoader(val dataset, batch size=4, shuffle=T
    # Get a batch of images
    images, labels = next(iter(val_loader))
    # Display images
    def imshow(img):
        # Unnormalize
        inv normalize = transforms.Normalize(
            mean=[-0.485/0.229, -0.456/0.224, -0.406/0.225],
            std=[1/0.229, 1/0.224, 1/0.225]
        img = inv_normalize(img)
        npimg = img.numpy()
        plt.imshow(np.transpose(npimg, (1, 2, 0)))
    plt.figure(figsize=(12, 6))
    imshow(torchvision.utils.make_grid(images))
    plt.title('ImageNet Samples')
    plt.show()
except Exception as e:
    print(f"Error loading ImageNet: {e}")
    print("Note: You need to download the ImageNet dataset separately due to its
```

# CIFAR-10/100

- **Description**: Small-scale image classification datasets
- **Contents**: 60,000 32x32 color images in 10 or 100 classes
- Access: <a href="https://www.cs.toronto.edu/~kriz/cifar.html">https://www.cs.toronto.edu/~kriz/cifar.html</a>
- Documentation: Built into most deep learning frameworks

```
# Example: Using CIFAR-10 with PyTorch
import torch
import torchvision
import torchvision.transforms as transforms
import matplotlib.pyplot as plt
import numpy as np
# Define transformations
transform = transforms.Compose([
    transforms.ToTensor(),
    transforms.Normalize((0.5, 0.5, 0.5), (0.5, 0.5, 0.5))
1)
# Load CIFAR-10 (downloaded automatically if not present)
train_dataset = torchvision.datasets.CIFAR10(root='./data', train=True,
                                             download=True, transform=transform)
train loader = torch.utils.data.DataLoader(train dataset, batch size=4,
                                           shuffle=True, num workers=2)
test dataset = torchvision.datasets.CIFAR10(root='./data', train=False,
                                            download=True, transform=transform)
test loader = torch.utils.data.DataLoader(test dataset, batch size=4,
                                          shuffle=False, num workers=2)
# Class labels
classes = ('plane', 'car', 'bird', 'cat', 'deer',
           'dog', 'frog', 'horse', 'ship', 'truck')
# Function to display images
def imshow(ima):
    img = img / 2 + 0.5 \# Unnormalize
    npimg = img.numpy()
    plt.imshow(np.transpose(npimg, (1, 2, 0)))
# Get some random training images
dataiter = iter(train loader)
images, labels = next(dataiter)
# Show images
plt.figure(figsize=(10, 5))
imshow(torchvision.utils.make grid(images))
plt.title(' '.join('%5s' % classes[labels[j]] for j in range(4)))
plt.show()
```

#### MS COCO

- Description: Dataset for object detection, segmentation, and captioning
- Contents: 330,000+ images with 80 object categories

- Access: <a href="https://cocodataset.org/">https://cocodataset.org/</a>
- Documentation: 🖸 cocodataset/cocoapi

```
# Example: Working with COCO dataset
# !pip install pycocotools
from pycocotools.coco import COCO
import numpy as no
import skimage.io as io
import matplotlib.pyplot as plt
import requests
from io import BytesIO
# Initialize COCO API for instance annotations
dataType = 'val2017'
annFile = f'./annotations/instances_{dataType}.json' # You need to download this
try:
    coco = COCO(annFile)
    # Get categories and supercategories
    cats = coco.loadCats(coco.getCatIds())
    cat_names = [cat['name'] for cat in cats]
    print(f"COCO categories: {cat names}")
    # Get all images containing specific categories
    catIds = coco.getCatIds(catNms=['person', 'dog', 'skateboard'])
    imgIds = coco.getImgIds(catIds=catIds)
    print(f"Found {len(imgIds)} images with specified categories")
    # Load and display an image
    if imaIds:
        img = coco.loadImgs(imgIds[np.random.randint(0, len(imgIds))])[0]
        # Use URL to load image
        I = io.imread(img['coco url'])
        plt.figure(figsize=(12, 9))
        plt.imshow(I)
        plt.axis('off')
        # Load and display instance annotations
        annIds = coco.getAnnIds(imgIds=img['id'], catIds=catIds, iscrowd=None)
        anns = coco.loadAnns(annIds)
        plt.figure(figsize=(12, 9))
        plt.imshow(I)
        plt.axis('off')
        coco.showAnns(anns)
        plt.show()
except Exception as e:
    print(f"Error working with COCO: {e}")
    print("Note: You need to download the COCO dataset separately.")
    # Display a sample image from the COCO website instead
    try:
        url = "http://images.cocodataset.org/val2017/000000013729.jpg"
        response = requests.get(url)
        I = io.imread(BytesIO(response.content))
```

```
plt.figure(figsize=(12, 9))
  plt.imshow(I)
  plt.axis('off')
  plt.title("Sample COCO Image")
  plt.show()
except:
  print("Could not display sample image")
```

# Natural Language Processing

#### **GLUE Benchmark**

- **Description**: General Language Understanding Evaluation benchmark
- Contents: Collection of 9 NLP tasks for evaluating language understanding
- Access: https://gluebenchmark.com/
- **Documentation**: nyu-mll/GLUE-baselines

```
# Example: Loading GLUE tasks with the datasets library
# !pip install datasets transformers
from datasets import load dataset
from transformers import AutoTokenizer
import pandas as pd
# Load a GLUE task (MNLI for example)
dataset = load dataset("glue", "mnli")
# Print dataset structure
print("Dataset structure:")
print(dataset)
# Load a tokenizer
tokenizer = AutoTokenizer.from pretrained("bert-base-uncased")
# Display some examples
print("\nExample data:")
df = pd.DataFrame(dataset["train"][:5])
print(df[["premise", "hypothesis", "label"]])
# Define a function to tokenize the data
def tokenize function(examples):
    return tokenizer(
        examples["premise"],
        examples["hypothesis"],
        padding="max_length",
        truncation=True,
    )
# Tokenize the dataset
tokenized_datasets = dataset.map(tokenize_function, batched=True)
print("\nTokenized dataset features:")
print(tokenized datasets["train"].column names)
```

#### **SQuAD**

- **Description**: Stanford Question Answering Dataset
- Contents: 100,000+ question-answer pairs on Wikipedia articles
- Access: <a href="https://rajpurkar.github.io/SQuAD-explorer/">https://rajpurkar.github.io/SQuAD-explorer/</a>
- **Documentation**: rajpurkar/SQuAD-explorer

```
# Example: Working with SQuAD dataset
# !pip install datasets transformers
from datasets import load dataset
import pandas as pd
import random
# Load SQuAD dataset
squad = load dataset("squad")
# Print dataset structure
print("Dataset structure:")
print(squad)
# Convert a few examples to DataFrame for easier viewing
train\ examples = []
for example in squad["train"][:5]:
    train examples.append({
        "question": example["question"],
        "context": example["context"][:100] + "...", # Truncate for display
        "answers": example["answers"]["text"][0]
    })
df = pd.DataFrame(train examples)
print("\nSample questions and answers:")
print(df)
# Select a random example and show full context
random idx = random.randint(0, len(squad["train"]) - 1)
example = squad["train"][random_idx]
print("\nRandom example:")
print(f"Question: {example['question']}")
print(f"Answer: {example['answers']['text'][0]}")
print(f"\nContext: {example['context']}")
```

#### WikiText

- **Description**: Language modeling dataset of Wikipedia articles
- **Contents**: 103 million words with long-context dependencies
- Access: <a href="https://blog.salesforceairesearch.com/the-wikitext-long-term-dependency-language-modeling-dataset/">https://blog.salesforceairesearch.com/the-wikitext-long-term-dependency-language-modeling-dataset/</a>
- **Documentation**: salesforce/awd-lstm-lm

```
# Example: Loading WikiText dataset
# !pip install datasets
from datasets import load dataset
import matplotlib.pyplot as plt
import numpy as np
# Load WikiText-2 dataset
wikitext = load_dataset("wikitext", "wikitext-2-raw-v1")
# Print dataset structure
print("Dataset structure:")
print(wikitext)
# Display a sample from the training set
print("\nSample from training set:")
print(wikitext["train"][100]["text"])
# Compute statistics
train_lens = [len(sample["text"].split()) for sample in wikitext["train"] if samp
# Plot distribution of sequence lengths
plt.figure(figsize=(10, 6))
plt.hist(train_lens, bins=50)
plt.xlabel('Sequence Length (words)')
plt.ylabel('Count')
plt.title('Distribution of Sequence Lengths in WikiText-2')
plt.axvline(np.mean(train_lens), color='r', linestyle='dashed', linewidth=1, labe
plt.axvline(np.median(train_lens), color='g', linestyle='dashed', linewidth=1, la
plt.legend()
plt.show()
# Count total words
total words = sum(train lens)
print(f"\nTotal words in training set: {total words:,}")
# Compute vocabulary size (unique words)
all_text = " ".join([sample["text"] for sample in wikitext["train"] if sample["te
vocab = set(all text.split())
print(f"Vocabulary size: {len(vocab):,} unique words")
```

### Reinforcement Learning

#### OpenAl Gym

- **Description**: Toolkit for developing and comparing RL algorithms
- **Contents**: Collection of environments from simple to complex

- Access: https://gym.openai.com/
- **Documentation**: openai/gym

```
# Example: Basic usage of OpenAI Gym
# !pip install gym
import gym
import numpy as np
import matplotlib.pyplot as plt
from matplotlib import animation
# Create an environment
env = gym.make('CartPole-v1')
# Reset the environment
observation = env.reset()
# Run a simple random policy
frames = []
done = False
for _ in range(100):
    # Render the environment
    frames.append(env.render(mode='rgb_array'))
    # Take a random action
    action = env.action space.sample()
    # Step in the environment
    observation, reward, done, info = env.step(action)
    if done:
        observation = env.reset()
env.close()
# Display the animation
plt.figure(figsize=(8, 6))
def update frame(i):
    plt.clf()
    plt.imshow(frames[i])
    plt.axis('off')
    return [plt.gca()]
ani = animation.FuncAnimation(plt.gcf(), update_frame, frames=len(frames), interv
plt.show()
# Print environment details
print(f"Observation space: {env.observation_space}")
print(f"Action space: {env.action space}")
```

# DeepMind Lab

- Description: 3D learning environment based on Quake III Arena
- Contents: Navigation and puzzle-solving tasks in 3D environment
- Access: 🖸 deepmind/lab
- Documentation: 🖸 deepmind/lab

```
# Example: Using DeepMind Lab (note: requires separate installation)
try:
   import deepmind lab
    import numpy as np
    import matplotlib.pyplot as plt
   # Create a simple DeepMind Lab environment
   lab = deepmind lab.Lab(
        'seekavoid_arena_01', # Level name
        ['RGB INTERLEAVED'], # Observations to return
       config={
            'width': 320,
            'height': 240,
            'fps': 60
       }
   )
   # Reset the environment
   lab.reset()
   # Run a few random actions and display observations
   for in range(10):
        # Generate a random action (forward/backward, strafe, look)
       action = np.zeros([7], dtype=np.intc) # DeepMind Lab actions are 7D
       action[0] = np.random.choice([-1, 0, 1]) # Forward/backward
       action[2] = np.random.choice([-1, 0, 1]) # Strafe left/right
       # Step in the environment
       reward = lab.step(action, num_steps=4) # Execute 4 frames
       # Get observation
       obs = lab.observations()['RGB_INTERLEAVED']
       # Display observation
       plt.figure(figsize=(8, 6))
       plt.imshow(obs)
       plt.title(f"Reward: {reward}")
       plt.axis('off')
       plt.show()
   lab.close()
except ImportError:
   print("DeepMind Lab not installed. It requires a separate installation.")
   print("See: https://github.com/deepmind/lab for installation instructions.")
```

#### MuJoCo

- **Description**: Physics-based robot simulation environment
- Contents: Various robot control tasks

- Access: O openai/mujoco-py
- **Documentation**: https://www.gymlibrary.ml/environments/mujoco/

```
# Example: Using MuJoCo environments in Gym
# !pip install gym mujoco-py
try:
    import gym
    import numpy as np
    import matplotlib.pyplot as plt
    from matplotlib import animation
    # Create a MuJoCo environment
    env = gym.make('HalfCheetah-v2')
    observation = env.reset()
    # Run a random policy
    frames = []
    for in range(100):
        # Render
        frames.append(env.render(mode='rgb array'))
        # Random action
        action = env.action space.sample()
        # Step
        observation, reward, done, info = env.step(action)
        if done:
            observation = env.reset()
    env.close()
    # Display the animation
    plt.figure(figsize=(8, 6))
    def update frame(i):
        plt.clf()
        plt.imshow(frames[i])
        plt.axis('off')
        return [plt.gca()]
    ani = animation.FuncAnimation(plt.gcf(), update_frame, frames=len(frames), in
    plt.show()
    # Print environment details
    print(f"Observation space: {env.observation space}")
    print(f"Action space: {env.action_space}")
except ImportError as e:
    print(f"Error: {e}")
    print("MuJoCo requires a separate installation and license.")
    print("See: https://github.com/openai/mujoco-py for installation instructions
```

# **B.3 NeuroAl Specific Datasets**

#### Brain-Score

- **Description**: Benchmark for neural network models of the visual system
- Contents: Neural and behavioral data for evaluating computational models
- Access: https://www.brain-score.org/
- **Documentation**: brain-score/brain-score

```
# Example: Using Brain-Score to evaluate models
# !pip install brain-score
try:
    import brainscore
    from brainscore.benchmarks.public benchmarks import MajajHongITPublicBenchmar
    from brainscore.model_interface import BrainModel
    # Create a simple mock model for illustration
    class MockModel(BrainModel):
        def __init__(self):
            self.recording_target = None
            self.stimuli identifier = None
        def look_at(self, stimuli, number_of_trials=1):
            import numpy as np
            # Return random activations
            return np.random.rand(len(stimuli), 10) # 10 neurons
        def start_task(self, task, fitting_stimuli=None):
            pass
        def start_recording(self, recording_target='IT', time_bins=None):
            self.recording_target = recording_target
    # Create the model
    model = MockModel()
    # Create benchmark
    benchmark = MajajHongITPublicBenchmark()
    # Score the model
    score = benchmark(model)
    print(f"Model score: {score}")
except ImportError:
    print("brain-score not installed. Use: pip install brain-score")
    print("Note: brain-score may have specific dependencies and requirements.")
```

# Neural Data Challenge

• **Description**: Neural decoding competitions

• Contents: Neural recordings with specific decoding tasks

• Access: https://neurodatachallenge.org/

• Documentation: Varies by challenge

```
# Example: Generic neural decoding pipeline (using simulated data)
import numpy as no
from sklearn.model selection import train test split
from sklearn.preprocessing import StandardScaler
from sklearn.linear model import LogisticRegression
from sklearn.metrics import accuracy score, classification report
import matplotlib.pyplot as plt
# Simulate neural data with 3 target classes
def simulate neural data(n samples=1000, n neurons=100, n classes=3):
    # Create class-specific patterns
    X = np.zeros((n samples, n neurons))
    y = np.zeros(n_samples, dtype=int)
    # Generate different patterns for each class
    patterns = np.random.randn(n_classes, n_neurons)
    samples per class = n samples // n classes
    for c in range(n_classes):
        start idx = c * samples per class
        end_idx = (c + 1) * samples_per_class if c < n_classes - 1 else n_samples
        # Assign class labels
        y[start idx:end idx] = c
        # Generate neural activity
        X[start_idx:end_idx] = patterns[c] + np.random.randn(end_idx - start_idx,
    return X, y
# Generate simulated data
X, y = simulate_neural_data()
# Split data
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_s
# Preprocess: Standardize features
scaler = StandardScaler()
X train scaled = scaler.fit transform(X train)
X test scaled = scaler.transform(X test)
# Train a decoder
model = LogisticRegression(max iter=1000)
model.fit(X train scaled, y train)
# Evaluate
y_pred = model.predict(X_test_scaled)
accuracy = accuracy score(y test, y pred)
print(f"Decoding accuracy: {accuracy:.3f}")
print(classification_report(y_test, y_pred))
# Plot confusion matrix
from sklearn.metrics import confusion matrix
```

```
import seaborn as sns
cm = confusion_matrix(y_test, y_pred)
plt.figure(figsize=(8, 6))
sns.heatmap(cm, annot=True, fmt='d', cmap='Blues')
plt.xlabel('Predicted Label')
plt.ylabel('True Label')
plt.title('Confusion Matrix')
plt.show()
# Visualize feature importance
coef = model.coef_
plt.figure(figsize=(12, 4))
for i, c in enumerate(coef):
    plt.subplot(1, len(coef), i+1)
    plt.bar(range(len(c)), c)
    plt.title(f'Class {i} vs Rest')
    plt.xlabel('Neuron ID')
    plt.ylabel('Weight')
plt.tight_layout()
plt.show()
```

# Algonauts Project

- **Description**: Bridging human and machine vision
- Contents: fMRI data for training and evaluating computer vision models
- Access: http://algonauts.csail.mit.edu/
- Documentation: 

  Brain-Bridge-Lab/Algonauts2023

```
# Example: Working with Algonauts data (simulated)
import numpy as no
import matplotlib.pyplot as plt
from sklearn.decomposition import PCA
from sklearn.cross decomposition import PLSRegression
from sklearn.model selection import KFold
from sklearn.metrics import r2 score
# Simulate fMRI data and CNN features
def simulate algonauts data(n samples=500, n voxels=1000, n cnn features=4096):
    # Simulate CNN features
    cnn features = np.random.randn(n samples, n cnn features)
    # Create "true" mapping weights
    true weights = np.random.randn(n cnn features, n voxels) * 0.01
    # Generate fMRI data using the features and weights
    fmri data = cnn features @ true weights + np.random.randn(n samples, n voxels
    return cnn features, fmri data
# Generate simulated data
cnn_features, fmri_data = simulate_algonauts_data()
# Reduce dimensionality of CNN features for visualization
pca = PCA(n components=100)
cnn_reduced = pca.fit_transform(cnn_features)
# Set up cross-validation
kf = KFold(n splits=5, shuffle=True, random state=42)
# Train a model to predict fMRI responses from CNN features
pls = PLSRegression(n components=20)
# For simplicity, just train on a subset of voxels
voxel subset = np.random.choice(fmri data.shape[1], 100, replace=False)
fmri_subset = fmri_data[:, voxel_subset]
# Cross-validation
print("Running cross-validation...")
r2 scores = []
for train_idx, test_idx in kf.split(cnn_features):
    # Train/test split
    X train, X test = cnn features[train idx], cnn features[test idx]
    y_train, y_test = fmri_subset[train_idx], fmri_subset[test_idx]
    # Fit model
    pls.fit(X_train, y_train)
    # Predict
    y_pred = pls.predict(X_test)
    # Compute R<sup>2</sup> for each voxel
```

```
r2 = [r2_score(y_test[:, i], y_pred[:, i]) for i in range(y_test.shape[1])]
    r2_scores.append(np.mean(r2))

print(f"Mean R² across folds: {np.mean(r2_scores):.3f}")

# Visualize distribution of R² scores
plt.figure(figsize=(10, 6))
plt.hist(r2_scores, bins=10)
plt.xlabel('Mean R² Score')
plt.ylabel('Count')
plt.title('Distribution of R² Scores Across CV Folds')
plt.grid(True)
plt.show()
```

# **B.4 Data Loading Examples**

The examples below demonstrate how to load specific file formats common in neuroscience research.

# **Loading NIfTI Files**

```
# Example: Working with NIfTI files (neuroimaging data)
# !pip install nibabel matplotlib
import nibabel as nib
import matplotlib.pyplot as plt
import numpy as np
import requests
import os
# Download a sample NIfTI file if needed
nifti_url = "https://ndownloader.figshare.com/files/12965017"
filename = "example.nii.gz"
if not os.path.exists(filename):
    try:
        print(f"Downloading {filename}...")
        response = requests.get(nifti url)
        with open(filename, 'wb') as f:
            f.write(response.content)
        print("Download complete.")
    except Exception as e:
        print(f"Error downloading file: {e}")
# Load NIfTI file
trv:
    img = nib.load(filename)
    # Get data as numpy array
    data = img.get_fdata()
    print(f"Image shape: {data.shape}")
    print(f"Data type: {data.dtype}")
    # Get header information
    header = img.header
    print(f"\nVoxel dimensions: {header.get zooms()}")
    print(f"Units: {header.get xyzt units()}")
    # Display central slices
    if len(data.shape) >= 3:
        middle x = data.shape[0] // 2
        middle y = data.shape[1] // 2
        middle z = data.shape[2] // 2
        plt.figure(figsize=(12, 4))
        plt.subplot(131)
        plt.imshow(data[middle_x, :, :].T, cmap='gray')
        plt.title('Sagittal View')
        plt.axis('off')
        plt.subplot(132)
        plt.imshow(data[:, middle_y, :].T, cmap='gray')
```

```
plt.title('Coronal View')
       plt.axis('off')
       plt.subplot(133)
       plt.imshow(data[:, :, middle_z].T, cmap='gray')
       plt.title('Axial View')
       plt.axis('off')
       plt.tight_layout()
       plt.show()
except Exception as e:
   print(f"Error processing NIfTI file: {e}")
   print("Will use a simulated 3D volume instead for demonstration:")
   # Create a simple simulated volume
   sim data = np.zeros((30, 30, 30))
   # Add a sphere in the middle
   x, y, z = np.ogrid[-15:15, -15:15]
   sphere = x*x + y*y + z*z <= 10*10
   sim_data[sphere] = 1
   # Display central slices
   middle x = sim data.shape[0] // 2
   middle_y = sim_data.shape[1] // 2
   middle z = sim data.shape[2] // 2
   plt.figure(figsize=(12, 4))
   plt.subplot(131)
   plt.imshow(sim_data[middle_x, :, :].T, cmap='gray')
   plt.title('Sagittal View (Simulated)')
   plt.axis('off')
   plt.subplot(132)
   plt.imshow(sim_data[:, middle_y, :].T, cmap='gray')
   plt.title('Coronal View (Simulated)')
   plt.axis('off')
   plt.subplot(133)
   plt.imshow(sim_data[:, :, middle_z].T, cmap='gray')
   plt.title('Axial View (Simulated)')
   plt.axis('off')
   plt.tight_layout()
   plt.show()
```

### Working with EEG Data

```
# Example: Working with EEG data
# !pip install mne
try:
   import mne
    import numpy as np
    import matplotlib.pyplot as plt
    # Download sample EEG data
    sample_data_folder = mne.datasets.sample.data_path()
    sample_data_raw_file = f"{sample_data_folder}/MEG/sample/sample_audvis_raw.fi
    # Load the data
    raw = mne.io.read raw fif(sample data raw file, preload=True)
    # Print information about the dataset
   print(raw.info)
    # Extract EEG channels only
    raw.pick_types(meg=False, eeg=True, eog=False, stim=False)
    # Filter the data
    raw.filter(l_freq=1.0, h_freq=40.0)
    # Plot EEG data
    raw.plot(duration=5, n_channels=10, scalings='auto')
   # Extract events
    events = mne.find_events(raw, stim_channel='STI 014')
    event_id = {'auditory/left': 1, 'auditory/right': 2, 'visual/left': 3, 'visual/
    # Create epochs
    epochs = mne.Epochs(raw, events, event id, tmin=-0.2, tmax=0.5,
                      proj=True, baseline=(None, 0), preload=True)
    # Plot evoked responses
    evoked = epochs['auditory/left'].average()
    evoked.plot()
    # Create and plot a topographic map
    evoked.plot_topomap(times=[0.1], ch_type='eeg')
    # Plot time-frequency representation
    frequencies = np.arange(6, 20, 2) # Frequencies from 6-20Hz
    power = mne.time frequency.tfr morlet(epochs['auditory/left'], freqs=frequenc
                                        n cycles=2, return itc=False)
   power.plot([0])
except Exception as e:
    print(f"Error processing EEG data: {e}")
    print("\nNote: MNE-Python requires sample data to be downloaded.")
    print("You can still work with EEG data by following these steps:")
    print("1. Install MNE: pip install mne")
```

```
print("2. Download sample data: mne.datasets.sample.data path()")
print("3. Load and process EEG data as shown in the example")
# Create a simulated EEG dataset for demonstration
try:
    import numpy as np
    import matplotlib.pyplot as plt
    # Parameters
    n channels = 16
    sfreq = 256 \# Hz
    duration = 5 # seconds
    t = np.arange(0, duration, 1/sfreq)
    n \text{ samples} = len(t)
    # Generate simulated EEG signals
    sim_eeg = np.zeros((n_channels, n_samples))
    for i in range(n channels):
        # Combine different frequency components with channel-specific weight
        alpha = 0.5 * np.sin(2 * np.pi * 10 * t) # 10 Hz alpha
        beta = 0.25 * np.sin(2 * np.pi * 20 * t) # 20 Hz beta
        theta = 0.3 * np.sin(2 * np.pi * 5 * t) # 5 Hz theta
        delta = 0.4 * np.sin(2 * np.pi * 2 * t) # 2 Hz delta
        # Mix components with random weights and add noise
        weights = np.random.rand(4)
        weights = weights / np.sum(weights)
        sim_eeg[i] = (weights[0] * alpha +
                     weights[1] * beta +
                     weights[2] * theta +
                     weights[3] * delta +
                     0.1 * np.random.randn(n_samples))
    # Plot simulated EEG data
    plt.figure(figsize=(10, 8))
    for i in range(min(8, n_channels)):
        plt.subplot(8, 1, i+1)
        plt.plot(t, sim_eeg[i])
        plt.ylabel(f'Ch {i+1}')
        if i == 0:
            plt.title('Simulated EEG Data')
        if i == 7:
            plt.xlabel('Time (s)')
    plt.tight layout()
    plt.show()
    # Create a simulated topographic map
    plt.figure(figsize=(8, 8))
    channel locs = []
    for i in range(n_channels):
        # Create circular layout
        angle = i * 2 * np.pi / n_channels
        x = 0.8 * np.cos(angle)
```

```
y = 0.8 * np.sin(angle)
        channel_locs.append((x, y))
   # Create a grid for interpolation
   grid_size = 100
   xi = np.linspace(-1, 1, grid_size)
   yi = np.linspace(-1, 1, grid_size)
   X, Y = np.meshgrid(xi, yi)
   # Simulate some values at a given time point
   values = np.random.rand(n_channels)
   # Plot values at channel locations
   plt.scatter([loc[0] for loc in channel_locs], [loc[1] for loc in channel_
               c=values, s=200, cmap='RdBu_r')
   # Add channel labels
   for i, loc in enumerate(channel locs):
        plt.text(loc[0], loc[1], f'{i+1}', ha='center', va='center', color='w
   plt.title('Simulated EEG Topographic Map')
   plt.colorbar(label='Activation')
   plt.axis('equal')
   plt.axis('off')
   plt.tight_layout()
   plt.show()
except Exception as e:
   print(f"Error creating simulated EEG data: {e}")
```

### Working with Spike Train Data

```
# Example: Working with spike train data
import numpy as np
import matplotlib.pyplot as plt
from matplotlib.ticker import MaxNLocator
# Simulate spike train data for multiple neurons
def simulate_spike_trains(n_neurons=10, rate=10, duration=5.0, bin_size=0.001):
    """Simulate Poisson spike trains for multiple neurons"""
    # Create time bins
    bins = np.arange(0, duration + bin size, bin size)
    n bins = len(bins) - 1
    # Initialize spike rasters and times
    spike_rasters = np.zeros((n_neurons, n_bins), dtype=bool)
    spike_times = [[] for _ in range(n_neurons)]
    # Generate spikes for each neuron
    for i in range(n neurons):
        # Rate might vary between neurons
        neuron_rate = rate * (0.5 + np.random.rand())
        # Probability of spike in each bin
        spike prob = neuron rate * bin size
        # Generate spikes
        for j in range(n bins):
            if np.random.rand() < spike_prob:</pre>
                spike_rasters[i, j] = True
                spike times[i].append(bins[j])
    return spike rasters, spike times, bins
# Simulate data
n neurons = 10
duration = 5.0 # seconds
bin size = 0.001 # 1 ms bins
spike rasters, spike times, bins = simulate spike trains(
    n_neurons=n_neurons, rate=10, duration=duration, bin_size=bin_size)
# Plot spike raster
plt.figure(figsize=(14, 6))
# Raster plot
plt.subplot(2, 1, 1)
for i in range(n neurons):
    plt.plot(spike_times[i], np.ones_like(spike_times[i]) * i, '|', markersize=3)
plt.ylabel('Neuron ID')
plt.title('Spike Raster Plot')
plt.ylim(-0.5, n neurons -0.5)
plt.gca().yaxis.set_major_locator(MaxNLocator(integer=True))
```

```
# PSTH (Population Peristimulus Time Histogram)
plt.subplot(2, 1, 2)
bin edges = np.arange(0, duration + 0.1, 0.1) # 100 ms bins for PSTH
population rate = np.zeros((n neurons, len(bin edges) - 1))
for i in range(n neurons):
    # Count spikes in each time bin
    counts, = np.histogram(spike times[i], bins=bin edges)
    # Convert to firing rate (Hz)
    population rate[i] = counts / 0.1 # 0.1s bin size
# Average across neurons
avg population rate = np.mean(population rate, axis=0)
# Plot PSTH
plt.bar(bin_edges[:-1], avg_population_rate, width=0.1, alpha=0.7)
plt.xlabel('Time (s)')
plt.ylabel('Firing Rate (Hz)')
plt.title('Population PSTH')
plt.tight layout()
plt.show()
# Calculate and plot various spike train statistics
plt.figure(figsize=(14, 8))
# 1. Calculate ISI (Inter-Spike Intervals) for all neurons
isis = [np.diff(times) for times in spike times if len(times) > 1]
# 2. Calculate CV (Coefficient of Variation) of ISIs
cvs = [np.std(isi)/np.mean(isi) if len(isi) > 0 else np.nan for isi in isis]
# 3. Calculate mean firing rates
rates = [len(times)/duration for times in spike_times]
# Plot ISI histogram
plt.subplot(2, 2, 1)
all isis = np.concatenate(isis) if isis else np.array([])
if len(all isis) > 0:
    plt.hist(all isis, bins=50, density=True, alpha=0.7)
    plt.xlabel('Inter-Spike Interval (s)')
    plt.ylabel('Density')
    plt.title('ISI Distribution')
else:
    plt.text(0.5, 0.5, "No sufficient spikes for ISI", ha='center')
# Plot CV distribution
plt.subplot(2, 2, 2)
valid cvs = [cv for cv in cvs if not np.isnan(cv)]
if valid_cvs:
    plt.hist(valid cvs, bins=10, alpha=0.7)
    plt.axvline(x=1.0, color='r', linestyle='--', label='Poisson')
    plt.xlabel('CV of ISI')
```

```
plt.ylabel('Count')
    plt.title('CV Distribution')
    plt.legend()
    plt.text(0.5, 0.5, "No sufficient spikes for CV", ha='center')
# Plot firing rate distribution
plt.subplot(2, 2, 3)
plt.hist(rates, bins=10, alpha=0.7)
plt.xlabel('Firing Rate (Hz)')
plt.ylabel('Count')
plt.title('Firing Rate Distribution')
# Plot autocorrelation for the first neuron
plt.subplot(2, 2, 4)
if len(spike rasters[0]) > 0:
    # Convert spike raster to 0/1 array
    spikes 01 = spike rasters[0].astype(int)
    # Compute autocorrelation
    max_{lag} = int(0.5 / bin_{size}) # 500 ms max lag
    lags = np.arange(-max_lag, max_lag + 1) * bin_size
    # Manual autocorrelation (numpy.correlate doesn't handle this well)
    autocorr = np.zeros(len(lags))
    for i, lag in enumerate(range(-max lag, max lag + 1)):
        if lag < 0:
            autocorr[i] = np.sum(spikes 01[:-lag] * spikes 01[-lag:])
            autocorr[i] = np.sum(spikes_01[lag:] * spikes_01[:-lag if lag > 0 els
    # Plot autocorrelation
    plt.plot(lags, autocorr)
    plt.xlabel('Lag (s)')
    plt.ylabel('Autocorrelation')
    plt.title(f'Autocorrelation (Neuron 0)')
else:
    plt.text(0.5, 0.5, "No spikes for autocorrelation", ha='center')
plt.tight_layout()
plt.show()
```

These examples and resources should provide a comprehensive starting point for working with a wide range of neuroscience and AI datasets.

# B.5 Neurological and Neurodegenerative Disease

## **Datasets**

This section catalogs datasets specifically focused on neurological and neurodegenerative diseases, which are valuable resources for developing diagnostic, prognostic, and treatment monitoring AI systems.

### Alzheimer's Disease

### Alzheimer's Disease Neuroimaging Initiative (ADNI)

- Description: Longitudinal multimodal data to track Alzheimer's progression
- Contents: MRI, PET, cognitive assessments, biospecimen data from 800+ subjects
- Access: http://adni.loni.usc.edu/ (requires application)
- Documentation: http://adni.loni.usc.edu/methods/

```
# Example: Working with ADNI MRI data
# Note: You need approved access to ADNI to download data
import nibabel as nib
import matplotlib.pyplot as plt
import numpy as np
import os
def visualize adni mri(filepath):
    """Visualize a 3D MRI from ADNI dataset."""
    # Load MRI volume
    img = nib.load(filepath)
    data = img.get fdata()
    # Get middle slices for each plane
    x \text{ mid} = data.shape[0] // 2
    y = data.shape[1] // 2
    z_{mid} = data.shape[2] // 2
    # Create a figure with three subplots
    fig, axes = plt.subplots(1, 3, figsize=(15, 5))
    # Sagittal view (YZ plane)
    axes[0].imshow(data[x_mid, :, :].T, cmap='gray', origin='lower')
    axes[0].set_title('Sagittal View')
    # Coronal view (XZ plane)
    axes[1].imshow(data[:, y_mid, :].T, cmap='gray', origin='lower')
    axes[1].set title('Coronal View')
    # Axial view (XY plane)
    axes[2].imshow(data[:, :, z_mid].T, cmap='gray', origin='lower')
    axes[2].set title('Axial View')
    plt.tight_layout()
    plt.show()
# Example of preprocessing ADNI data for machine learning
def preprocess adni for ml(mri files, diagnoses):
    0.00
    Preprocess ADNI MRI files for ML classification.
    Parameters:
    mri files: List of paths to MRI files
    diagnoses: List of diagnoses (e.g., 'CN', 'MCI', 'AD')
    Returns:
    features: Extracted features for each MRI
    labels: Corresponding diagnostic labels
    # For demonstration - would require actual data
    features = []
    for mri file in mri files:
```

```
# Load MRI
        img = nib.load(mri file)
        data = img.get fdata()
        # Example feature extraction (hippocampal ROI)
        # This is simplified - real implementation would use proper segmentation
        x_range = slice(90, 110) # Example hippocampal coordinates
        y range = slice(60, 80)
        z_range = slice(60, 80)
        hippocampus_roi = data[x_range, y_range, z_range]
        # Extract features (e.g., mean, variance, etc.)
        features.append([
            np.mean(hippocampus_roi),
            np.var(hippocampus roi),
            np.max(hippocampus_roi),
            np.min(hippocampus roi)
        ])
    # Convert diagnoses to numeric labels
    label_map = {'CN': 0, 'MCI': 1, 'AD': 2} # Control, Mild Cognitive Impairmen
    labels = [label_map[d] for d in diagnoses]
    return np.array(features), np.array(labels)
# Usage example (pseudocode)
# mri_files = ['/path/to/adni/001.nii', '/path/to/adni/002.nii', ...]
\# diagnoses = ['CN', 'MCI', 'AD', ...]
# features, labels = preprocess_adni_for_ml(mri_files, diagnoses)
```

#### OASIS (Open Access Series of Imaging Studies)

- **Description**: Free datasets for brain aging and Alzheimer's research
- Contents: MRI scans, demographics, clinical data for 400+ subjects
- Access: https://www.oasis-brains.org/
- **Documentation**: https://www.oasis-brains.org/#data

```
# Example: Downloading and visualizing OASIS-3 data
# !pip install nilearn nibabel matplotlib
import os
import nibabel as nib
import numpy as np
import matplotlib.pyplot as plt
from nilearn import datasets, plotting
# OASIS data can be accessed via nilearn
oasis dataset = datasets.fetch oasis vbm(n subjects=5)
print(f"Dataset directory: {oasis_dataset.dirname}")
print(f"Gray matter anatomy: {len(oasis dataset.gray matter maps)}")
# Visualize a gray matter map
gm_img = oasis_dataset.gray_matter_maps[0]
plotting.plot_img(gm_img, title="OASIS Gray Matter Map")
plt.show()
# Plot the first 5 subject MRIs
fig, axes = plt.subplots(1, min(5, len(oasis dataset.gray matter maps)), figsize=
for i, gm file in enumerate(oasis dataset.gray matter maps[:5]):
    # Load the image
    img = nib.load(gm file)
    # Get a slice at the middle of the 3rd dimension
    data = img.get fdata()
    slice idx = data.shape[2] // 2
    # Plot on the corresponding axis
    axes[i].imshow(data[:, :, slice_idx], cmap='gray')
    axes[i].set title(f'Subject {i+1}')
    axes[i].axis('off')
plt.tight layout()
plt.show()
# Plot clinical information
y = oasis_dataset.ext_vars['y'] # Mini-Mental State Examination (MMSE) scores
age = oasis dataset.ext vars['age']
plt.figure(figsize=(10, 6))
plt.scatter(age, y)
plt.xlabel('Age')
plt.ylabel('MMSE Score')
plt.title('Age vs. Cognitive Function in OASIS Dataset')
plt.grid(True)
plt.show()
```

## Parkinson's Disease

## Parkinson's Progression Markers Initiative (PPMI)

- **Description**: Comprehensive dataset for Parkinson's disease biomarkers
- Contents: Clinical, imaging, genetic, and biospecimen data from 1,400+ subjects
- Access: https://www.ppmi-info.org/ (requires application)
- **Documentation**: https://www.ppmi-info.org/access-data-specimens/

```
# Example: Analyzing PPMI clinical data
# Note: You need approved access to PPMI to download data
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns
def analyze ppmi clinical(clinical file):
    """Analyze PPMI clinical data from CSV file."""
    # Load clinical data
    df = pd.read csv(clinical file)
    # Example exploratory analysis
    print(f"Number of participants: {df['PATNO'].nunique()}")
    print(f"Average age: {df['AGE'].mean():.1f} years")
   # UPDRS (Unified Parkinson's Disease Rating Scale) analysis
    if 'UPDRS TOT' in df.columns:
        # Distribution of UPDRS scores
        plt.figure(figsize=(10, 6))
        sns.histplot(df['UPDRS TOT'].dropna(), kde=True)
        plt.xlabel('Total UPDRS Score')
        plt.ylabel('Frequency')
        plt.title('Distribution of UPDRS Scores in PPMI Dataset')
        plt.show()
    # Compare scores between different disease stages
    if 'DIAGNOSIS' in df.columns and 'UPDRS_TOT' in df.columns:
        plt.figure(figsize=(10, 6))
        sns.boxplot(x='DIAGNOSIS', y='UPDRS_TOT', data=df)
        plt.xlabel('Diagnosis Group')
        plt.ylabel('UPDRS Score')
        plt.title('UPDRS Scores by Diagnosis Group')
        plt.show()
    return df
# Example of accessing DaTscan images
def visualize datscan(datscan file):
    """Visualize a DaTscan SPECT image from PPMI."""
    # Load DaTscan image
    img = nib.load(datscan file)
    data = img.get fdata()
    # Get middle slice
    slice_idx = data.shape[2] // 2
   # Display
    plt.figure(figsize=(8, 8))
    plt.imshow(data[:, :, slice_idx], cmap='hot')
    plt.colorbar(label='Intensity')
   plt.title('DaTscan SPECT Image (Axial Slice)')
    plt.axis('off')
    plt.show()
```

```
# Usage example (pseudocode)
# clinical_data = analyze_ppmi_clinical('/path/to/ppmi/clinical.csv')
# visualize_datscan('/path/to/ppmi/datscan.nii')
```

### PhysioNet Freeze-Gait in Parkinson's Disease

- Description: Gait data for patients with and without freezing of gait
- Contents: Accelerometer data from 16 PD patients during various activities
- Access: https://physionet.org/content/freeze-gait-pd/1.0.0/
- **Documentation**: https://physionet.org/content/freeze-gait-pd/1.0.0/

```
# Example: Analyzing freezing of gait data from PhysioNet
# !pip install wfdb
import wfdb
import numpy as np
import matplotlib.pyplot as plt
from scipy import signal
# Path to the dataset (update with your path or download)
# data_path = '/path/to/physionet/freeze-gait-pd/1.0.0/'
data path = 'freeze-gait-pd/1.0.0/'
def analyze freeze gait(record name, data path):
    """Analyze a freeze of gait recording."""
        # Load the data
        record = wfdb.rdrecord(data_path + record_name)
        signals = record.p_signal
        # Extract the acceleration channels
        # Typically channels are x, y, z acceleration
        x_{acc} = signals[:, 0]
        y acc = signals[:, 1]
        z_{acc} = signals[:, 2]
        # Time vector
        fs = record.fs # Sampling frequency
        t = np.arange(len(x_acc)) / fs
        # Compute the frequency spectrum
        f, Pxx x = signal.welch(x acc, fs, nperseg=1024)
        _, Pxx_y = signal.welch(y_acc, fs, nperseg=1024)
        _, Pxx_z = signal.welch(z_acc, fs, nperseg=1024)
        # Plot time domain signals
        plt.figure(figsize=(15, 10))
        plt.subplot(2, 1, 1)
        plt.plot(t, x acc, label='X')
        plt.plot(t, y_acc, label='Y')
        plt.plot(t, z acc, label='Z')
        plt.xlabel('Time (s)')
        plt.ylabel('Acceleration')
        plt.title(f'Acceleration Data - {record name}')
        plt.legend()
        plt.grid(True)
        # Plot frequency domain - useful for detecting freezing frequency bands
        plt.subplot(2, 1, 2)
        plt.semilogy(f, Pxx_x, label='X')
        plt.semilogy(f, Pxx_y, label='Y')
        plt.semilogy(f, Pxx_z, label='Z')
        plt.xlabel('Frequency (Hz)')
        plt.ylabel('PSD (g^2/Hz)')
```

```
plt.title('Power Spectral Density')
        plt.axvspan(3, 8, color='r', alpha=0.3, label='Freezing Band')
        plt.legend()
        plt.grid(True)
        plt.tight_layout()
        plt.show()
        # Calculate freezing index
        # Freezing Index = Power in freezing band (3-8 Hz) / Power in locomotion
        freeze_band = (f \ge 3) & (f \le 8)
        locomotion band = (f \ge 0.5) & (f \le 3)
        freeze power = np.mean(Pxx y[freeze band])
        locomotion power = np.mean(Pxx y[locomotion band])
        freezing_index = freeze_power / locomotion_power if locomotion_power > 0
        print(f"Freezing Index: {freezing index:.2f}")
        return freezing index
    except Exception as e:
        print(f"Error processing record {record_name}: {e}")
        return None
# Example usage (uncomment to use)
# analyze freeze gait('S01R01', data path)
```

## **Epilepsy**

#### Temple University EEG Corpus

- **Description**: Massive clinical EEG dataset from Temple University Hospital
- Contents: 30,000+ EEG recordings from 16,000+ patients, including many with epilepsy
- Access: <a href="https://isip.piconepress.com/projects/tuh\_eeg/">https://isip.piconepress.com/projects/tuh\_eeg/</a>
- **Documentation**: https://isip.piconepress.com/projects/tuh\_eeg/html/downloads.shtml

```
# Example: Working with Temple University EEG data
# !pip install pyedflib numpy matplotlib
import pyedflib
import numpy as np
import matplotlib.pyplot as plt
def read eeg file(edf file):
    """Read and plot EEG data from EDF file."""
        # Open the EDF file
        f = pyedflib.EdfReader(edf_file)
        # Get information about the file
        n channels = f.signals in file
        channel_names = f.getSignalLabels()
        fs = f.getSampleFrequency(0) # Assuming same sampling rate for all chann
        print(f"File: {edf file}")
        print(f"Number of channels: {n_channels}")
        print(f"Channel names: {channel names}")
        print(f"Sampling frequency: {fs} Hz")
        # Read a subset of channels (e.g., first 8)
        n_{display} = \min(8, n_{channels})
        signals = np.zeros((n_display, f.getNSamples()[0]))
        for i in range(n_display):
            signals[i, :] = f.readSignal(i)
        # Calculate time vector
        seconds = len(signals[0]) / fs
        t = np.linspace(0, seconds, len(signals[0]))
        # Plot the signals
        plt.figure(figsize=(15, 10))
        # Default display: 10 seconds window with appropriate scaling
        window size = min(10, seconds)
        samples to display = int(window size * fs)
        for i in range(n display):
            plt.subplot(n_display, 1, i+1)
            # Z-score normalize for display
            signal_std = np.std(signals[i][:samples_to_display])
            signal mean = np.mean(signals[i][:samples to display])
            normalized_signal = (signals[i][:samples_to_display] - signal_mean) /
            plt.plot(t[:samples_to_display], normalized_signal)
            plt.ylabel(channel names[i])
            plt.xlim(∅, window_size)
            # Only show x-axis for bottom plot
```

```
if i < n \text{ display } -1:
                plt.xticks([])
        plt.xlabel('Time (s)')
        plt.tight layout()
        plt.show()
        # Close the file
        f.close()
        return signals, channel names, fs
    except Exception as e:
        print(f"Error reading EEG file: {e}")
        return None, None, None
# Example seizure detection function
def detect seizures(eeg signal, fs, window size=1.0, overlap=0.5):
    Basic seizure detection based on signal energy and line length.
    Parameters:
    eeg signal: EEG signal (single channel)
    fs: Sampling frequency
    window_size: Analysis window size in seconds
    overlap: Window overlap fraction
    Returns:
    detection times: List of potential seizure onset times
    # Convert window size from seconds to samples
    window samples = int(window size * fs)
    step_samples = int(window_samples * (1 - overlap))
    # Features
    energy = []
    line length = []
    # Process windows
    for i in range(0, len(eeg_signal) - window_samples, step_samples):
        window = eeg signal[i:i + window samples]
        # Calculate energy (sum of squared amplitudes)
        energy.append(np.sum(window**2) / window samples)
        # Calculate line length (sum of absolute differences)
        line_length.append(np.sum(np.abs(np.diff(window))) / window_samples)
    # Normalize features
    energy = np.array(energy)
    energy_norm = (energy - np.mean(energy)) / (np.std(energy) if np.std(energy)
    line_length = np.array(line_length)
    11_norm = (line_length - np.mean(line_length)) / (np.std(line_length) if np.s
```

```
# Combined feature
    combined = energy norm + 11 norm
    # Threshold for detection (this is a simple approach, real detection would us
    threshold = 3.0
    # Find potential seizures
    detection_indices = np.where(combined > threshold)[0]
    # Convert indices to times
    detection times = []
    if len(detection indices) > 0:
        # Group consecutive detections
        groups = np.split(detection_indices, np.where(np.diff(detection_indices)
        for group in groups:
            if len(group) > 0:
                # Convert from window index to time
                onset_time = (group[0] * step_samples) / fs
                detection_times.append(onset_time)
    return detection times, combined
# Example usage (pseudocode)
# signals, channel_names, fs = read_eeg_file('/path/to/tuh_eeg/patient/eeg.edf')
# if signals is not None:
      # Analyze a specific channel (e.g., channel index 2)
      channel idx = 2
      detection_times, feature = detect_seizures(signals[channel_idx], fs)
#
      print(f"Potential seizure onsets at: {detection times} seconds")
```

#### Stroke

#### International Stroke Perfusion Imaging Registry (INSPIRE)

- Description: Multi-center dataset for stroke imaging
- **Contents**: CT, MRI, and clinical data from stroke patients
- Access: <a href="https://inspire.app/">https://inspire.app/</a>
- **Documentation**: Available upon registration

```
# Example: Simulated analysis of stroke perfusion images
import numpy as no
import matplotlib.pyplot as plt
from skimage import filters, measure
import nibabel as nib
def analyze stroke perfusion(perfusion file, t max threshold=6.0):
   Analyze perfusion imaging (Tmax map) to identify potential stroke penumbra.
   Parameters:
   perfusion file: Path to the perfusion Tmax map
   t_max_threshold: Threshold in seconds for penumbra definition
   Returns:
   penumbra_volume: Volume of tissue with Tmax > threshold
   # Load the perfusion map
   img = nib.load(perfusion_file)
   tmax data = img.get fdata()
   # Get voxel dimensions to calculate volume
   voxel_size = np.prod(img.header.get_zooms())
   # Threshold the image to identify penumbra
   penumbra mask = tmax data > t max threshold
   # Calculate penumbra volume in ml
   penumbra_volume = np.sum(penumbra_mask) * voxel_size / 1000.0
   # Get a middle slice for visualization
   slice_idx = tmax_data.shape[2] // 2
   # Visualize the perfusion map and penumbra
   fig, (ax1, ax2) = plt.subplots(1, 2, figsize=(15, 6))
   # Perfusion map
   im1 = ax1.imshow(tmax data[:, :, slice idx], cmap='jet')
   ax1.set_title('Tmax Perfusion Map')
   ax1.axis('off')
   plt.colorbar(im1, ax=ax1, label='Tmax (s)')
   # Penumbra overlay
   ax2.imshow(tmax_data[:, :, slice_idx], cmap='gray')
   penumbra_overlay = np.ma.masked_where(
       ~penumbra mask[:, :, slice idx],
       np.ones_like(tmax_data[:, :, slice_idx])
   ax2.imshow(penumbra_overlay, cmap='autumn', alpha=0.7)
    ax2.set_title(f'Penumbra (Tmax > {t_max_threshold}s)')
   ax2.axis('off')
   plt.tight_layout()
```

```
plt.show()
    print(f"Estimated penumbra volume: {penumbra volume:.2f} ml")
    return penumbra volume
# Example of decision support for stroke treatment
def stroke treatment decision(penumbra volume, core volume, onset time):
    Simple decision support for stroke treatment based on imaging and clinical fa
    Parameters:
    penumbra volume: Volume of salvageable tissue (ml)
    core volume: Volume of infarcted core (ml)
    onset_time: Time since stroke onset (hours)
    Returns:
    recommendation: Treatment recommendation
    mismatch_ratio = penumbra_volume / core_volume if core_volume > 0 else float(
    # Decision logic (simplified for demonstration)
    if onset time <= 4.5:</pre>
        if core volume < 70:
            recommendation = "Consider IV thrombolysis"
        else:
            recommendation = "Large core infarct. Consider mechanical thrombectom
    elif onset time <= 24:</pre>
        if mismatch_ratio >= 1.8 and penumbra_volume >= 15:
            recommendation = "Consider extended window mechanical thrombectomy"
        else:
            recommendation = "Insufficient penumbra for endovascular treatment"
        recommendation = "Outside treatment window"
    # Visualization
    plt.figure(figsize=(10, 6))
    plt.bar(['Core', 'Penumbra'], [core_volume, penumbra_volume], color=['red', '
    plt.axhline(y=70, linestyle='--', color='black', label='Core Volume Threshold
    plt.ylabel('Volume (ml)')
    plt.title(f'Stroke Tissue Volumes\nMismatch Ratio: {mismatch ratio:.1f}, Time
    plt.leaend()
    plt.show()
    print(f"Treatment recommendation: {recommendation}")
    return recommendation
# Usage example (pseudocode)
# penumbra volume = analyze stroke perfusion('/path/to/tmax.nii.gz')
# stroke_treatment_decision(penumbra_volume=45, core_volume=15, onset_time=3.5)
```

# Multiple Sclerosis

## Multiple Sclerosis Lesion Segmentation Challenge Dataset

- Description: Public dataset for evaluating MS lesion segmentation algorithms
- Contents: MRI scans with manual lesion segmentations from multiple centers
- Access: https://smart-stats-tools.org/lesion-challenge
- **Documentation**: https://smart-stats-tools.org/lesion-challenge-dataset

```
# Example: MS lesion segmentation
import nibabel as nib
import numpy as np
import matplotlib.pyplot as plt
from skimage import measure, filters
def segment ms lesions(flair file, t1 file, t2 file, threshold factor=2.5):
    Simple MS lesion segmentation using FLAIR hyperintensities.
    Parameters:
    flair file: Path to FLAIR MRI
    t1 file: Path to T1 MRI
    t2 file: Path to T2 MRI
    threshold_factor: Multiplier for intensity threshold
    Returns:
    lesion_mask: Binary mask of detected lesions
    # Load MRI sequences
    flair_img = nib.load(flair_file)
    flair data = flair img.get fdata()
    t1_data = nib.load(t1_file).get_fdata()
    t2_data = nib.load(t2_file).get_fdata()
    # Simple lesion segmentation based on FLAIR hyperintensities
    # Real segmentation would use more sophisticated methods
    # Calculate brain mask (simple thresholding)
    brain mask = t1 data > np.percentile(t1 data, 15)
    # Threshold FLAIR within brain mask
    flair brain = flair data * brain mask
    flair threshold = np.mean(flair brain[flair brain > 0]) + threshold factor *
    # Initial lesion segmentation
    lesion mask = (flair data > flair threshold) & brain mask
    # Remove small lesions (noise)
    # Label connected components
    labels = measure.label(lesion mask)
    # Calculate properties for each lesion candidate
    props = measure.regionprops(labels)
   # Filter by size (remove very small objects)
   min size = 10 # voxels
    lesion_mask = np.zeros_like(lesion_mask)
    for prop in props:
        if prop.area >= min size:
            lesion_mask[labels == prop.label] = 1
```

```
# Visualize results (middle slice)
    slice idx = flair data.shape[2] // 2
    fig, axes = plt.subplots(2, 2, figsize=(15, 12))
    # FLAIR image
    axes[0, 0].imshow(flair_data[:, :, slice_idx], cmap='gray')
    axes[0, 0].set_title('FLAIR Image')
    axes[0, 0].axis('off')
    # T1 image
    axes[0, 1].imshow(t1 data[:, :, slice idx], cmap='gray')
    axes[0, 1].set title('T1 Image')
    axes[0, 1].axis('off')
   # T2 image
    axes[1, 0].imshow(t2 data[:, :, slice idx], cmap='gray')
    axes[1, 0].set_title('T2 Image')
    axes[1, 0].axis('off')
    # FLAIR with lesion overlay
    axes[1, 1].imshow(flair_data[:, :, slice_idx], cmap='gray')
    lesion overlay = np.ma.masked where(
        ~lesion_mask[:, :, slice_idx],
        np.ones like(flair data[:, :, slice idx])
    )
    axes[1, 1].imshow(lesion overlay, cmap='hot', alpha=0.7)
    axes[1, 1].set_title('Lesion Segmentation')
    axes[1, 1].axis('off')
    plt.tight_layout()
   plt.show()
    # Calculate total lesion volume
    voxel size = np.prod(flair img.header.get zooms())
    lesion volume = np.sum(lesion mask) * voxel size / 1000.0 # ml
    print(f"Estimated MS lesion volume: {lesion volume:.2f} ml")
    return lesion mask, lesion volume
# Example of lesion load progression analysis
def analyze lesion progression(lesion volumes, scan dates):
    """Analyze progression of MS lesion volumes over time."""
    # Convert dates to numeric (days since first scan)
    from datetime import datetime
    dates = [datetime.strptime(d, '%Y-%m-%d') for d in scan_dates]
    days = [(d - dates[0]).days for d in dates]
    # Visualize progression
    plt.figure(figsize=(10, 6))
    plt.plot(days, lesion_volumes, 'o-', linewidth=2)
    plt.xlabel('Days from Baseline')
```

```
plt.ylabel('Lesion Volume (ml)')
    plt.title('MS Lesion Volume Progression')
    plt.grid(True)
    # Calculate rate of change
    if len(lesion volumes) > 1:
        annual_change = 365.0 * (lesion_volumes[-1] - lesion_volumes[0]) / days[-
        plt.axline((0, lesion_volumes[0]), slope=annual_change/365.0,
                  color='r', linestyle='--', label=f'Trend: {annual_change:.2f} m
        plt.legend()
    plt.show()
# Usage example (pseudocode)
# lesion_mask, volume = segment_ms_lesions(
      '/path/to/flair.nii.gz',
      '/path/to/t1.nii.gz',
      '/path/to/t2.nii.gz'
# )
# Lesion progression analysis example
# analyze_lesion_progression(
     [10.5, 12.3, 15.7, 14.9, 18.2],
      ['2020-01-15', '2020-07-20', '2021-01-10', '2021-07-05', '2022-01-12']
# )
```

## Other Neurological Disorders

### Brain Tumor Segmentation (BraTS) Challenge

- Description: Multimodal brain tumor MRI dataset
- Contents: T1, T1ce, T2, and FLAIR MRIs with expert segmentations
- Access: <a href="http://braintumorsegmentation.org/">http://braintumorsegmentation.org/</a>
- **Documentation**: https://www.med.upenn.edu/cbica/brats2020/data.html

```
# Example: Brain tumor visualization and segmentation
import nibabel as nib
import numpy as np
import matplotlib.pyplot as plt
def visualize_brain_tumor(t1_file, t1ce_file, t2_file, flair_file, seg_file=None)
    Visualize multimodal brain tumor MRI data.
    Parameters:
    t1 file: Path to T1 MRI
    t1ce file: Path to T1 contrast-enhanced MRI
    t2 file: Path to T2 MRI
    flair_file: Path to FLAIR MRI
    seg_file: Path to segmentation (optional)
    # Load MRI data
    t1 = nib.load(t1_file).get_fdata()
    t1ce = nib.load(t1ce_file).get_fdata()
    t2 = nib.load(t2 file).get fdata()
    flair = nib.load(flair_file).get_fdata()
    # Load segmentation if available
    if seg_file:
        seg = nib.load(seg_file).get_fdata()
        has_seg = True
    else:
        has seg = False
    # Get the middle slice for each dimension
    axial slice = t1.shape[2] // 2
    # Create a figure
    fig, axes = plt.subplots(2, 3 if has_seg else 2, figsize=(15, 10))
    # Display T1
    axes[0, 0].imshow(t1[:, :, axial_slice], cmap='gray')
    axes[0, 0].set_title('T1')
    axes[0, 0].axis('off')
    # Display T1ce
    axes[0, 1].imshow(t1ce[:, :, axial_slice], cmap='gray')
    axes[0, 1].set title('T1ce')
    axes[0, 1].axis('off')
    # Display T2
    axes[1, 0].imshow(t2[:, :, axial_slice], cmap='gray')
    axes[1, 0].set_title('T2')
    axes[1, 0].axis('off')
    # Display FLAIR
    axes[1, 1].imshow(flair[:, :, axial_slice], cmap='gray')
    axes[1, 1].set_title('FLAIR')
```

```
axes[1, 1].axis('off')
    # Display segmentation if available
    if has seg:
        axes[0, 2].imshow(t1ce[:, :, axial slice], cmap='gray')
        # Create a colored overlay for different tumor components
        # In BraTS: 1=necrotic core, 2=edema, 4=enhancing tumor
        seg slice = seg[:, :, axial slice]
        # Necrotic core (red)
        necrotic = np.ma.masked where(seg slice != 1, np.ones like(seg slice))
        axes[0, 2].imshow(necrotic, cmap='Reds', alpha=0.7)
        # Edema (green)
        edema = np.ma.masked where(seg slice != 2, np.ones like(seg slice))
        axes[0, 2].imshow(edema, cmap='Greens', alpha=0.7)
        # Enhancing tumor (blue)
        enhancing = np.ma.masked_where(seg_slice != 4, np.ones_like(seg_slice))
        axes[0, 2].imshow(enhancing, cmap='Blues', alpha=0.7)
        axes[0, 2].set_title('Tumor Segmentation')
        axes[0, 2].axis('off')
        # Calculate tumor volumes
        voxel_size = 1.0 # mm³, adjust based on actual voxel size
        necrotic vol = np.sum(seg == 1) * voxel size / 1000.0 # ml
        edema_vol = np.sum(seg == 2) * voxel_size / 1000.0 # ml
        enhancing vol = np.sum(seq == 4) * voxel size / 1000.0 # ml
        total vol = necrotic vol + edema vol + enhancing vol
        # Display volumes
        text = (f"Tumor Components:\n"
                f"Necrotic core: {necrotic vol:.2f} ml\n"
                f"Edema: {edema vol:.2f} ml\n"
                f"Enhancing tumor: {enhancing_vol:.2f} ml\n"
                f"Total volume: {total vol:.2f} ml")
        axes[1, 2].axis('off')
        axes[1, 2].text(0.1, 0.5, text, fontsize=12)
    plt.tight_layout()
    plt.show()
# Usage example (pseudocode)
# visualize brain tumor(
      '/path/to/t1.nii.gz',
#
      '/path/to/t1ce.nii.gz',
      '/path/to/t2.nii.gz',
#
#
      '/path/to/flair.nii.gz',
      '/path/to/seg.nii.gz'
#
# )
```

## Sleep-EDF Database

- Description: Sleep recordings with annotations for normal and abnormal sleep
- Contents: PSG and EEG recordings with sleep stage annotations
- Access: <a href="https://physionet.org/content/sleep-edfx/1.0.0/">https://physionet.org/content/sleep-edfx/1.0.0/</a>
- **Documentation**: https://physionet.org/content/sleep-edfx/1.0.0/

```
# Example: Sleep stage classification from EEG
import numpy as np
import matplotlib.pyplot as plt
import pyedflib
from scipy import signal
def analyze sleep edf(edf file, hypnogram file):
   Analyze Sleep-EDF data with sleep stage annotations.
   Parameters:
   edf file: Path to EEG recording
   hypnogram_file: Path to sleep stage annotations
   # Load EEG data
   try:
       f = pyedflib.EdfReader(edf file)
       # Get signal information
       n channels = f.signals in file
       channel_names = f.getSignalLabels()
       # Find EEG channels (typically Fpz-Cz and Pz-Oz in Sleep-EDF)
       eeg_channels = [i for i, name in enumerate(channel_names) if 'EEG' in nam
       if not eeg_channels:
            print("No EEG channels found")
            return
        # Read EEG data from first EEG channel
       eeg_channel = eeg_channels[0]
       fs = f.getSampleFrequency(eeg channel)
        eeg_signal = f.readSignal(eeg_channel)
       # Load hypnogram (sleep stages)
       with open(hypnogram file, 'r') as hyp file:
            hypnogram = hyp_file.readlines()
       # Parse sleep stages (simplified, actual parsing depends on hypnogram for
       # In Sleep-EDF annotations: 0=Wake, 1=REM, 2=N1, 3=N2, 4=N3/N4
        sleep stages = []
       for line in hypnogram:
            if line.strip() and line[0].isdigit():
                sleep stages.append(int(line[0]))
       sleep_stages = np.array(sleep_stages)
       # Calculate time vectors
       eeg_time = np.arange(len(eeg_signal)) / fs # seconds
        stage\_time = np.arange(len(sleep\_stages)) * 30 # 30-second epochs
       # Extract features for visualization
       # (in real application, would extract features for each epoch)
```

```
# Power spectral density (whole signal)
f, psd = signal.welch(eeg signal, fs, nperseg=fs*4)
# Find frequency bands
delta mask = (f >= 0.5) & (f <= 4)
theta_mask = (f >= 4) & (f <= 8)
alpha mask = (f >= 8) & (f <= 13)
beta mask = (f >= 13) & (f <= 30)
# Visualize data
fig, axes = plt.subplots(3, 1, figsize=(15, 12))
# Plot EEG signal (first 60 seconds)
plot_duration = min(60, len(eeg_signal) / fs)
plot samples = int(plot duration * fs)
axes[0].plot(eeg time[:plot samples], eeg signal[:plot samples])
axes[0].set_xlabel('Time (s)')
axes[0].set_ylabel('Amplitude')
axes[0].set_title(f'EEG Signal ({channel_names[eeg_channel]})')
# Plot sleep stages
stage_labels = ['Wake', 'REM', 'N1', 'N2', 'N3/N4']
axes[1].step(stage time / 3600, sleep stages, where='post')
axes[1].set_yticks(range(5))
axes[1].set yticklabels(stage labels)
axes[1].set_xlabel('Time (hours)')
axes[1].set_ylabel('Sleep Stage')
axes[1].set_title('Hypnogram')
axes[1].grid(True)
# Plot power spectral density
axes[2].semilogy(f, psd)
axes[2].set xlabel('Frequency (Hz)')
axes[2].set_ylabel('PSD (\mu V^2/Hz)')
axes[2].set_title('Power Spectral Density')
# Highlight frequency bands
axes[2].fill_between(f[delta_mask], 0, psd[delta_mask], alpha=0.3, label=
axes[2].fill_between(f[theta_mask], 0, psd[theta_mask], alpha=0.3, label=
axes[2].fill_between(f[alpha_mask], 0, psd[alpha_mask], alpha=0.3, label=
axes[2].fill between(f[beta mask], 0, psd[beta mask], alpha=0.3, label='B
axes[2].legend()
plt.tight_layout()
plt.show()
# Calculate sleep metrics
total_epochs = len(sleep_stages)
total sleep time = sum(sleep stages != 0) * 30 / 60 # minutes
# Sleep efficiency (total sleep time / time in bed)
```

```
sleep_efficiency = sum(sleep_stages != 0) / total_epochs * 100
        # Time in each stage
        stage minutes = {
            stage_labels[i]: sum(sleep_stages == i) * 30 / 60
            for i in range(5)
        }
        # Print sleep metrics
        print(f"Sleep Metrics:")
        print(f"Total recording time: {total epochs * 30 / 60:.1f} minutes")
        print(f"Total sleep time: {total sleep time:.1f} minutes")
        print(f"Sleep efficiency: {sleep efficiency:.1f}%")
        print("Time in each stage:")
        for stage, minutes in stage_minutes.items():
            print(f" {stage}: {minutes:.1f} minutes ({minutes/total_sleep_time*1
        f.close()
    except Exception as e:
        print(f"Error analyzing Sleep-EDF data: {e}")
# Usage example (pseudocode)
# analyze_sleep_edf('/path/to/sleep_edf.edf', '/path/to/hypnogram.txt')
```

These neurological and neurodegenerative disease datasets provide valuable resources for developing and validating AI systems for diagnosis, treatment planning, and monitoring disease progression. The examples demonstrate basic approaches to working with these diverse data types and highlight the potential for AI applications in clinical neurology.