data loading

```
import os
import shutil
import random
import numpy as np
import pandas as pd
import cv2
import matplotlib.pyplot as plt
from google.colab import drive
try:
   drive.mount('/content/drive')
   print("Google Drive mounted successfully!")
printpt:
   prprintint("Running locally or Drive already mounted")
lgg_dataset_path = '/content/drive/MyDrive/ddd'
output_dir = '/content/drive/MyDrive/brain_tumor_dataset'
os.makedirs(os.path.join(output_dir, 'images'), exist_ok=True)
os.makedirs(os.path.join(output_dir, 'masks'), exist_ok=True)
def prepare_dataset(num_samples=10, random_selection=True):
   Prepare a subset of the LGG dataset for segmentation
   Parameters:
    -----
   num_samples : int
        Number of image-mask pairs to extract
   random_selection : bool
       Whether to select random samples or the first ones
   case_dirs = [d for d in os.listdir(lgg_dataset_path)
                if os.path.isdir(os.path.join(lgg_dataset_path, d))]
   print(f"Found {len(case_dirs)} case directories")
   if random_selection:
        selected_cases = random.sample(case_dirs, min(len(case_dirs), num_samples))
   else:
        selected_cases = case_dirs[:min(len(case_dirs), num_samples)]
   print(f"Selected {len(selected_cases)} cases")
   processed count = 0
    for case_id in selected_cases:
        case_dir = os.path.join(lgg_dataset_path, case_id)
        files = os.listdir(case_dir)
        image_files = [f for f in files if f.endswith('.tif') and '_mask' not in f]
        for img_file in image_files:
           mask_file = img_file.replace('.tif', '_mask.tif')
            if mask_file in files:
```

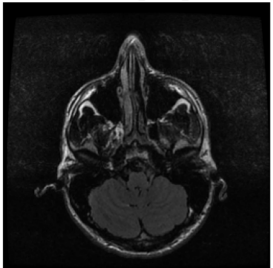
```
img_path = os.path.join(case_dir, img_file)
               mask path = os.path.join(case dir, mask file)
               img = cv2.imread(img_path)
               mask = cv2.imread(mask_path, cv2.IMREAD_GRAYSCALE)
               if img is not None and mask is not None:
                    img = cv2.cvtColor(img, cv2.COLOR_BGR2RGB)
                    flair = img[:, :, 1]
                    out_img_name = f"brain_tumor_{processed_count:03d}.png"
                    out_mask_name = f"brain_tumor_{processed_count:03d}_mask.png"
                    out_img_path = os.path.join(output_dir, 'images', out_img_name)
                    out mask path = os.path.join(output dir, 'masks', out mask name)
                    cv2.imwrite(out_img_path, flair)
                    cv2.imwrite(out_mask_path, mask)
                    processed_count += 1
                    print(f"Processed {processed_count}/{num_samples}: {out_img_name}")
                    if processed count >= num samples:
                        return
def visualize_dataset(dataset_path, num_samples=5):
   Visualize the prepared dataset
   Parameters:
    -----
   dataset_path : str
       Path to the prepared dataset
   num_samples : int
       Number of samples to visualize
    images_dir = os.path.join(dataset_path, 'images')
   masks_dir = os.path.join(dataset_path, 'masks')
    image files = sorted(os.listdir(images dir))
    image_files = image_files[:min(len(image_files), num_samples)]
   plt.figure(figsize=(12, 4 * len(image_files)))
   for i, img_file in enumerate(image_files):
        if img_file.replace('.png', '_mask.png') in os.listdir(masks_dir):
           mask_file = img_file.replace('.png', '_mask.png')
        else:
           mask_file = next((m for m in os.listdir(masks_dir) if m.startswith(img_file.split('.')[")
        if mack file.
```

```
II Mask_IIIe.
            img_path = os.path.join(images_dir, img_file)
            mask_path = os.path.join(masks_dir, mask_file)
            img = cv2.imread(img_path, cv2.IMREAD_GRAYSCALE)
            mask = cv2.imread(mask_path, cv2.IMREAD_GRAYSCALE)
            plt.subplot(len(image_files), 2, i * 2 + 1)
            plt.imshow(img, cmap='gray')
            plt.title(f"MRI Image: {img_file}")
            plt.axis('off')
            plt.subplot(len(image_files), 2, i * 2 + 2)
            plt.imshow(mask, cmap='gray')
            plt.title(f"Tumor Mask: {mask_file}")
            plt.axis('off')
    plt.tight_layout()
   plt.show()
print("Preparing dataset...")
prepare_dataset(num_samples=10, random_selection=True)
print("Dataset preparation completed!")
print("Visualizing prepared dataset...")
visualize_dataset(output_dir, num_samples=5)
print("\nDataset is ready to use with the brain tumor segmentation code!")
print(f"Images directory: {os.path.join(output_dir, 'images')}")
print(f"Masks directory: {os.path.join(output_dir, 'masks')}")
print("\nUpdate these paths in the main code:")
print("base_dir = '", output_dir, "'")
```

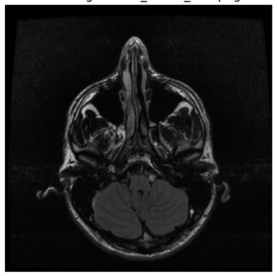
print("images_dir = os.path.join(base_dir, 'images')")
print("masks dir = os.path.join(base dir, 'masks')")

Google Drive mounted successfully! Preparing dataset... Found 1 case directories Selected 1 cases Dataset preparation completed! Visualizing prepared dataset...

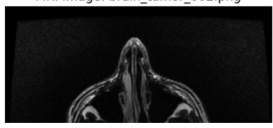
MRI Image: brain_tumor_000.png



MRI Image: brain_tumor_001.png



MRI Image: brain_tumor_002.png

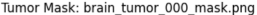


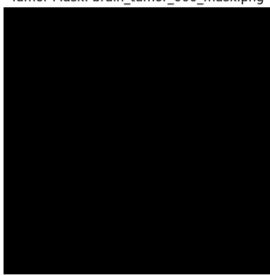
from google.colab import drive drive.mount('/content/drive')



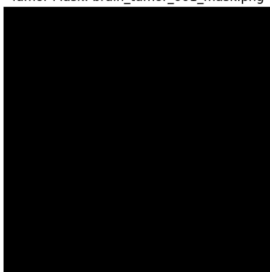
to attempt to



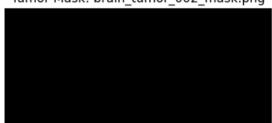




Tumor Mask: brain tumor 001 mask.png



Tumor Mask: brain_tumor_002_mask.png



import os import cv2 import numpy as np import matplotlib.pyplot as plt from google.colab import drive from sklearn.model_selection import train_test_split

from sklearn.metrics import accuracy_score, jaccard_score, f1_score, precision_score, recall_scor

```
try:
   drive.mount('/content/drive')
   print("Google Drive mounted successfully!")
   print("Running locally or Drive already mounted")
class BrainTumorSegmentation:
   def __init__(self, base_dir=None):
        Initialize the Brain Tumor Segmentation class
        Parameters:
        _____
        base_dir : str
           Base directory containing the dataset
        self.base_dir = base_dir
        self.images = []
        self.masks = []
        self.processed_images = []
        self.segmented_masks = []
       self.metrics = {}
   def load_dataset(self, images_dir, masks_dir, max_samples=None):
        Load dataset images and their corresponding masks
        Parameters:
        -----
        images_dir : str
           Directory containing MRI scan images
        masks_dir : str
           Directory containing corresponding mask images
        max_samples : int, optional
           Maximum number of samples to load
        print("Loading dataset...")
        image_files = sorted(os.listdir(images_dir))
        loaded count = 0
        for img_file in image_files:
            if not img_file.endswith(('.jpg', '.png', '.jpeg', '.tif')):
               continue
            if '_mask' not in img_file:
               mask_file = img_file.replace('.png', '_mask.png')
               mask_file = mask_file.replace('.tif', '_mask.tif')
               mask_file = mask_file.replace('.jpg', '_mask.jpg')
                image_path = os.path.join(images_dir, img_file)
               mask_path = os.path.join(masks_dir, mask_file)
               if not os.path.exists(mask_path):
                    print(f"Warning: No mask found for {img_file}")
                    continue
```

```
mask = cv2.imread(mask path, cv2.IMREAD GRAYSCALE)
           if image is not None and mask is not None:
                if len(image.shape) == 3 and image.shape[2] == 3:
                    gray_image = image[:, :, 1]
                else:
                    gray_image = image.copy()
                if len(gray_image.shape) == 3:
                    gray_image = cv2.cvtColor(gray_image, cv2.COLOR_BGR2GRAY)
                _, binary_mask = cv2.threshold(mask, 127, 255, cv2.THRESH_BINARY)
                self.images.append(gray_image)
                self.masks.append(binary_mask)
                loaded count += 1
                if max_samples is not None and loaded_count >= max_samples:
                    break
    print(f"Loaded {len(self.images)} images and {len(self.masks)} masks.")
def preprocess_images(self):
    """Preprocess the loaded MRI images"""
    print("Preprocessing images...")
    self.processed_images = []
    for image in self.images:
        if len(image.shape) > 2:
           gray image = cv2.cvtColor(image, cv2.COLOR RGB2GRAY)
        else:
           gray_image = image.copy()
        if gray_image.max() > 0:
           normalized = ((gray_image - gray_image.min()) /
                         (gray_image.max() - gray_image.min()) * 255).astype(np.uint8)
        else:
           normalized = gray_image
        clahe = cv2.createCLAHE(clipLimit=2.0, tileGridSize=(8, 8))
        enhanced = clahe.apply(normalized)
       blurred = cv2.GaussianBlur(enhanced, (5, 5), 0)
        self.processed_images.append(blurred)
    print(f"Preprocessed {len(self.processed_images)} images.")
def segment_tumors(self, method='watershed'):
    Segment tumor regions from preprocessed images
    Parameters:
    ------
```

image = cv2.imread(image_path)

```
method : str
   Segmentation method to use ('threshold', 'watershed', 'kmeans')
print(f"Segmenting tumors using {method} method...")
self.segmented_masks = []
for image in self.processed_images:
    if method == 'threshold':
        _, segmented = cv2.threshold(image, 0, 255, cv2.THRESH_BINARY + cv2.THRESH OTSU)
   elif method == 'watershed':
        _, thresholded = cv2.threshold(image, 0, 255, cv2.THRESH_BINARY + cv2.THRESH_OTSU
       kernel = np.ones((3, 3), np.uint8)
       opening = cv2.morphologyEx(thresholded, cv2.MORPH_OPEN, kernel, iterations=2)
       sure_bg = cv2.dilate(opening, kernel, iterations=3)
       dist transform = cv2.distanceTransform(opening, cv2.DIST L2, 5)
        _, sure_fg = cv2.threshold(dist_transform, 0.7*dist_transform.max(), 255, 0)
       sure_fg = np.uint8(sure_fg)
       unknown = cv2.subtract(sure_bg, sure_fg)
       _, markers = cv2.connectedComponents(sure_fg)
       markers = markers + 1
       markers[unknown == 255] = 0
       markers = cv2.watershed(cv2.cvtColor(image, cv2.COLOR_GRAY2BGR), markers)
        segmented = np.zeros_like(image)
        segmented[markers > 1] = 255
    elif method == 'kmeans':
        image_data = image.reshape((-1, 1))
        image_data = np.float32(image_data)
       criteria = (cv2.TERM_CRITERIA_EPS + cv2.TERM_CRITERIA_MAX_ITER, 100, 0.2)
       k = 3
        _, labels, centers = cv2.kmeans(image_data, k, None, criteria, 10, cv2.KMEANS_RAN
       centers = np.uint8(centers)
       brightest_cluster = np.argmax(centers)
        segmented = np.zeros_like(image)
       segmented[labels.reshape(image.shape) == brightest_cluster] = 255
    else:
        raise ValueError(f"Unknown segmentation method: {method}")
```

```
segmented = self.post_process_mask(segmented)
        self.segmented_masks.append(segmented)
    print(f"Segmented {len(self.segmented_masks)} images.")
def post_process_mask(self, mask):
    Apply post-processing to improve the segmentation mask
   Parameters:
    -----
    mask : numpy.ndarray
       Binary segmentation mask
    Returns:
    _ _ _ _ _ _ _
    numpy.ndarray
       Post-processed binary mask
    if mask.dtype != np.uint8:
       mask = mask.astype(np.uint8)
    contours, _ = cv2.findContours(mask, cv2.RETR_EXTERNAL, cv2.CHAIN_APPROX_SIMPLE)
    filled_mask = np.zeros_like(mask)
    cv2.drawContours(filled_mask, contours, -1, 255, -1)
    nb_components, output, stats, _ = cv2.connectedComponentsWithStats(filled_mask, connectiv
    sizes = stats[1:, -1]
    min size = 100
    processed mask = np.zeros like(filled mask)
    for i in range(1, nb_components):
        if sizes[i - 1] >= min_size:
           processed_mask[output == i] = 255
    return processed_mask
def evaluate_segmentation(self, ground_truth_masks=None):
    Evaluate segmentation performance against ground truth masks
    Parameters:
    -----
    ground_truth_masks : list
       List of ground truth mask images
    Returns:
    dict
       Dictionary containing evaluation metrics
    if ground_truth_masks is None:
       ground_truth_masks = self.masks
    if len(ground_truth_masks) != len(self.segmented_masks):
        raise ValueError("Mismatch between number of ground truth masks and segmented masks")
```

dice scores = []

```
jaccard_scores = []
   precision scores = []
   recall scores = []
   for gt_mask, pred_mask in zip(ground_truth_masks, self.segmented_masks):
       gt_binary = np.where(gt_mask > 0, 1, 0).flatten()
       pred_binary = np.where(pred_mask > 0, 1, 0).flatten()
       dice = f1_score(gt_binary, pred_binary, zero_division=1)
       dice_scores.append(dice)
       iou = jaccard_score(gt_binary, pred_binary, zero_division=1)
       jaccard_scores.append(iou)
       precision = precision_score(gt_binary, pred_binary, zero_division=1)
       recall = recall_score(gt_binary, pred_binary, zero_division=1)
       precision scores.append(precision)
       recall_scores.append(recall)
   self.metrics = {
        'dice_coefficient': np.mean(dice_scores),
        'jaccard_index': np.mean(jaccard_scores),
        'precision': np.mean(precision_scores),
        'recall': np.mean(recall_scores)
   }
   print("Segmentation Evaluation Metrics:")
   print(f" Dice Coefficient (F1-Score): {self.metrics['dice_coefficient']:.4f}")
   print(f" Jaccard Index (IoU): {self.metrics['jaccard_index']:.4f}")
   print(f" Precision: {self.metrics['precision']:.4f}")
   print(f" Recall: {self.metrics['recall']:.4f}")
   return self.metrics
def visualize_results(self, num_samples=5):
   Visualize original images, ground truth masks, and segmented masks
   Parameters:
   num_samples : int
       Number of samples to visualize
   num_samples = min(num_samples, len(self.images))
   plt.figure(figsize=(15, 4 * num_samples))
   for i in range(num_samples):
       plt.subplot(num_samples, 3, i * 3 + 1)
       plt.imshow(self.images[i], cmap='gray')
       plt.title(f"Original Image {i+1}")
       plt.axis('off')
       plt.subplot(num_samples, 3, i * 3 + 2)
       plt.imshow(self.masks[i], cmap='gray')
       plt.title(f"Ground Truth Mask {i+1}")
       plt.axis('off')
```

```
plt.subplot(num_samples, 3, i * 3 + 3)
        plt.imshow(self.segmented_masks[i], cmap='gray')
        plt.title(f"Segmented Mask {i+1}")
        plt.axis('off')
    plt.tight_layout()
    plt.show()
def overlay_results(self, num_samples=5):
    Overlay segmentation results on original images for visualization
    Parameters:
    -----
    num_samples : int
       Number of samples to visualize
    num_samples = min(num_samples, len(self.images))
    plt.figure(figsize=(12, 4 * num_samples))
    for i in range(num_samples):
        display_img = cv2.cvtColor(self.images[i], cv2.COLOR_GRAY2RGB)
        plt.subplot(num_samples, 2, i * 2 + 1)
        overlay = display_img.copy()
        green_mask = np.zeros_like(overlay)
        green_mask[:, :, 1] = self.masks[i]
        alpha = 0.5
        cv2.addWeighted(green_mask, alpha, overlay, 1 - alpha, 0, overlay)
        plt.imshow(overlay)
        plt.title(f"Original + Ground Truth {i+1}")
        plt.axis('off')
        plt.subplot(num_samples, 2, i * 2 + 2)
        overlay = display_img.copy()
        red_mask = np.zeros_like(overlay)
        red_mask[:, :, 0] = self.segmented_masks[i]
        cv2.addWeighted(red_mask, alpha, overlay, 1 - alpha, 0, overlay)
        plt.imshow(overlay)
        plt.title(f"Original + Segmentation {i+1}")
        plt.axis('off')
    plt.tight_layout()
    plt.show()
```

```
def run_full_pipeline(self, images_dir, masks_dir, max_samples=None, segmentation_method='wat
        Run the full segmentation pipeline
        Parameters:
        images_dir : str
           Directory containing MRI scan images
        masks_dir : str
            Directory containing corresponding mask images
        max_samples : int, optional
            Maximum number of samples to load
        segmentation_method : str
           Method to use for segmentation ('threshold', 'watershed', 'kmeans')
        self.load_dataset(images_dir, masks_dir, max_samples)
        self.preprocess_images()
        self.segment_tumors(method=segmentation_method)
        self.evaluate segmentation()
        self.visualize_results()
        self.overlay_results()
        return self.metrics
if __name__ == "__main__":
   base dir = '/content/drive/MyDrive/brain tumor dataset'
   images_dir = os.path.join(base_dir, 'images')
   masks_dir = os.path.join(base_dir, 'masks')
   tumor_segmentation = BrainTumorSegmentation(base_dir)
   methods = ['threshold', 'watershed', 'kmeans']
   results = {}
    for method in methods:
        print(f"\n{'-'*50}")
        print(f"Running segmentation with {method.upper()} method")
       print(f"{'-'*50}")
        metrics = tumor_segmentation.run_full_pipeline(
            images dir=images dir,
            masks_dir=masks_dir,
            max_samples=10,
            segmentation_method=method
        )
        results[method] = metrics
   print("\nComparison of Segmentation Methods:")
   print(f"{'Method':<12} {'Dice':<8} {'IoU':<8} {'Precision':<10} {'Recall':<8}")</pre>
   print("-" * 50)
```

Running segmentation with THRESHOLD method

Loading dataset...

Loaded 7 images and 7 masks.

Preprocessing images... Preprocessed 7 images.

Segmenting tumors using threshold method...

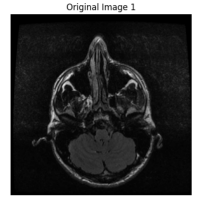
Segmented 7 images.

Segmentation Evaluation Metrics:

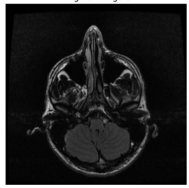
Dice Coefficient (F1-Score): 0.0000

Jaccard Index (IoU): 0.0000

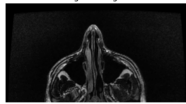
Precision: 0.0000 Recall: 1.0000



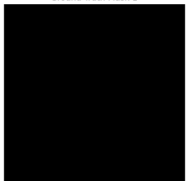




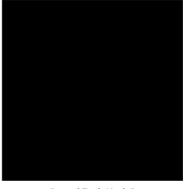
Original Image 3



Ground Truth Mask 1



Ground Truth Mask 2



Ground Truth Mask 3



Segmented Mask 1



Segmented Mask 2



Segmented Mask 3

