## 2D Glioma classification

This notebook demonstrates classification of brain tumors with MONAI. To accelerate training, 2D dataset is used based on the 3D one.

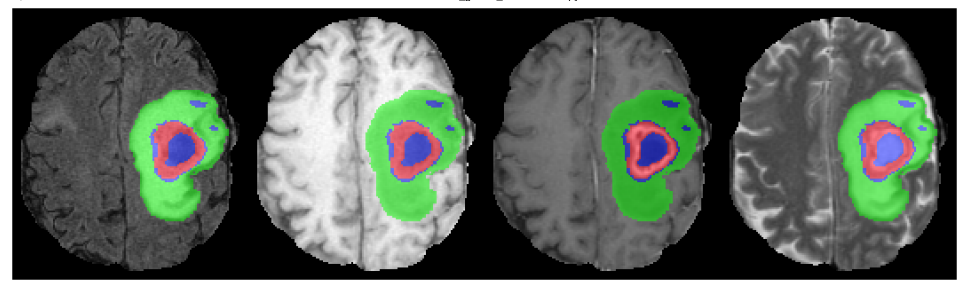
### Notebook structure

Here's a rough outline of this notebook:

- 1. Check MONAI is installed and install if not (plus any extra dependencies)
- 2. Import libraries
- 3. Set up the data (download or create)
- 4. Prepare for training -- create transforms, datasets, dataloaders
- 5. Do some quick visualisations
- 6. Create our model, loss function, etc.
- 7. Train 🞉 😚 👭
- 8. Check results

### Dataset

The dataset used here is the Decathlon 3D brain tumor dataset. We generate a 2D dataset by combining slices containing tumour, and those containing no tumourous voxels (considered healthy). We'll download the pre-computed dataset from Google Drive, but the script is available in case you're interested.



# User input

Some sections have been left blank for you, the user, to complete. Suggested solutions are hidden in a collapsible text box. Feel free to rely on these as much as necessary, but you will get the most of the experience if you think it through or search the internet prior to displaying the results. Remember there are lots of ways of achieving the same goal, so it's OK if your solutions are different!

## **Extensions**

- 1. We have a small amount of acceleration thanks to the CacheDataset, can you think of any other ways to make this training faster?
- 2. What methods could we use to reduce overfitting? Think about transforms and our model.
- 3. Could you use a different model or loss function altogether? Does this give better results?



### 1. Check MONAL is installed.

This checks if MONAI is installed, and if not installs it (plus any optional extras that might be needed for this notebook).

## 2. Import libraries

```
# Copyright 2022 MONAI Consortium
# Licensed under the Apache License, Version 2.0 (the "License");
# you may not use this file except in compliance with the License.
# You may obtain a copy of the License at
      http://www.apache.org/licenses/LICENSE-2.0
# Unless required by applicable law or agreed to in writing, software
# distributed under the License is distributed on an "AS IS" BASIS,
# WITHOUT WARRANTIES OR CONDITIONS OF ANY KIND, either express or implied.
# See the License for the specific language governing permissions and
# limitations under the License.
from glob import glob
import matplotlib.pyplot as plt
import numpy as np
import os
import tempfile
import torch
from tqdm import tqdm, trange
```

from sklearn.metrics import confusion matrix, ConfusionMatrixDisplay import monai from monai.apps import download and extract from monai.data import CacheDataset, DataLoader, pad list data collate, partition dataset from monai.networks import eval mode import monai.transforms as mt from monai.utils import set\_determinism monai.config.print\_config() # Use GPU if possible, else CPU (not recommended) device = torch.device("cuda" if torch.cuda.is available() else "cpu") # Set deterministic training for reproducibility set\_determinism(seed=42) → MONAI version: 1.5.dev2446 Numpy version: 1.26.4 Pytorch version: 2.5.1+cu121 MONAI flags: HAS EXT = False, USE\_COMPILED = False, USE\_META\_DICT = False MONAI rev id: 218216250ce297265400abe56ee915898d75a2ec MONAI file : /usr/local/lib/python3.10/dist-packages/monai/\_\_init\_\_.py Optional dependencies: Pytorch Ignite version: NOT INSTALLED or UNKNOWN VERSION. ITK version: NOT INSTALLED or UNKNOWN VERSION. Nibabel version: 5.3.2 scikit-image version: 0.24.0 scipy version: 1.13.1 Pillow version: 11.0.0 Tensorboard version: 2.17.1 gdown version: 5.2.0 TorchVision version: 0.20.1+cu121 tqdm version: 4.66.6 1mdb version: NOT INSTALLED or UNKNOWN VERSION. psutil version: 5.9.5 pandas version: 2.2.2 einops version: 0.8.0 transformers version: 4.46.2 mlflow version: NOT INSTALLED or UNKNOWN VERSION.

```
pynrrd version: NOT INSTALLED or UNKNOWN VERSION.
    clearml version: NOT INSTALLED or UNKNOWN VERSION.

For details about installing the optional dependencies, please visit:
        https://docs.monai.io/en/latest/installation.html#installing-the-recommended-dependencies

# figure out if we're running in Google Colab. Set paths accordingly.
try:
    from google.colab import drive
    drive.mount('/content/drive', force_remount=True)
    best_model_folder = "/content/drive/MyDrive/saved_models/"
    os.makedirs(best_model_folder, exist_ok=True)

except:
    best_model_folder = ""

best_model_path = os.path.join(best_model_folder, "best_model_2d_glioma_classification.pth")
```

## 3. Setup data

Mounted at /content/drive

You can specify a directory with the MONAI\_DATA\_DIRECTORY environment variable.

This allows you to save results and reuse downloads.

If not specified a temporary directory will be used.

#### 2D data

We'll download the pre-computed dataset from Google Drive, but the script is available in case you're interested.

```
directory = os.environ.get("MONAI_DATA_DIRECTORY")
root_dir = tempfile.mkdtemp() if directory is None else os.path.expanduser(directory)
print(root_dir)

_____ /tmp/tmpp_jjvnbj
```

```
download_from_gdrive = True
task = "Task01 BrainTumour"
output dir = os.path.join(root dir, task + "2D")
if download_from_gdrive:
    resource = "https://drive.google.com/uc?id=1BB0S2PcY6yUR7TK-AeyCFoh6PyoJiH0E&export=download"
    md5 = "214a338a26778c84ddebca29822add56"
    compressed file = os.path.join(root dir, task + "2D.tar")
    download and extract(resource, compressed file, root dir, hash val=md5)
else:
    %run -i ../utils/2d slice creator.py --path {output dir} --download path {root dir} --task {task}
    pass
# get all the 2d images
images_healthy = glob(os.path.join(output_dir, "image_healthy", "*.nii.gz"))
images tumour = glob(os.path.join(output dir, "image tumour", "*.nii.gz"))
data_dicts = []
data_dicts += [{"image": i, "label": [1, 0]} for i in images_healthy]
data_dicts += [{"image": i, "label": [0, 1]} for i in images_tumour]
# shuffle the data and sort into training and validation
train_files, val_files = partition_dataset(data_dicts, ratios=(8, 2), shuffle=True)
print("total num files:", len(data_dicts))
print("num training files:", len(train files))
print("num validation files:", len(val files))
→ Downloading...
     From (original): https://drive.google.com/uc?id=1BB0S2PcY6yUR7TK-AeyCFoh6PyoJiH0E
     From (redirected): https://drive.google.com/uc?id=1BB0S2PcY6yUR7TK-AeyCFoh6PyoJiH0E&confirm=t&uuid=80e206a1-45c6-4e16-8
     To: /tmp/tmpqf12jxwn/Task01 BrainTumour2D.tar
     100%| 122M/122M [00:03<00:00, 33.8MB/s]2024-11-19 23:45:17,283 - INFO - Downloaded: /tmp/tmpp_jjvnbj/Task01_
     2024-11-19 23:45:17,513 - INFO - Verified 'Task01 BrainTumour2D.tar', md5: 214a338a26778c84ddebca29822add56.
     2024-11-19 23:45:17,514 - INFO - Writing into directory: /tmp/tmpp jjvnbj.
     total num files: 968
     num training files: 774
     num validation files: 194
```

# 4. Prepare for training -- create transforms, datasets, dataloaders

### **Transforms**

In terms of transforms, we first load both the image and its corresponding label. We then rotate by 90 degrees, crop out superfluous zeros around the edge of the images, scale the image between 0 and 1 and then convert from numpy to pytorch.

In MONAI, we have two forms of transforms -- those that act on arrays of data, and those that act on dictionaries of data. For classification tasks, both are acceptable. We've used dictionaries here (note the d suffix in our transform names) -- such that each image and its corresponding class are stored together.

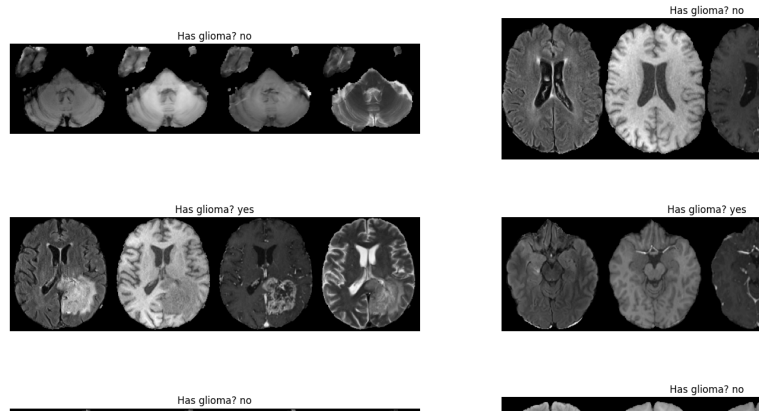
#### Datasets and dataloaders

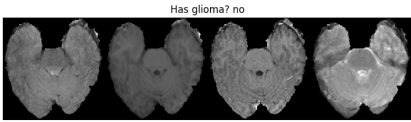
We use the CacheDataset which accelerates things by pre-computing the results of our deterministic transforms. We then use a batch size of 10 in our dataloader.

## 5. Display some examples

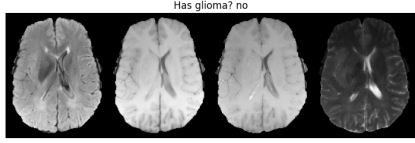
```
nrow, ncol = 5, 2
num_files = nrow * ncol
files_to_visualize = np.random.choice(train_files, size=num_files, replace=False)
fig, axes = plt.subplots(nrow, ncol, figsize=(20, 20), facecolor='white')
for f, ax in zip(files_to_visualize, axes.flatten()):
    data = train_transforms(f)
    # different modalities side by side
    img = np.concatenate(list(data["image"]), axis=1)
    im_show = ax.imshow(img, cmap="gray")
    ax.set_title(f"Has glioma? {['yes', 'no'][data['label'][0]]}")
    ax.axis("off")
```

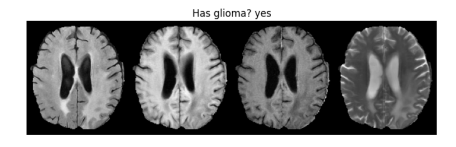
















6. Create the model, loss function, etc.

### The model

Which network do you want to use? How many channels should go in and how many should come out?

```
model = monai.networks.nets.DenseNet121(spatial_dims=2, in_channels=4, out_channels=2)
```

### The loss function

Which loss function should you use? Remember that you're doing classification. How many channels are you dealing with? What kind of activation is needed?

```
loss_function = torch.nn.CrossEntropyLoss()
```

```
# Insert your model here
#layers=(3, 8, 36, 3), groups=64, reduction=16, pretrained=False, progress=True
model = monai.networks.nets.SENet154(spatial_dims = 2, in_channels = 4)
model.to(device)

# Insert your loss function here
loss_function = torch.nn.CrossEntropyLoss()

# Helper function for inferring during validation
def infer_seg(images, model):
    return model(images).argmax(0)
```

```
# use Adam optimizer
optimizer = torch.optim.Adam(model.parameters(), 1e-3)
```

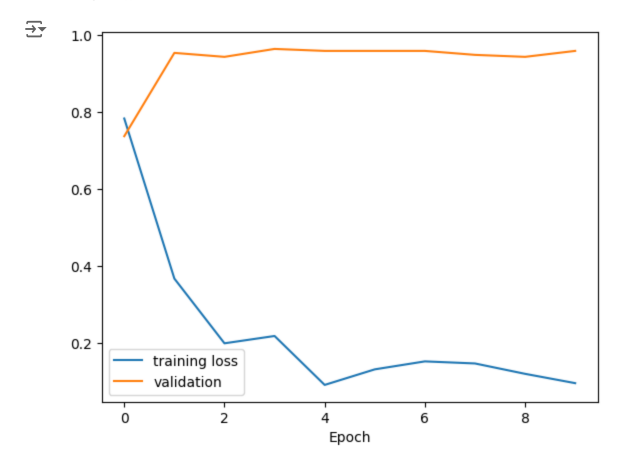
## 7. Train!

```
# Start a typical PyTorch training
max_epochs = 10
val_interval = 1
best_metric = -1
best_metric_epoch = -1
losses, metrics = [], []
tr = trange(max_epochs)
for epoch in tr:
    epoch loss = 0
    # Training phase
    model.train()
    for batch data in train loader:
        inputs, labels = batch_data["image"], batch_data["label"]
        inputs, labels = inputs.to(device), labels.to(device)
        # Convert one-hot encoded labels to class indices, if necessary
        if len(labels.shape) > 1 and labels.shape[1] > 1: # Check if labels are one-hot encoded
            labels = labels.argmax(dim=1) # Convert to class indices
        # If labels have extra dimensions, squeeze them
        labels = labels.squeeze() # Ensures labels are 1D tensor of class indices
        optimizer.zero_grad()
        outputs = model(inputs)
        # CrossEntropyLoss already applies LogSoftmax, so no need for Softmax
        loss = loss function(outputs, labels.long()) # Ensure labels are long type for CrossEntropyLoss
        loss.backward()
        optimizer.step()
```

```
epoch loss += loss.item()
epoch loss /= len(train loader)
losses.append(epoch loss)
# Validation phase
if (epoch + 1) % val interval == 0:
    model.eval() # Set model to evaluation mode
    num_correct = 0
    metric_count = 0
    with torch.no_grad(): # Disable gradient calculation during validation
        for val_data in val_loader:
            val_images, val_labels = val_data["image"], val_data["label"]
            val_images, val_labels = val_images.to(device), val_labels.to(device)
            # Convert one-hot encoded labels to class indices, if necessary
            if len(val labels.shape) > 1 and val labels.shape[1] > 1:
                val labels = val_labels.argmax(dim=1) # Convert to class indices
            val labels = val labels.squeeze() # Ensure labels are 1D tensor of class indices
            val outputs = model(val images)
            # Compare the predicted class (argmax) with the true class
            value = torch.eq(val outputs.argmax(dim=1), val labels)
            metric_count += len(value)
            num_correct += int(value.sum().item())
    metric = num_correct / metric_count
    metrics.append(metric)
    # If the current validation metric is better, save the model
    if metric > best_metric:
        best metric = metric
        best metric epoch = epoch + 1
       torch.save(model.state_dict(), best_model_path)
tr.set_description(f"Loss: {epoch_loss:.4f}, best val metric: {best_metric:.4f} at epoch {best_metric_epoch}")
```

```
→ Loss: 0.0952, best val metric: 0.9639 at epoch 4: 100%| 10/10 [07:16<00:00, 43.69s/it]
```

```
fig, ax = plt.subplots(facecolor='white')
ax.plot(losses, label="training loss")
ax.plot(metrics, label="validation")
ax.set_xlabel("Epoch")
_ = ax.legend()
```



```
model.load_state_dict(torch.load(best_model_path))
_ = model.eval()
```

<ipython-input-25-df4a6896f45c>:1: FutureWarning: You are using `torch.load` with `weights\_only=False` (the current def model.load\_state\_dict(torch.load(best\_model\_path))