

<u>Image Source (https://medium.com/stanford-ai-for-healthcare/its-a-no-brainer-deep-learning-for-brain-mr-images-f60116397472)</u>

# **Brain Tumor Auto-Segmentation for Magnetic Resonance Imaging (MRI)**

Welcome to the final part of the "Artificial Intelligence for Medicine" course 1!

You will learn how to build a neural network to automatically segment tumor regions in brain, using MRI (Magnetic Resonance Imaging (https://en.wikipedia.org/wiki/Magnetic resonance imaging)) scans

The MRI scan is one of the most common image modalities that we encounter in the radiology field.

Other data modalities include:

- Computer Tomography (CT) (https://en.wikipedia.org/wiki/CT\_scan),
- <u>Ultrasound (https://en.wikipedia.org/wiki/Ultrasound)</u>
- X-Rays (https://en.wikipedia.org/wiki/X-ray).

In this assignment we will be focusing on MRIs but many of our learnings applies to other mentioned modalities as well. We'll walk you through some of the steps of training a deep learning model for segmentation.

#### You will learn:

- What is in an MR image
- Standard data preparation techniques for MRI datasets
- Metrics and loss functions for segmentation
- Visualizing and evaluating segmentation models

### **Outline**

Use these links to jump to particular sections of this assignment!

- 1. Dataset
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- 1.2 MRI Data Processing
- 1.3 Exploring the Dataset
- 1.4 Data Preprocessing
  - 1.4.1 Sub-volume Sampling
  - 1.4.2 Standardization
- 2. Model: 3D U-Net
- 3. Metrics
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  - 3.2 Soft Dice Loss
- 4. Training
- <u>5. Evaluation</u>
  - 5.1 Overall Performance
  - 5.2 Patch-level Predictions
  - 5.3 Running on Entire Scans

# **Packages**

In this assignment, we'll make use of the following packages:

- keras is a framework for building deep learning models.
- keras.backend allows us to perform math operations on tensors.
- nibabel will let us extract the images and labels from the files in our dataset.
- numpy is a library for mathematical and scientific operations.
- pandas is what we'll use to manipulate our data.

# **Import Packages**

Run the next cell to import all the necessary packages, dependencies and custom util functions.

```
In [1]:
    import keras
    import json
    import numpy as np
    import pandas as pd
    import nibabel as nib
    import matplotlib.pyplot as plt

from tensorflow.keras import backend as K

import util
```

Using TensorFlow backend.

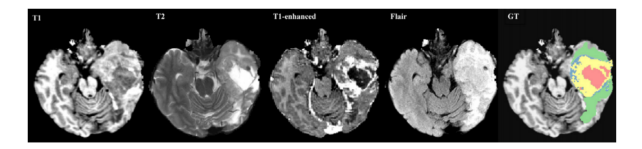
### 1 Dataset

### 1.1 What is an MRI?

Magnetic resonance imaging (MRI) is an advanced imaging technique that is used to observe a variety of diseases and parts of the body.

As we will see later, neural networks can analyze these images individually (as a radiologist would) or combine them into a single 3D volume to make predictions.

At a high level, MRI works by measuring the radio waves emitting by atoms subjected to a magnetic field.



In this assignment, we'll build a multi-class segmentation model. We'll identify 3 different abnormalities in each image: edemas, non-enhancing tumors, and enhancing tumors.

## 1.2 MRI Data Processing

We often encounter MR images in the <u>DICOM format</u> (https://en.wikipedia.org/wiki/DICOM).

 The DICOM format is the output format for most commercial MRI scanners. This type of data can be processed using the <u>pydicom</u> (<a href="https://pydicom.github.io/pydicom/stable/getting\_started.html">https://pydicom.github.io/pydicom/stable/getting\_started.html</a>) Python library.

In this assignment, we will be using the data from the <u>Decathlon 10 Challenge</u> (<a href="https://decathlon-10.grand-challenge.org">https://decathlon-10.grand-challenge.org</a>). This data has been mostly preprocessed for the competition participants, however in real practice, MRI data needs to be significantly pre-preprocessed before we can use it to train our models.

# 1.3 Exploring the Dataset

Our dataset is stored in the <u>NifTI-1 format (https://nifti.nimh.nih.gov/nifti-1/)</u> and we will be using the <u>NiBabel library (https://github.com/nipy/nibabel)</u> to interact with the files. Each training sample is composed of two separate files:

The first file is an image file containing a 4D array of MR image in the shape of (240, 240, 155, 4).

- The first 3 dimensions are the X, Y, and Z values for each point in the 3D volume, which is commonly called a voxel.
- The 4th dimension is the values for 4 different sequences
  - 0: FLAIR: "Fluid Attenuated Inversion Recovery" (FLAIR)
  - 1: T1w: "T1-weighted"
  - 2: t1gd: "T1-weighted with gadolinium contrast enhancement" (T1-Gd)
  - 3: T2w: "T2-weighted"

The second file in each training example is a label file containing a 3D array with the shape of (240, 240, 155).

- The integer values in this array indicate the "label" for each voxel in the corresponding image files:
  - 0: background
  - 1: edema
  - 2: non-enhancing tumor
  - 3: enhancing tumor

We have access to a total of 484 training images which we will be splitting into a training (80%) and validation (20%) dataset.

Let's begin by looking at one single case and visualizing the data! You have access to 10 different cases via this notebook and we strongly encourage you to explore the data further on your own.

We'll use the <u>NiBabel library (https://nipy.org/nibabel/nibabel\_images.html)</u> to load the image and label for a case. The function is shown below to give you a sense of how it works.

```
In [2]: # set home directory and data directory
    HOME_DIR = "./BraTS-Data/"
    DATA_DIR = HOME_DIR

def load_case(image_nifty_file, label_nifty_file):
    # load the image and label file, get the image content and return a numpy arr
    ay for each
    image = np.array(nib.load(image_nifty_file).get_fdata())
    label = np.array(nib.load(label_nifty_file).get_fdata())
    return image, label
```

We'll now visualize an example. For this, we use a pre-defined function we have written in the util.py file that uses matplotlib to generate a summary of the image.

The colors correspond to each class.

- Red is edema
- Green is a non-enhancing tumor
- Blue is an enhancing tumor.

Do feel free to look at this function at your own time to understand how this is achieved.

```
In [3]: image, label = load_case(DATA_DIR + "imagesTr/BRATS_003.nii.gz", DATA_DIR + "labe
lsTr/BRATS_003.nii.gz")
image = util.get_labeled_image(image, label)

util.plot_image_grid(image)
```

We've also written a utility function which generates a GIF that shows what it looks like to iterate over each axis.

```
In [4]: image, label = load_case(DATA_DIR + "imagesTr/BRATS_003.nii.gz", DATA_DIR + "labe
    lsTr/BRATS_003.nii.gz")
    util.visualize_data_gif(util.get_labeled_image(image, label))
```

```
Out[4]:
```

**Reminder:** You can explore more images in the imagesTr directory by changing the image name file.

# 1.4 Data Preprocessing using patches

While our dataset is provided to us post-registration and in the NIfTI format, we still have to do some minor pre-processing before feeding the data to our model.

#### Generate sub-volumes

We are going to first generate "patches" of our data which you can think of as sub-volumes of the whole MR images.

- The reason that we are generating patches is because a network that can process the entire volume at once will simply not fit inside our current environment's memory/GPU.
- Therefore we will be using this common technique to generate spatially consistent sub-volumes of our data, which can be fed into our network.
- Specifically, we will be generating randomly sampled sub-volumes of shape [160, 160, 16] from our images.
- Furthermore, given that a large portion of the MRI volumes are just brain tissue or black background without any tumors, we want to make sure that we pick patches that at least include some amount of tumor data.
- Therefore, we are only going to pick patches that have at most 95% non-tumor regions (so at least 5% tumor).
- We do this by filtering the volumes based on the values present in the background labels.

Lastly, given that the values in MR images cover a very wide range, we will standardize the values to have a mean of zero and standard deviation of 1.

• This is a common technique in deep image processing since standardization makes it much easier for the network to learn.

Let's walk through these steps in the following exercises.

### 1.4.1 Sub-volume Sampling

Fill in the function below takes in:

- a 4D image (shape: [240, 240, 155, 4])
- its 3D label (shape: [240, 240, 155]) arrays,

#### The function returns:

- A randomly generated sub-volume of size [160, 160, 16]
- Its corresponding label in a 1-hot format which has the shape [3, 160, 160, 16]

### Additionally:

- 1. Make sure that at most 95% of the returned patch is non-tumor regions.
- 2. Given that our network expects the channels for our images to appear as the first dimension (instead of the last one in our current setting) reorder the dimensions of the image to have the channels appear as the first dimension.
- 3. Reorder the dimensions of the label array to have the first dimension as the classes (instead of the last one in our current setting)
- 4. Reduce the labels array dimension to only include the non-background classes (total of 3 instead of 4)

#### Hints

```
In [5]:
        # UNQ C1 (UNIQUE CELL IDENTIFIER, DO NOT EDIT)
         def get sub volume(image, label,
                            orig_x = 240, orig_y = 240, orig_z = 155,
                            output_x = 160, output_y = 160, output_z = 16,
                            num classes = 4, max tries = 1000,
                            background_threshold=0.95):
             .. .. ..
             Extract random sub-volume from original images.
             Args:
                 image (np.array): original image,
                     of shape (orig_x, orig_y, orig_z, num_channels)
                 label (np.array): original label.
                     labels coded using discrete values rather than
                     a separate dimension,
                     so this is of shape (orig_x, orig_y, orig_z)
                 orig_x (int): x_dim of input image
                 orig_y (int): y_dim of input image
                 orig z (int): z dim of input image
                 output_x (int): desired x_dim of output
                 output_y (int): desired y_dim of output
                 output z (int): desired z dim of output
                 num_classes (int): number of class labels
                 max_tries (int): maximum trials to do when sampling
                 background_threshold (float): limit on the fraction
                     of the sample which can be the background
             returns:
                 X (np.array): sample of original image of dimension
                     (num_channels, output_x, output_y, output_z)
                 y (np.array): labels which correspond to X, of dimension
                     (num_classes, output_x, output_y, output_z)
             11 11 11
             # Initialize features and labels with `None`
             X = None
             y = None
             ### START CODE HERE (REPLACE INSTANCES OF 'None' with your code) ###
             tries = 0
             while tries < max_tries:</pre>
                 # randomly sample sub-volume by sampling the corner voxel
                 # hint: make sure to leave enough room for the output dimensions!
                 start_x = np.random.randint(0, orig_x - output_x+1) #None
                 start_y = np.random.randint(0, orig_y - output_y+1) #None
                 start_z = np.random.randint(0, orig_z - output_z+1) #None
                 # extract relevant area of label
                 y = label[start_x: start_x + output_x,
                           start_y: start_y + output_y,
```

```
start_z: start_z + output_z]
    # One-hot encode the categories.
    # This adds a 4th dimension, 'num_classes'
    # (output_x, output_y, output_z, num_classes)
    y = keras.utils.to_categorical(y, num_classes=num_classes) #None
    # compute the background ratio
    bgrd_ratio = np.sum(y[:, :, :, 0])/(output_x * output_y * output_z) #None
    # increment tries counter
    tries += 1
    # if background ratio is below the desired threshold,
    # use that sub-volume.
    # otherwise continue the loop and try another random sub-volume
    if bgrd_ratio < background_threshold:</pre>
        # make copy of the sub-volume
        X = np.copy(image[start_x: start_x + output_x,
                          start_y: start_y + output_y,
                          start_z: start_z + output_z, :])
        # change dimension of X
        # from (x_dim, y_dim, z_dim, num_channels)
        # to (num_channels, x_dim, y_dim, z_dim)
        X = np.moveaxis(X, 3, 0) #None
        # change dimension of y
        # from (x_dim, y_dim, z_dim, num_classes)
        # to (num_classes, x_dim, y_dim, z_dim)
        y = np.moveaxis(y, 3, 0) #None
        ### END CODE HERE ###
        # take a subset of y that excludes the background class
        # in the 'num_classes' dimension
        y = y[1:, :, :, :]
        return X, y
# if we've tried max_tries number of samples
# Give up in order to avoid looping forever.
print(f"Tried {tries} times to find a sub-volume. Giving up...")
```

```
C1M3 Assignment slides
In [6]:
         np.random.seed(3)
         image = np.zeros((4, 4, 3, 1))
         label = np.zeros((4, 4, 3))
         for i in range(4):
             for j in range(4):
                 for k in range(3):
                     image[i, j, k, 0] = i*j*k
                     label[i, j, k] = k
         print("image:")
         for k in range(3):
             print(f"z = \{k\}")
             print(image[:, :, k, 0])
         print("\n")
         print("label:")
         for k in range(3):
             print(f"z = \{k\}")
             print(label[:, :, k])
        image:
        z = 0
        [[0. 0. 0. 0.]
         [0. 0. 0. 0.]
         [0. 0. 0. 0.]
         [0. 0. 0. 0.]]
        z = 1
        [[0. 0. 0. 0.]
         [0. 1. 2. 3.]
         [0. 2. 4. 6.]
         [0.3.6.9.]]
        z = 2
        [[ 0. 0. 0. 0.]
         [0. 2. 4. 6.]
         [ 0. 4. 8. 12.]
         [ 0. 6. 12. 18.]]
        label:
```

```
тарет.
```

```
z = 0
[[0. 0. 0. 0.]
  [0. 0. 0. 0.]
  [0. 0. 0. 0.]
  [0. 0. 0. 0.]]
z = 1
[[1. 1. 1. 1.]
  [1. 1. 1. 1.]
  [1. 1. 1. 1.]
  [2. 2. 2. 2.]
```

```
[2. 2. 2. 2.]
[2. 2. 2. 2.]
[2. 2. 2. 2.]]
```

### Test: Extracting (2, 2, 2) sub-volume

#### Sampled Image:

```
z = 0
[[0. 2.]
[0. 3.]]
z = 1
[[0. 4.]
[0. 6.]]
```

### **Expected output:**

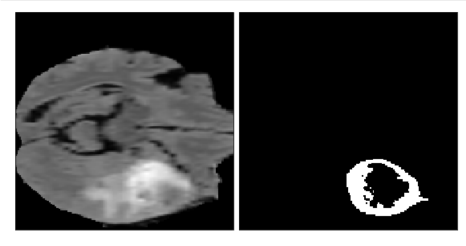
```
Sampled Image:
z = 0
[[0. 2.]
  [0. 3.]]
z = 1
[[0. 4.]
  [0. 6.]]
```

```
Sampled Label:
class = 0
z = 0
[[1. 1.]
  [1. 1.]]
z = 1
[[0. 0.]
  [0. 0.]]
class = 1
z = 0
[[0. 0.]
  [0. 0.]]
z = 1
[[1. 1.]
  [1. 1.]]
```

```
Sampled Label:
class = 0
z = 0
[[1. 1.]
  [1. 1.]]
z = 1
[[0. 0.]
  [0. 0.]]
class = 1
z = 0
[[0. 0.]
  [0. 0.]]
z = 1
[[1. 1.]
  [1. 1.]]
```

You can run the following cell to look at a candidate patch and ensure that the function works correctly. We'll look at the enhancing tumor part of the label.

```
In [9]: image, label = load_case(DATA_DIR + "imagesTr/BRATS_001.nii.gz", DATA_DIR + "labe
lsTr/BRATS_001.nii.gz")
X, y = get_sub_volume(image, label)
# enhancing tumor is channel 2 in the class label
# you can change indexer for y to look at different classes
util.visualize_patch(X[0, :, :, :], y[2])
```



### 1.4.2 Standardization

Next, fill in the following function that given a patch (sub-volume), standardizes the values across each channel and each Z plane to have a mean of zero and standard deviation of 1.

#### Hints

```
In [10]:
         # UNQ_C2 (UNIQUE CELL IDENTIFIER, DO NOT EDIT)
         def standardize(image):
              Standardize mean and standard deviation
                  of each channel and z dimension.
              Args:
                  image (np.array): input image,
                      shape (num_channels, dim_x, dim_y, dim_z)
              Returns:
                  standardized_image (np.array): standardized version of input image
              ### START CODE HERE (REPLACE INSTANCES OF 'None' with your code) ###
              # initialize to array of zeros, with same shape as the image
              standardized_image = np.zeros(image.shape) #None
              # iterate over channels
              for c in range(image.shape[0]):
                  # iterate over the `z` dimension
                  for z in range(image.shape[3]):
                      # get a slice of the image
                      # at channel c and z-th dimension `z`
                      image_slice = image[c,:,:,z]
                      # subtract the mean from image_slice
                      centered = image_slice - np.mean(image_slice) #None
                      # divide by the standard deviation (only if it is different from zer
         o)
                      centered_scaled = centered / np.std(centered) #None
                      # update the slice of standardized image
                      # with the scaled centered and scaled image
                      standardized_image[c, :, :, z] = centered_scaled #None
              ### END CODE HERE ###
              return standardized_image
```

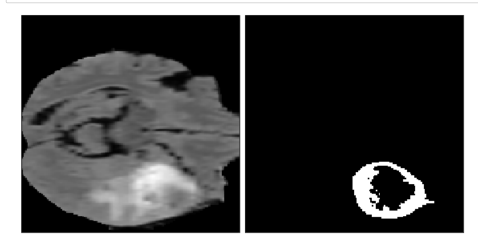
And to sanity check, let's look at the output of our function:

```
In [11]: X_norm = standardize(X)
    print("standard deviation for a slice should be 1.0")
    print(f"stddv for X_norm[0, :, :, 0]: {X_norm[0,:,:,0].std():.2f}")

standard deviation for a slice should be 1.0
stddv for X norm[0, :, :, 0]: 1.00
```

Let's visualize our patch again just to make sure (it won't look different since the imshow function we use to visualize automatically normalizes the pixels when displaying in black and white).

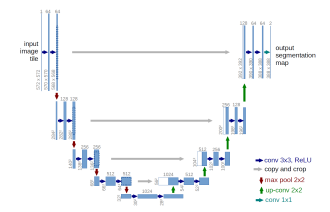
```
In [12]: util.visualize_patch(X_norm[0, :, :, :], y[2])
```



# 2 Model: 3D U-Net

Now let's build our model. In this assignment we will be **building a <u>3D U-net</u>** (https://arxiv.org/abs/1606.06650).

- This architecture will take advantage of the volumetric shape of MR images and is one of the best performing models for this task.
- Feel free to familiarize yourself with the architecture by reading this paper (https://arxiv.org/abs/1606.06650).



### 3 Metrics

# 3.1 Dice Similarity Coefficient

Aside from the architecture, one of the most important elements of any deep learning method is the choice of our loss function.

A natural choice that you may be familiar with is the cross-entropy loss function.

 However, this loss function is not ideal for segmentation tasks due to heavy class imbalance (there are typically not many positive regions).

A much more common loss for segmentation tasks is the **Dice similarity** coefficient, which is a measure of how well two contours overlap.

- The Dice index ranges from 0 (complete mismatch)
- To 1 (perfect match).

In general, for two sets A and B, the Dice similarity coefficient is defined as:

$$\mathrm{DSC}(A,B) = rac{2 imes |A \cap B|}{|A| + |B|}.$$

Here we can interpret A and B as sets of voxels, A being the predicted tumor region and B being the ground truth.

Our model will map each voxel to 0 or 1

- 0 means it is a background voxel
- 1 means it is part of the segmented region.

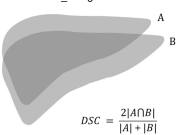
In the dice coefficient, the variables in the formula are:

- x: the input image
- f(x): the model output (prediction)
- y: the label (actual ground truth)

The dice coefficient "DSC" is:

$$ext{DSC}(f,x,y) = rac{2 imes \sum_{i,j} f(x)_{ij} imes y_{ij} + \epsilon}{\sum_{i,j} f(x)_{ij} + \sum_{i,j} y_{ij} + \epsilon}$$

ullet is a small number that is added to avoid division by zero



DSC: Dice similarity coefficient

<u>Image Source (https://www.researchgate.net/figure/Calculation-of-the-Dicesimilarity-coefficient-The-deformed-contour-of-the-liver-from fig4 328671987)</u>

Implement the dice coefficient for a single output class below.

Please use the <u>Keras.sum(x,axis=)</u>
 (<a href="https://www.tensorflow.org/api">https://www.tensorflow.org/api</a> docs/python/tf/keras/backend/sum)
function to compute the numerator and denominator of the dice coefficient.

```
In [13]:
          # UNQ_C3 (UNIQUE CELL IDENTIFIER, DO NOT EDIT)
          def single_class_dice_coefficient(y_true, y_pred, axis=(0, 1, 2),
                                            epsilon=0.00001):
              .....
              Compute dice coefficient for single class.
              Args:
                 y true (Tensorflow tensor): tensor of ground truth values for single clas
          s.
                                              shape: (x_dim, y_dim, z_dim)
                 y_pred (Tensorflow tensor): tensor of predictions for single class.
                                              shape: (x_dim, y_dim, z_dim)
                  axis (tuple): spatial axes to sum over when computing numerator and
                                denominator of dice coefficient.
                                Hint: pass this as the 'axis' argument to the K.sum functio
          n.
                  epsilon (float): small constant added to numerator and denominator to
                                  avoid divide by 0 errors.
              Returns:
                  dice_coefficient (float): computed value of dice coefficient.
              ### START CODE HERE (REPLACE INSTANCES OF 'None' with your code) ###
              dice_numerator = 2.0 * K.sum(y_true * y_pred, axis=axis) + epsilon #None
              dice_denominator = K.sum(y_true, axis=axis) + K.sum(y_pred, axis=axis) + epsi
          lon #None
              dice_coefficient = (dice_numerator) / (dice_denominator) #None
              ### FND CODE HERE ###
              return dice_coefficient
```

```
In [14]:
         # TEST CASES
          sess = K.get session()
          #sess = tf.compat.v1.Session()
          with sess.as_default() as sess:
              pred = np.expand dims(np.eye(2), -1)
              label = np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), -1)
              print("Test Case #1")
              print("pred:")
              print(pred[:, :, 0])
              print("label:")
              print(label[:, :, 0])
              # choosing a large epsilon to help check for implementation errors
              dc = single_class_dice_coefficient(pred, label,epsilon=1)
              print(f"dice coefficient: {dc.eval():.4f}")
              print("\n")
              print("Test Case #2")
              pred = np.expand_dims(np.eye(2), -1)
              label = np.expand_dims(np.array([[1.0, 1.0], [0.0, 1.0]]), -1)
              print("pred:")
              print(pred[:, :, 0])
              print("label:")
              print(label[:, :, 0])
              # choosing a large epsilon to help check for implementation errors
              dc = single_class_dice_coefficient(pred, label,epsilon=1)
              print(f"dice_coefficient: {dc.eval():.4f}")
         Test Case #1
         pred:
         [[1. 0.]
          [0. 1.]]
         label:
         [[1. 1.]
           [0. 0.]]
         dice coefficient: 0.6000
         Test Case #2
         pred:
         [[1. 0.]
```

[0. 1.]]
label:
[[1. 1.]
[0. 1.]]

dice\_coefficient: 0.8333

If you get a different result, please check that you implemented the equation completely.

```
Test Case #1
pred:
[[1. 0.]
 [0. 1.]]
label:
[[1. 1.]
 [0. 0.]]
dice coefficient: 0.6000
Test Case #2
pred:
[[1. 0.]
 [0. 1.]]
label:
[[1. 1.]
 [0. 1.]]
dice_coefficient: 0.8333
```

### **Dice Coefficient for Multiple classes**

Now that we have the single class case, we can think about how to approach the multi class context.

- Remember that for this task, we want segmentations for each of the 3 classes of abnormality (edema, enhancing tumor, non-enhancing tumor).
- This will give us 3 different dice coefficients (one for each abnormality class).
- To combine these, we can just take the average. We can write that the overall dice coefficient is:

$$DC(f,x,y) = rac{1}{3}(DC_1(f,x,y) + DC_2(f,x,y) + DC_3(f,x,y))$$

•  $DC_1$ ,  $DC_2$  and  $DC_3$  are edema, enhancing tumor, and non-enhancing tumor dice coefficients.

For any number of classes, the equation becomes:

$$DC(f,x,y) = rac{1}{N} \sum_{c=1}^{C} \left( DC_c(f,x,y) 
ight)$$

In this case, with three categories, C=3

Implement the mean dice coefficient below. This should not be very different from your singe-class implementation.

Please use the <u>K.mean (https://keras.io/backend/#mean)</u> function to take the average of the three classes.

 Apply the mean to the ratio that you calculate in the last line of code that you'll implement.

```
In [15]:
         # UNO C4 (UNIQUE CELL IDENTIFIER, DO NOT EDIT)
          def dice_coefficient(y_true, y_pred, axis=(1, 2, 3),
                               epsilon=0.00001):
              .....
              Compute mean dice coefficient over all abnormality classes.
              Args:
                  y_true (Tensorflow tensor): tensor of ground truth values for all classe
          s.
                                              shape: (num_classes, x_dim, y_dim, z_dim)
                  y_pred (Tensorflow tensor): tensor of predictions for all classes.
                                              shape: (num_classes, x_dim, y_dim, z_dim)
                  axis (tuple): spatial axes to sum over when computing numerator and
                                denominator of dice coefficient.
                                Hint: pass this as the 'axis' argument to the K.sum
                                      and K.mean functions.
                  epsilon (float): small constant add to numerator and denominator to
                                  avoid divide by 0 errors.
              Returns:
                  dice_coefficient (float): computed value of dice coefficient.
              ### START CODE HERE (REPLACE INSTANCES OF 'None' with your code) ###
              dice_numerator = 2.0 * K.sum(y_true * y_pred, axis=axis) + epsilon #None
              dice_denominator = K.sum(y_true, axis=axis) + K.sum(y_pred, axis=axis) + epsi
          lon #None
              dice_coefficient = K.mean((dice_numerator)/(dice_denominator)) #None
              ### END CODE HERE ###
              return dice_coefficient
```

```
In [16]:
         # TEST CASES
          sess = K.get_session()
          with sess.as_default() as sess:
              pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
              label = np.expand dims(np.expand dims(np.array([[1.0, 1.0], [0.0, 0.0]]), 0),
          -1)
              print("Test Case #1")
              print("pred:")
              print(pred[0, :, :, 0])
              print("label:")
              print(label[0, :, :, 0])
              dc = dice_coefficient(label, pred, epsilon=1)
              print(f"dice coefficient: {dc.eval():.4f}")
              print("\n")
              print("Test Case #2")
              pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
              label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 1.0]]), 0),
          -1)
              print("pred:")
              print(pred[0, :, :, 0])
              print("label:")
              print(label[0, :, :, 0])
              dc = dice_coefficient(pred, label,epsilon=1)
              print(f"dice coefficient: {dc.eval():.4f}")
              print("\n")
              print("Test Case #3")
              pred = np.zeros((2, 2, 2, 1))
              pred[0, :, :, :] = np.expand_dims(np.eye(2), -1)
              pred[1, :, :, :] = np.expand_dims(np.eye(2), -1)
              label = np.zeros((2, 2, 2, 1))
              label[0, :, :, :] = np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), -1)
              label[1, :, :, :] = np.expand_dims(np.array([[1.0, 1.0], [0.0, 1.0]]), -1)
              print("pred:")
              print("class = 0")
              print(pred[0, :, :, 0])
              print("class = 1")
              print(pred[1, :, :, 0])
              print("label:")
              print("class = 0")
              print(label[0, :, :, 0])
```

```
print("class = 1")
  print(label[1, :, :, 0])

dc = dice_coefficient(pred, label,epsilon=1)
  print(f"dice coefficient: {dc.eval():.4f}")

Test Case #1
```

```
Test Case #1
pred:
[[1. 0.]
[0. 1.]]
label:
[[1. 1.]
[0. 0.]]
dice coefficient: 0.6000
Test Case #2
pred:
[[1. 0.]
[0. 1.]]
label:
[[1. 1.]
[0. 1.]]
dice coefficient: 0.8333
Test Case #3
pred:
class = 0
[[1. 0.]
[0. 1.]]
class = 1
[[1. 0.]
[0. 1.]]
label:
class = 0
[[1. 1.]
[0. 0.]]
class = 1
[[1. 1.]
[0. 1.]]
dice coefficient: 0.7167
```

```
Test Case #1
pred:
[[1. 0.]
[0. 1.]]
label:
[[1. 1.]
[0. 0.]]
dice coefficient: 0.6000
Test Case #2
pred:
[[1. 0.]
[0. 1.]]
label:
[[1, 1,]
 [0. 1.]]
dice coefficient: 0.8333
Test Case #3
pred:
class = 0
[[1. 0.]
[0. 1.]]
class = 1
[[1. 0.]
[0. 1.]]
label:
class = 0
[[1. 1.]
 [0. 0.]]
class = 1
[[1. 1.]
 [0. 1.]]
dice coefficient: 0.7167
```

### 3.2 Soft Dice Loss

While the Dice Coefficient makes intuitive sense, it is not the best for training.

- This is because it takes in discrete values (zeros and ones).
- The model outputs *probabilities* that each pixel is, say, a tumor or not, and we want to be able to backpropagate through those outputs.

Therefore, we need an analogue of the Dice loss which takes real valued input. This is where the **Soft Dice loss** comes in. The formula is:

$$\mathcal{L}_{Dice}(p,q) = 1 - rac{2 imes \sum_{i,j} p_{ij} q_{ij} + \epsilon}{\left(\sum_{i,j} p_{ij}^2
ight) + \left(\sum_{i,j} q_{ij}^2
ight) + \epsilon}$$

- *p* is our predictions
- q is the ground truth
- In practice each  $q_i$  will either be 0 or 1.
- $\epsilon$  is a small number that is added to avoid division by zero

The soft Dice loss ranges between

- ullet 0: perfectly matching the ground truth distribution q
- 1: complete mismatch with the ground truth.

You can also check that if  $p_i$  and  $q_i$  are each 0 or 1, then the soft Dice loss is just one minus the dice coefficient.

### **Multi-Class Soft Dice Loss**

We've explained the single class case for simplicity, but the multi-class generalization is exactly the same as that of the dice coefficient.

Since you've already implemented the multi-class dice coefficient, we'll
have you jump directly to the multi-class soft dice loss.

For any number of categories of diseases, the expression becomes:

$$\mathcal{L}_{Dice}(p,q) = 1 - rac{1}{N} \sum_{c=1}^{C} rac{2 imes \sum_{i,j} p_{cij} q_{cij} + \epsilon}{\left(\sum_{i,j} p_{cij}^2
ight) + \left(\sum_{i,j} q_{cij}^2
ight) + \epsilon}$$

Please implement the soft dice loss below!

As before, you will use K.mean()

 Apply the average the mean to ratio that you'll calculate in the last line of code that you'll implement.

```
In [17]:
          # UNQ_C5 (UNIQUE CELL IDENTIFIER, DO NOT EDIT)
          def soft_dice_loss(y_true, y_pred, axis=(1, 2, 3),
                             epsilon=0.00001):
              Compute mean soft dice loss over all abnormality classes.
              Args:
                 y_true (Tensorflow tensor): tensor of ground truth values for all classe
          s.
                                              shape: (num_classes, x_dim, y_dim, z_dim)
                 y_pred (Tensorflow tensor): tensor of soft predictions for all classes.
                                              shape: (num_classes, x_dim, y_dim, z_dim)
                 axis (tuple): spatial axes to sum over when computing numerator and
                                denominator in formula for dice loss.
                                Hint: pass this as the 'axis' argument to the K.sum
                                      and K.mean functions.
                  epsilon (float): small constant added to numerator and denominator to
                                  avoid divide by 0 errors.
              Returns:
                  dice_loss (float): computed value of dice loss.
              ### START CODE HERE (REPLACE INSTANCES OF 'None' with your code) ###
              dice_numerator = 2.0 * K.sum(y_true * y_pred, axis=axis) + epsilon #None
              dice_denominator = K.sum(y_true**2, axis=axis) + K.sum(y_pred**2, axis=axis)
          + epsilon #None
              dice_loss = 1 - K.mean((dice_numerator)/(dice_denominator)) #None
              ### END CODE HERE ###
              return dice_loss
```

```
In [18]: # TEST CASES
sess = K.get_session()
with sess.as_default() as sess:
    pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
    label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), 0),
-1)

    print("Test Case #1")
    print("pred:")
    print(pred[0, :, :, 0])
    print("label:")
    print(label[0, :, :, 0])

    dc = soft_dice_loss(pred, label, epsilon=1)
    print(f"soft dice loss:{dc.eval():.4f}")
```

```
Test Case #1
pred:
[[1. 0.]
  [0. 1.]]
label:
[[1. 1.]
  [0. 0.]]
soft dice loss:0.4000
```

```
Test Case #1
pred:
[[1. 0.]
[0. 1.]]
label:
[[1. 1.]
[0. 0.]]
soft dice loss:0.4000
```

```
In [19]:
    sess = K.get_session()
    with sess.as_default() as sess:
        pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
        label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), 0),
        -1)

        print("Test Case #2")
        pred = np.expand_dims(np.expand_dims(0.5*np.eye(2), 0), -1)
        print("pred:")
        print(pred[0, :, :, 0])
        print([label:")
        print(label[0, :, :, 0])
        dc = soft_dice_loss(pred, label, epsilon=1)
        print(f"soft dice loss: {dc.eval():.4f}")
```

```
Test Case #2
pred:
[[0.5 0. ]
  [0. 0.5]]
label:
[[1. 1.]
  [0. 0.]]
soft dice loss: 0.4286
```

```
Test Case #2
pred:
[[0.5 0.]
[0. 0.5]]
label:
[[1. 1.]
[0. 0.]]
soft dice loss: 0.4286
```

```
In [20]:
          sess = K.get_session()
          with sess.as default() as sess:
              pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
              label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), 0),
          -1)
              print("Test Case #3")
              pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
              label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 1.0]]), 0),
          -1)
              print("pred:")
              print(pred[0, :, :, 0])
              print("label:")
              print(label[0, :, :, 0])
              dc = soft_dice_loss(pred, label, epsilon=1)
              print(f"soft dice loss: {dc.eval():.4f}")
```

```
Test Case #3
pred:
[[1. 0.]
[0. 1.]]
label:
[[1. 1.]
[0. 1.]]
soft dice loss: 0.1667
```

```
Test Case #3
pred:
[[1. 0.]
[0. 1.]]
label:
[[1. 1.]
[0. 1.]]
soft dice loss: 0.1667
```

```
In [21]:
          sess = K.get_session()
          with sess.as default() as sess:
              pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
              label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), 0),
          -1)
              print("Test Case #4")
              pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
              pred[0, 0, 1, 0] = 0.8
              label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 1.0]]), 0),
          -1)
              print("pred:")
              print(pred[0, :, :, 0])
              print("label:")
              print(label[0, :, :, 0])
              dc = soft_dice_loss(pred, label, epsilon=1)
              print(f"soft dice loss: {dc.eval():.4f}")
```

```
Test Case #4
pred:
[[1. 0.8]
  [0. 1. ]]
label:
[[1. 1.]
  [0. 1.]]
soft dice loss: 0.0060
```

```
Test Case #4
pred:
[[1. 0.8]
  [0. 1. ]]
label:
[[1. 1.]
  [0. 1.]]
soft dice loss: 0.0060
```

```
In [22]:
          sess = K.get_session()
          with sess.as default() as sess:
              pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
              label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), 0),
          -1)
              print("Test Case #5")
              pred = np.zeros((2, 2, 2, 1))
              pred[0, :, :, :] = np.expand_dims(0.5*np.eye(2), -1)
              pred[1, :, :, :] = np.expand_dims(np.eye(2), -1)
              pred[1, 0, 1, 0] = 0.8
              label = np.zeros((2, 2, 2, 1))
              label[0, :, :, :] = np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), -1)
              label[1, :, :, :] = np.expand_dims(np.array([[1.0, 1.0], [0.0, 1.0]]), -1)
              print("pred:")
              print("class = 0")
              print(pred[0, :, :, 0])
              print("class = 1")
              print(pred[1, :, :, 0])
              print("label:")
              print("class = 0")
              print(label[0, :, :, 0])
              print("class = 1")
              print(label[1, :, :, 0])
              dc = soft dice loss(pred, label, epsilon=1)
              print(f"soft dice loss: {dc.eval():.4f}")
```

```
Test Case #5
pred:
class = 0
[[0.5 0.]
[0. 0.5]]
class = 1
[[1. 0.8]
[0. 1.]]
label:
class = 0
[[1. 1.]
[0. 0.]]
class = 1
[[1. 1.]
[0. 1.]]
soft dice loss: 0.2173
```

```
Test Case #5
pred:
class = 0
[[0.5 0.]
 [0. 0.5]]
class = 1
[[1. 0.8]
[0. 1.]]
label:
class = 0
[[1. 1.]
[0. 0.]]
class = 1
[[1. 1.]
[0. 1.]]
soft dice loss: 0.2173
```

```
In [23]:
          # Test case 6
          pred = np.array([
                               [
                                   Γ
                                        [1.0, 1.0], [0.0, 0.0]
                                   ],
                                   [1.0, 0.0], [0.0, 1.0]
                                   ]
                               ],
                               [
                                   [
                                       [1.0, 1.0], [0.0, 0.0]
                                   ],
                                   [
                                       [1.0, 0.0], [0.0, 1.0]
                                   ]
                               ],
                             1)
          label = np.array([
                               [
                                   [
                                        [1.0, 0.0], [1.0, 0.0]
                                   ],
                                   [1.0, 0.0], [0.0, 0.0]
                                   ]
                               ],
                               [
                                   [
                                       [0.0, 0.0], [0.0, 0.0]
                                   ],
                                   Γ
                                       [1.0, 0.0], [0.0, 0.0]
                                   ]
                               ]
                             1)
          sess = K.get_session()
          print("Test case #6")
          with sess.as_default() as sess:
              dc = soft_dice_loss(pred, label, epsilon=1)
              print(f"soft dice loss",dc.eval())
```

Test case #6 soft dice loss 0.4375

### **Expected Output**

Test case #6 soft dice loss: 0.4375

Note, if you don't have a scalar, and have an array with more than one value, please check your implementation!

# 4 Create and Train the model

Once you've finished implementing the soft dice loss, we can create the model!

We'll use the unet model 3d function in utils which we implemented for you.

- This creates the model architecture and compiles the model with the specified loss functions and metrics.
- Check out function util.unet\_model\_3d(loss\_function) in the util.py file.

```
In [24]: model = util.unet_model_3d(loss_function=soft_dice_loss, metrics=[dice_coefficien
t])
```

WARNING:tensorflow:From /opt/conda/lib/python3.6/site-packages/tensorflow\_cor e/python/ops/resource\_variable\_ops.py:1630: calling BaseResourceVariable.\_\_ini t\_\_ (from tensorflow.python.ops.resource\_variable\_ops) with constraint is deprecated and will be removed in a future version.

Instructions for updating:

If using Keras pass \*\_constraint arguments to layers.

# 4.1 Training on a Large Dataset

In order to facilitate the training on the large dataset:

- We have pre-processed the entire dataset into patches and stored the patches in the <u>h5py (http://docs.h5py.org/en/stable/)</u> format.
- We also wrote a custom Keras <u>Sequence</u>
   (<a href="https://www.tensorflow.org/api">https://www.tensorflow.org/api</a> docs/python/tf/keras/utils/Sequence)
   class which can be used as a Generator for the keras model to train on large datasets.
- Feel free to look at the VolumeDataGenerator class in util.py to learn about how such a generator can be coded.

Note: <u>Here (https://www.geeksforgeeks.org/keras-fit-and-keras-fit\_generator/)</u> you can check the difference between fit and fit\_generator functions.

To get a flavor of the training on the larger dataset, you can run the following cell to train the model on a small subset of the dataset (85 patches). You should see the loss going down and the dice coefficient going up.

Running model.fit() on the Coursera workspace may cause the kernel to die.

• Soon, we will load a pre-trained version of this model, so that you don't need to train the model on this workspace.

```
# Run this on your local machine only
# May cause the kernel to die if running in the Coursera platform
base_dir = HOME_DIR + "processed/"
with open(base_dir + "config.json") as json_file:
    config = json.load(json_file)
# Get generators for training and validation sets
train_generator = util.VolumeDataGenerator(config["train"], base_dir + "train/",
batch size=3, dim=(160, 160, 16), verbose=0)
valid_generator = util.VolumeDataGenerator(config["valid"], base_dir + "valid/",
batch_size=3, dim=(160, 160, 16), verbose=0)
steps_per_epoch = 20
n_epochs=10
validation_steps = 20
model.fit_generator(generator=train_generator,
        steps_per_epoch=steps_per_epoch,
        epochs=n_epochs,
        use multiprocessing=True,
        validation data=valid generator,
        validation_steps=validation_steps)
# run this cell if you to save the weights of your trained model in cell section
#model.save_weights(base_dir + 'my_model_pretrained.hdf5')
```

# 4.2 Loading a Pre-Trained Model

As in assignment 1, instead of having the model train for longer, we'll give you access to a pretrained version. We'll use this to extract predictions and measure performance.

```
In [25]: # run this cell if you didn't run the training cell in section 4.1
    base_dir = HOME_DIR + "processed/"
    with open(base_dir + "config.json") as json_file:
        config = json.load(json_file)
    # Get generators for training and validation sets
    train_generator = util.VolumeDataGenerator(config["train"], base_dir + "train/",
    batch_size=3, dim=(160, 160, 16), verbose=0)
    valid_generator = util.VolumeDataGenerator(config["valid"], base_dir + "valid/",
    batch_size=3, dim=(160, 160, 16), verbose=0)
```

In [26]: model.load\_weights(HOME\_DIR + "model\_pretrained.hdf5")

In [27]: model.summary()

Layer (type)	Output Shape		
input_1 (InputLayer)	(None, 4, 160, 160,		
conv3d_1 (Conv3D)	(None, 32, 160, 160	3488	input_1[0][0]
activation_1 (Activation) [0]	(None, 32, 160, 160	9, 0	conv3d_1[0]
 conv3d_2 (Conv3D) [0][0]	(None, 64, 160, 160	), 55360	activation_1
activation_2 (Activation) [0]	(None, 64, 160, 160	), 0	conv3d_2[0]
max_pooling3d_1 (MaxPooling3D) [0][0]	(None, 64, 80, 80,	8 0	activation_2
conv3d_3 (Conv3D) _1[0][0]	(None, 64, 80, 80,	8 110656	max_pooling3d
activation_3 (Activation) [0]	(None, 64, 80, 80,	8 0	conv3d_3[0]
 conv3d_4 (Conv3D) [0][0]	(None, 128, 80, 80,	, 221312	activation_3
activation_4 (Activation) [0]	(None, 128, 80, 80,	, 0	conv3d_4[0]
max_pooling3d_2 (MaxPooling3D) [0][0]	(None, 128, 40, 40,	, 0	activation_4
conv3d_5 (Conv3D)	(None, 128, 40, 40,	, 442496	max_pooling3d

_		
2	0	110
_		

activation_5 (Activation) [0]	(None,	128,	40,	40,	0	conv3d_5[0]
 conv3d_6 (Conv3D) [0][0]	(None,	256,	40,	40,	884992	activation_5
activation_6 (Activation) [0]	(None,	256,	40,	40,	0	conv3d_6[0]
<pre>max_pooling3d_3 (MaxPooling3D) [0][0]</pre>	(None,	256,	20,	20,	0	activation_6
conv3d_7 (Conv3D) _3[0][0]	(None,	256,	20,	20,	1769728	max_pooling3d
activation_7 (Activation) [0]	(None,	256,	20,	20,	0	conv3d_7[0]
conv3d_8 (Conv3D) [0][0]	(None,	512,	20,	20,	3539456	activation_7
activation_8 (Activation) [0]	(None,	512,	20,	20,	0	conv3d_8[0]
up_sampling3d_1 (UpSampling3D) [0][0]	(None,	512,	40,	40,	0	activation_8
concatenate_1 (Concatenate) _1[0][0]  [0][0]	(None,	768,	40,	40,	0	up_sampling3d activation_6
 conv3d_9 (Conv3D) [0][0]	(None,	256,	40,	40,	5308672	concatenate_1
activation_9 (Activation) [0]	(None,	256,	40,	40,	0	conv3d_9[0]

conv3d_10 (Conv3D) [0][0]		(None,	256,	40,	40,	1769728	activation_9
activation_10 (Activ	ation)	(None,	256,	40,	40,	0	conv3d_10[0]
up_sampling3d_2 (UpS	ampling3D)	(None,	256,	80,	80,	0	activation_10
concatenate_2 (Conca _2[0][0]	tenate)	(None,	384,	80,	80,	0	up_sampling3d
[0][0]							
		(None,	128,	80,	80,	1327232	concatenate_2
activation_11 (Activ	ation)	(None,	128,	80,	80,	0	conv3d_11[0]
conv3d_12 (Conv3D) [0][0]		(None,	128,	80,	80,	442496	activation_11
activation_12 (Activ	ation)	(None,	128,	80,	80,	0	conv3d_12[0]
up_sampling3d_3 (UpS	ampling3D)	(None,	128,	160,	, 160	0	activation_12
concatenate_3 (Conca _3[0][0]	tenate)	(None,	192,	160,	, 160	0	up_sampling3d
[0][0]							activation_2
conv3d_13 (Conv3D) [0][0]		(None,	64, 2	160,	160,	331840	concatenate_3
activation_13 (Activ	ation)	(None,	64, 2	160,	160,	0	conv3d_13[0]

conv3d_14 (Conv3D) [0][0]	(None,	64, 160	, 160,	110656	activation_13
activation_14 (Activation) [0]	(None,	64, 160	, 160,	0	conv3d_14[0]
conv3d_15 (Conv3D) [0][0]	(None,	3, 160,	160,	195	activation_14
activation_15 (Activation) [0]	(None,	3, 160,	160,	0	conv3d_15[0]
Total params: 16,318,307 Trainable params: 16,318,307 Non-trainable params: 0					

# 5 Evaluation

Now that we have a trained model, we'll learn to extract its predictions and evaluate its performance on scans from our validation set.

# 5.1 Overall Performance

First let's measure the overall performance on the validation set.

 We can do this by calling the keras <u>evaluate generator</u> (<u>https://keras.io/models/model/#evaluate generator</u>) function and passing in the validation generator, created in section 4.1.

#### Using the validation set for testing

- Note: since we didn't do cross validation tuning on the final model, it's okay to use the validation set.
- For real life implementations, however, you would want to do cross validation as usual to choose hyperparamters and then use a hold out test set to assess performance

Python Code for measuring the overall performance on the validation set:

```
val_loss, val_dice = model.evaluate_generator(valid_generator)
print(f"validation soft dice loss: {val_loss:.4f}")
print(f"validation dice coefficient: {val_dice:.4f}")
```

### **Expected output:**

validation soft dice loss: 0.4742 validation dice coefficient: 0.5152

**NOTE:** Do not run the code shown above on the Coursera platform as it will exceed the platform's memory limitations. However, you can run the code shown above locally on your machine or in Colab to practice measuring the overall performance on the validation set.

Like we mentioned above, due to memory limitiations on the Coursera platform we won't be runing the above code, however, you should take note of the **expected output** below it. We should note that due to the randomness in choosing sub-volumes, the values for soft dice loss and dice coefficient will be different each time that you run it.

# 5.2 Patch-level predictions

When applying the model, we'll want to look at segmentations for individual scans (entire scans, not just the sub-volumes)

- This will be a bit complicated because of our sub-volume approach.
- First let's keep things simple and extract model predictions for subvolumes.
- We can use the sub-volume which we extracted at the beginning of the assignment.

In [28]: util.visualize\_patch(X\_norm[0, :, :, :], y[2])





#### Add a 'batch' dimension

We can extract predictions by calling model.predict on the patch.

- We'll add an images per batch dimension, since the predict method is written to take in batches.
- The dimensions of the input should be (images per batch, num\_channels, x\_dim, y\_dim, z\_dim).
- Use numpy.expand dims (https://docs.scipy.org/doc/numpy/reference/generated/numpy.expand\_dims.ht to add a new dimension as the zero-th dimension by setting axis=0

```
In [29]:
```

```
X_norm_with_batch_dimension = np.expand_dims(X_norm, axis=0)
patch pred = model.predict(X norm with batch dimension)
```

WARNING:tensorflow:From /opt/conda/lib/python3.6/site-packages/keras/backend/t ensorflow backend.py:422: The name tf.global variables is deprecated. Please u se tf.compat.v1.global variables instead.

#### Convert prediction from probability into a category

Currently, each element of patch pred is a number between 0.0 and 1.0.

- Each number is the model's confidence that a voxel is part of a given class.
- You will convert these to discrete 0 and 1 integers by using a threshold.
- We'll use a threshold of 0.5.
- In real applications, you would tune this to achieve your required level of sensitivity or specificity.

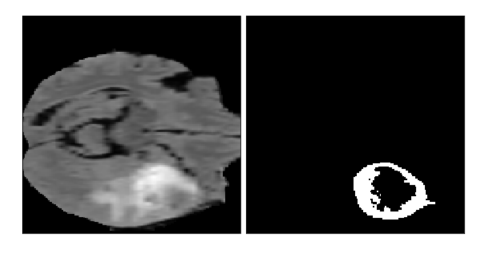
```
In [30]:
```

```
# set threshold.
threshold = 0.5
# use threshold to get hard predictions
patch pred[patch pred > threshold] = 1.0
patch_pred[patch_pred <= threshold] = 0.0</pre>
```

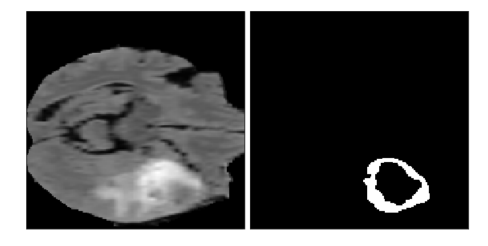
Now let's visualize the original patch and ground truth alongside our thresholded predictions.

```
In [31]: print("Patch and ground truth")
    util.visualize_patch(X_norm[0, :, :, :], y[2])
    plt.show()
    print("Patch and prediction")
    util.visualize_patch(X_norm[0, :, :, :], patch_pred[0, 2, :, :, :])
    plt.show()
```

#### Patch and ground truth



#### Patch and prediction



The model is covering some of the relevant areas, but it's definitely not perfect.

 To quantify its performance, we can use per-pixel sensitivity and specificity.

Recall that in terms of the true positives, true negatives, false positives, and false negatives,

$$sensitivity = \frac{true\ positives}{true\ positives + false\ negatives}$$
 
$$specificity = \frac{true\ negatives}{true\ negatives + false\ positives}$$

Below let's write a function to compute the sensitivity and specificity per output class.

▶ Hints

```
In [32]:
          # UNQ C6 (UNIQUE CELL IDENTIFIER, DO NOT EDIT)
          def compute class sens spec(pred, label, class num):
              Compute sensitivity and specificity for a particular example
              for a given class.
              Args:
                  pred (np.array): binary arrary of predictions, shape is
                                   (num classes, height, width, depth).
                  label (np.array): binary array of labels, shape is
                                    (num classes, height, width, depth).
                  class_num (int): number between 0 - (num_classes -1) which says
                                   which prediction class to compute statistics
                                   for.
              Returns:
                  sensitivity (float): precision for given class_num.
                  specificity (float): recall for given class_num
              # extract sub-array for specified class
              class_pred = pred[class_num]
              class_label = label[class_num]
              ### START CODE HERE (REPLACE INSTANCES OF 'None' with your code) ###
              # compute:
              # true positives
              tp = np.sum((class_pred == 1) & (class_label == 1)) #None
              # true negatives
              tn = np.sum((class_pred == 0) & (class_label == 1)) #None
              #false positives
              fp = np.sum((class_pred == 1) & (class_label == 0)) #None
              # false negatives
              fn = np.sum((class_pred == 0) & (class_label == 0)) #None
              # compute sensitivity and specificity
              sensitivity = tp / (tp + fn) #None
              specificity = tn / (tn + fp) #None
              ### END CODE HERE ###
              return sensitivity, specificity
```

```
In [33]: # TEST CASES
    pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
    label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 0.0]]), 0), -1)

    print("Test Case #1")
    print("pred:")
    print(pred[0, :, :, 0])
    print("label:")
    print(label[0, :, :, 0])

sensitivity, specificity = compute_class_sens_spec(pred, label, 0)
    print(f"sensitivity: {sensitivity:.4f}")
    print(f"specificity: {specificity:.4f}")
```

```
Test Case #1
pred:
[[1. 0.]
  [0. 1.]]
label:
[[1. 1.]
  [0. 0.]]
sensitivity: 0.5000
specificity: 0.5000
```

## **Expected output:**

```
Test Case #1
pred:
[[1. 0.]
  [0. 1.]]
label:
[[1. 1.]
  [0. 0.]]
sensitivity: 0.5000
specificity: 0.5000
```

```
In [34]: print("Test Case #2")

pred = np.expand_dims(np.expand_dims(np.eye(2), 0), -1)
    label = np.expand_dims(np.expand_dims(np.array([[1.0, 1.0], [0.0, 1.0]]), 0), -1)

print("pred:")
    print(pred[0, :, :, 0])
    print("label:")
    print(label[0, :, :, 0])

sensitivity, specificity = compute_class_sens_spec(pred, label, 0)
    print(f"sensitivity: {sensitivity:.4f}")
    print(f"specificity: {specificity:.4f}")
```

```
pred:
[[1. 0.]
  [0. 1.]]
label:
[[1. 1.]
  [0. 1.]]
sensitivity: 0.6667
specificity: 1.0000
```

Test Case #2

### **Expected output:**

```
Test Case #2
pred:
[[1. 0.]
  [0. 1.]]
label:
[[1. 1.]
  [0. 1.]]
sensitivity: 0.6667
specificity: 1.0000
```

```
In [35]:
         # Note: we must explicity import 'display' in order for the autograder to compile
          the submitted code
          # Even though we could use this function without importing it, keep this import i
          n order to allow the grader to work
          from IPython.display import display
          print("Test Case #3")
          df = pd.DataFrame({'y_test': [1,1,0,0,0,0,0,0,0,1,1,1,1,1],
                             'preds_test': [1,1,0,0,0,1,1,1,1,0,0,0,0,0],
                             'category': ['TP','TP','TN','TN','FP','FP','FP','FP','FN'
          ,'FN','FN','FN','FN']
                            })
          display(df)
          pred = np.array( [df['preds_test']])
          label = np.array( [df['y_test']])
          sensitivity, specificity = compute_class_sens_spec(pred, label, 0)
          print(f"sensitivity: {sensitivity:.4f}")
          print(f"specificity: {specificity:.4f}")
```

Test Case #3

	y_test	preds_test	category
0	1	1	TP
1	1	1	TP
2	0	0	TN
3	0	0	TN
4	0	0	TN
5	0	1	FP
6	0	1	FP
7	0	1	FP
8	0	1	FP
9	1	0	FN
10	1	0	FN
11	1	0	FN
12	1	0	FN
13	1	0	FN

sensitivity: 0.4000 specificity: 0.5556

#### **Expected Output**

```
Test case #3 ... sensitivity: 0.2857 specificity: 0.4286
```

### Sensitivity and Specificity for the patch prediction

Next let's compute the sensitivity and specificity on that patch for expanding tumors.

```
In [36]: sensitivity, specificity = compute_class_sens_spec(patch_pred[0], y, 2)
    print(f"Sensitivity: {sensitivity:.4f}")
    print(f"Specificity: {specificity:.4f}")
Sensitivity: 0.0211
```

Sensitivity: 0.0211 Specificity: 0.5867

#### **Expected output:**

Sensitivity: 0.7891 Specificity: 0.9960

We can also display the sensitivity and specificity for each class.

```
In [38]: df = get_sens_spec_df(patch_pred[0], y)
    print(df)
```

```
Edema Non-Enhancing Tumor Enhancing Tumor Sensitivity 0.0357 0.026 0.0211 Specificity 0.1944 0.2632 0.5867
```

#### **Expected output**

```
Edema Non-Enhancing Tumor Enhancing Tumor

Sensitivity 0.9085 0.9505 0.7891

Specificity 0.9848 0.9961 0.996
```

# 5.3 Running on entire scans

As of now, our model just runs on patches, but what we really want to see is our model's result on a whole MRI scan.

- To do this, generate patches for the scan.
- Then we run the model on the patches.
- Then combine the results together to get a fully labeled MR image.

The output of our model will be a 4D array with 3 probability values for each voxel in our data.

 We then can use a threshold (which you can find by a calibration process) to decide whether or not to report a label for each voxel.

We have written a function that stitches the patches together: predict\_and\_viz(image, label, model, threshold)

- Inputs: an image, label and model.
- Ouputs: the model prediction over the whole image, and a visual of the ground truth and prediction.

Run the following cell to see this function in action!

#### Note: the prediction takes some time!

- The first prediction will take about 7 to 8 minutes to run.
- You can skip running this first prediction to save time.

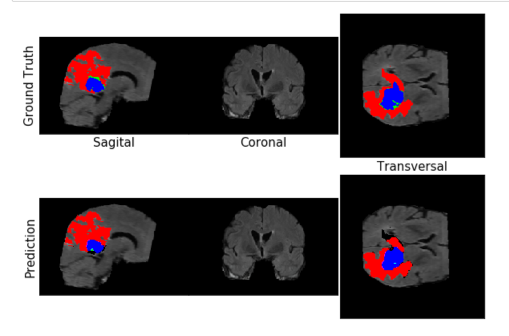
```
In [39]: # uncomment this code to run it
    # image, label = load_case(DATA_DIR + "imagesTr/BRATS_001.nii.gz", DATA_DIR + "la
    belsTr/BRATS_001.nii.gz")
# pred = util.predict_and_viz(image, label, model, .5, loc=(130, 130, 77))
```

Here's a second prediction.

Takes about 7 to 8 minutes to run

Please run this second prediction so that we can check the predictions.

```
image, label = load_case(DATA_DIR + "imagesTr/BRATS_003.nii.gz", DATA_DIR + "labe
lsTr/BRATS_003.nii.gz")
pred = util.predict_and_viz(image, label, model, .5, loc=(130, 130, 77))
```



### Check how well the predictions do

We can see some of the discrepancies between the model and the ground truth visually.

- We can also use the functions we wrote previously to compute sensitivity and specificity for each class over the whole scan.
- First we need to format the label and prediction to match our functions expect.

```
In [41]: whole_scan_label = keras.utils.to_categorical(label, num_classes = 4)
    whole_scan_pred = pred

# move axis to match shape expected in functions
    whole_scan_label = np.moveaxis(whole_scan_label, 3 ,0)[1:4]
    whole_scan_pred = np.moveaxis(whole_scan_pred, 3, 0)[1:4]
```

Now we can compute sensitivity and specificity for each class just like before.

```
In [42]: whole_scan_df = get_sens_spec_df(whole_scan_pred, whole_scan_label)
    print(whole_scan_df)
```

```
Edema Non-Enhancing Tumor Enhancing Tumor Sensitivity 0.0152 0.0001 0.002 Specificity 0.1347 0.6625 0.1687
```

# That's all for now!

Congratulations on finishing this challenging assignment! You now know all the basics for building a neural auto-segmentation model for MRI images. We hope that you end up using these skills on interesting and challenging problems that you face in the real world.