

## Lab3 – Human Pose Estimation

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講解影片: https://youtu.be/JFDif5F-94k

## What is Human Pose Estimation?



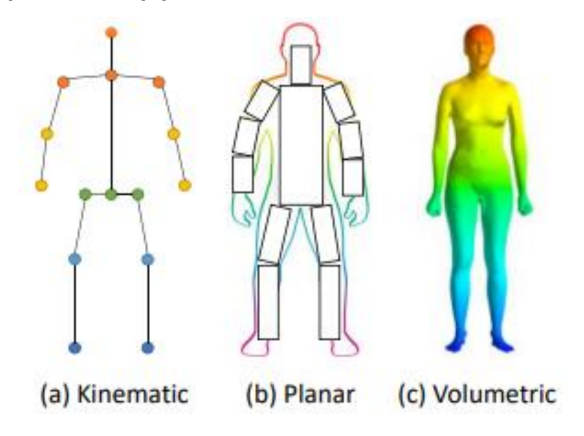
 Human Pose Estimation (HPE) is a way to capture a set of coordinates for each joint (arm, head, torso, etc.,) which is known as a keypoint that can describe a pose of a person. The connection between these points is known as a pair.



# Human body modeling



There are three types of approaches to model the human body:



## Deep Learning-based approaches to 2D HPE



 Deep learning-based approaches are well defined by their ability to generalize any function.

 When it comes to computer vision tasks, deep convolutional neural networks (CNN) surpass all other algorithms, and this is true in HPE as well.

## Deep Learning-based approaches to 2D HPE



DeepPose: Human Pose Estimation via Deep Neural Networks

Toshev, A., & Szegedy, C. (2014). Deeppose: Human pose estimation via deep neural networks. In Proceedings of the IEEE conference on computer vision and pattern recognition (pp. 1653-1660).

 In the paper that they had released, they defined the whole problem as a DNN-based regression problem towards body joints.

 With strong and promising results shown by DeepPose, the HPE research naturally gravitated towards the deep learning-based approaches.

# **HPE using Deep Neural Networks**



- DNNs are very proficient in estimating single human pose but when it comes to estimating multi-human they struggle because:
  - 1. An image can contain multiple numbers of people in different positions.
  - 2. As the number of people increases, the interaction between increases leads to computational complexities.
  - 3. An increase in computational complexities often leads to an increase in inference time in real-time.

## **HPE using Deep Neural Networks**

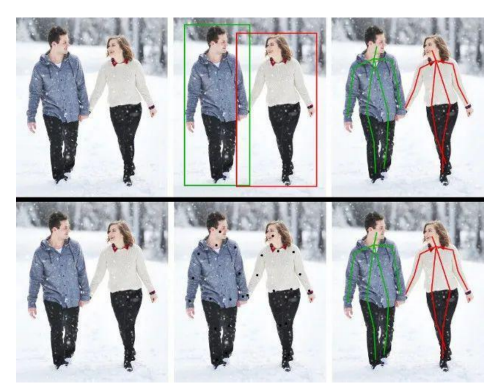


• In order to tackle these problems, we introduced two approaches:

1. Top-down: Localize the humans in the image or video and then estimate the parts followed by calculating the pose.

2. Bottom-up: Estimate the human body parts in the image followed by calculating

the pose.



# **OpenPose**



Cao, Z., Simon, T., Wei, S. E., & Sheikh, Y. (2017). Realtime multi-person 2d pose estimation using part affinity fields. In Proceedings of the IEEE conference on computer vision and pattern recognition (pp. 7291-7299).

An efficient method for multi-person pose estimation.

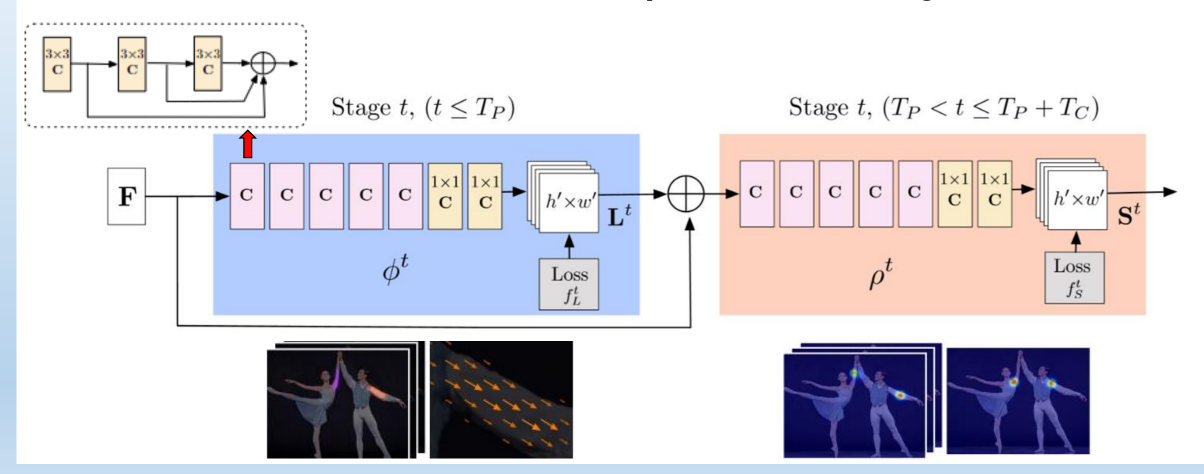
- Bottom-up approach.
- Part Affinity Fields (PAFs), a set of 2D vector fields that encode the location and orientation of limbs over the image domain.

**OpenPose** 

## **Network architecture**

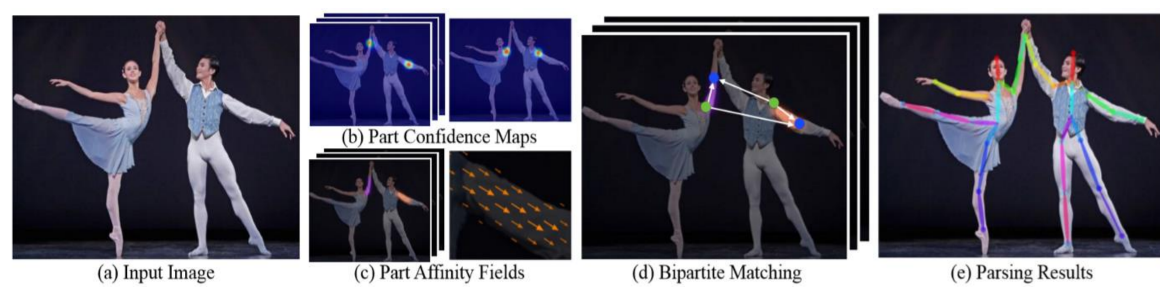


 Iteratively predicts PAFs that encode part-to-part association, shown in blue, and detection Confidence Maps, shown in beige.



## **Method**





- 2D confidence maps S of body part locations
  - S = (S1,S2,...,SJ) has J confidence maps, one per part.
- 2D vector fields L of part affinity fields (PAFs)
  - L = (L1,L2,...,LC) has C vector fields, one per limb.

#### Simultaneous detection and association



- First stage input: feature maps F
  - Initialized by the first 10 layers of VGG-19 and fine-tuned.
  - PAFs:  $L^1 = \phi^1(F)$
- Subsequent stage:

$$\mathbf{L}^t = \phi^t(\mathbf{F}, \mathbf{L}^{t-1}), \ \forall 2 \le t \le T_P, \tag{1}$$

Confidence maps stage:

$$\mathbf{S}^{T_P} = \rho^t(\mathbf{F}, \mathbf{L}^{T_P}), \ \forall t = T_P, \tag{2}$$

$$\mathbf{S}^t = \rho^t(\mathbf{F}, \mathbf{L}^{T_P}, \mathbf{S}^{t-1}), \ \forall T_P < t \le T_P + T_C, \quad (3)$$

#### Simultaneous detection and association



Loss function(L2 loss):

$$f_{\mathbf{L}}^{t_i} = \sum_{c=1}^{C} \sum_{\mathbf{p}} \mathbf{W}(\mathbf{p}) \cdot \|\mathbf{L}_c^{t_i}(\mathbf{p}) - \mathbf{L}_c^*(\mathbf{p})\|_2^2, \tag{4}$$

$$f_{\mathbf{S}}^{t_k} = \sum_{j=1}^{J} \sum_{\mathbf{p}} \mathbf{W}(\mathbf{p}) \cdot \|\mathbf{S}_j^{t_k}(\mathbf{p}) - \mathbf{S}_j^*(\mathbf{p})\|_2^2,$$
 (5)

The overall objective:

$$f = \sum_{t=1}^{T_P} f_{\mathbf{L}}^t + \sum_{t=T_P+1}^{T_P+T_C} f_{\mathbf{S}}^t.$$
 (6)

# Confidence maps for part detection

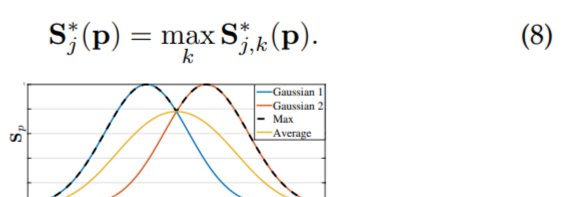


• The value at location p in  $S_{j,k}^*$ :

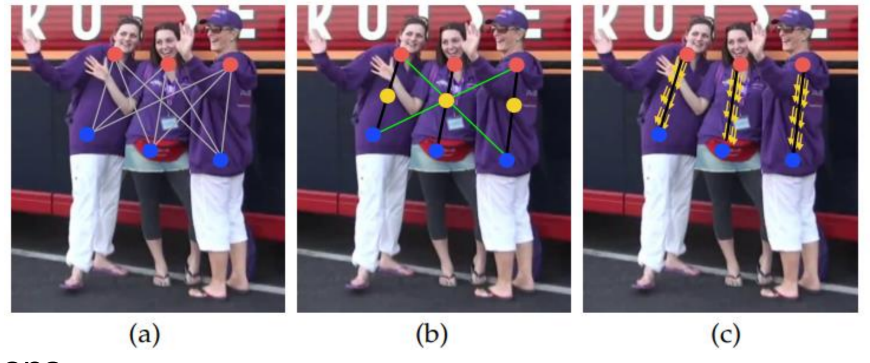
$$\mathbf{S}_{j,k}^{*}(\mathbf{p}) = \exp\left(-\frac{||\mathbf{p} - \mathbf{x}_{j,k}||_{2}^{2}}{\sigma^{2}}\right),\tag{7}$$

• σ: controls the spread of the peak

The groundtruth confidence map:

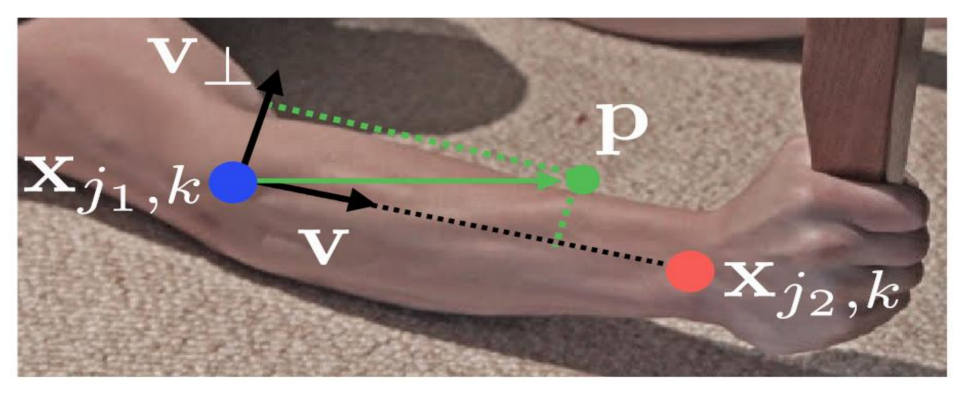






- Two limitations
  - It encodes only the position, and not the orientation, of each limb.
  - It reduces the region of support of a limb to a single point.





Groundtruth PAF at an image point p:

$$\mathbf{L}_{c,k}^*(\mathbf{p}) = \begin{cases} \mathbf{v} & \text{if } \mathbf{p} \text{ on limb } c, k \\ \mathbf{0} & \text{otherwise.} \end{cases}$$
 (9)



Groundtruth PAF at an image point p:

$$\mathbf{L}_{c,k}^*(\mathbf{p}) = \begin{cases} \mathbf{v} & \text{if } \mathbf{p} \text{ on limb } c, k \\ \mathbf{0} & \text{otherwise.} \end{cases}$$
 (9)

Points p:

$$0 \le \mathbf{v} \cdot (\mathbf{p} - \mathbf{x}_{j_1,k}) \le l_{c,k}$$
 and  $|\mathbf{v}_{\perp} \cdot (\mathbf{p} - \mathbf{x}_{j_1,k})| \le \sigma_l$ ,

• Groundtruth part affinity field averages the affinity fields:

$$\mathbf{L}_{c}^{*}(\mathbf{p}) = \frac{1}{n_{c}(\mathbf{p})} \sum_{k} \mathbf{L}_{c,k}^{*}(\mathbf{p}), \tag{10}$$



Score each candidate limb:

$$E = \int_{u=0}^{u=1} \mathbf{L}_c(\mathbf{p}(u)) \cdot \frac{\mathbf{d}_{j_2} - \mathbf{d}_{j_1}}{||\mathbf{d}_{j_2} - \mathbf{d}_{j_1}||_2} du,$$
(11)

where  $\mathbf{p}(\mathbf{u})$  interpolates the position of the two body parts:

$$\mathbf{p}(u) = (1 - u)\mathbf{d}_{j_1} + u\mathbf{d}_{j_2}. \tag{12}$$

# Multi-person parsing using PAFs



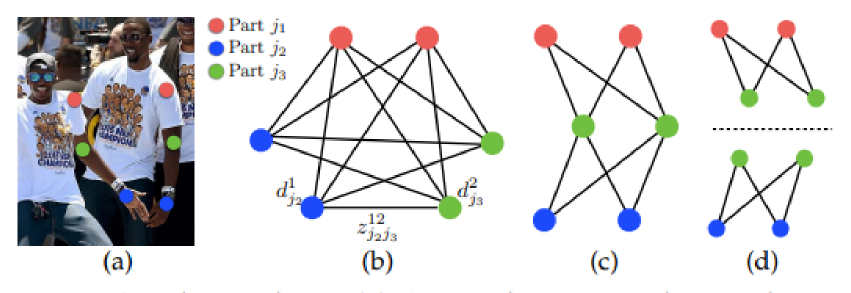


Fig. 6: Graph matching. (a) Original image with part detections. (b) K-partite graph. (c) Tree structure. (d) A set of bipartite graphs.

• These part candidates define a large set of possible limbs.

# Multi-person parsing using PAFs



Determine whether the two keypoints are connected:

$$z_{j_1j_2}^{mn} \in \{0,1\}$$

Final problem we need to solve:

$$\max_{\mathcal{Z}_c} E_c = \max_{\mathcal{Z}_c} \sum_{m \in \mathcal{D}_{j_1}} \sum_{n \in \mathcal{D}_{j_2}} E_{mn} \cdot z_{j_1 j_2}^{mn}, \tag{13}$$

s.t. 
$$\forall m \in \mathcal{D}_{j_1}, \sum_{n \in \mathcal{D}_{j_2}} z_{j_1 j_2}^{mn} \le 1,$$
 (14)

$$\forall n \in \mathcal{D}_{j_2}, \sum_{m \in \mathcal{D}_{j_1}} z_{j_1 j_2}^{mn} \le 1,$$
 (15)

# Multi-person parsing using PAFs



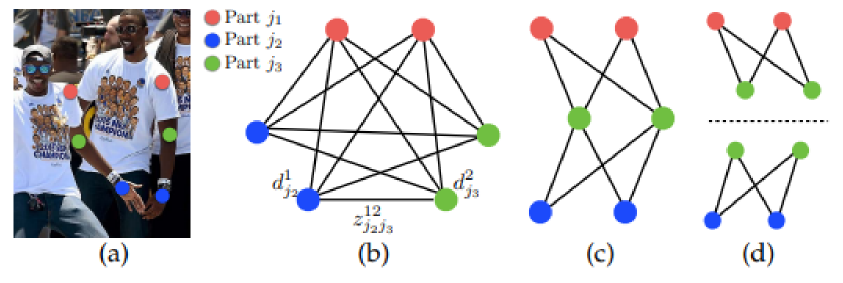
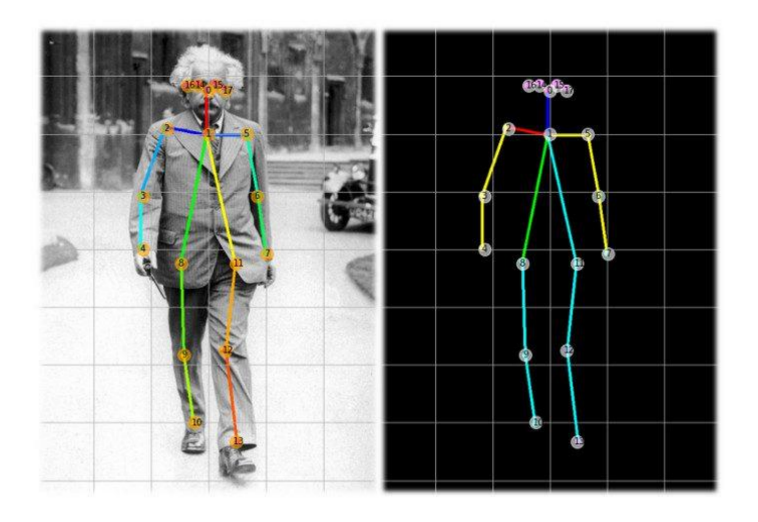


Fig. 6: Graph matching. (a) Original image with part detections. (b) K-partite graph. (c) Tree structure. (d) A set of bipartite graphs.

## **Human Pose Estimation**

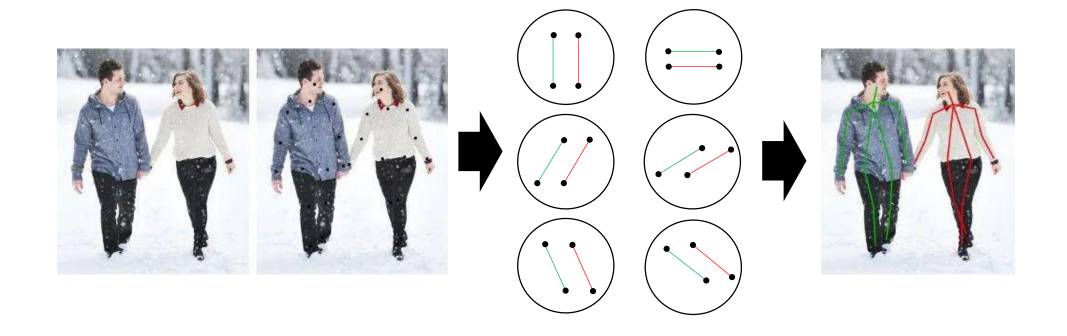




## **EAI Lab3 introduction**



Design your own grouping algorithm.



## **Environment setting**



#### Anaconda

Anaconda is a distribution of the Python programming languages for scientific computing (data science, machine learning applications, large-scale data processing, predictive analytics, etc.), that aims to simplify package management and deployment. (download link)



# **Environment setting**



- Download and extract EAI\_Lab3 file
- Open Anaconda Prompt and type the following command line below:
  - conda create --name poseLab python=3.8 (your own env name)
  - conda activate poseLab (your own env name)
  - conda install git
  - cd C:\Users\user\Desktop\ EAI\_Lab3 (You need to use your own path.)
  - pip install -r requirements.txt

#### Lab Task



```
position meaning:
                                                                    , right wrist , left shoulder
       [nose
                  , neck
                                  , right_shoulder , right_elbow
        left_elbow , left_wrist
                                  , right_hip
                                                , right_knee
                                                                    , right_ankle , left_hip
       left_knee , left_ankle
                                                , left_eye
                                                                    , right_ear
                                                                                 , left_ear
                                  , right_eye
        score, parts_num],
```

- Complete EAI\_Lab3/estimation/merge.py
- Input: subset = [[-1.

- 1.9540779 2.

1.76488239 2.

## Lab Task



[[-1.	1.	2.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
	2.	1			
	1.	-1.	-1.	-1.	5.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
1.76488239	2.	1			
[-1.	-1.	2.	3.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
1.69892728	2.	1			
[ 0.	1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
1.89980185	2.	]			
[ 0.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	6.	-1.	-1.	-1.
1.78199434	2.	]			
[-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	6.	-1.	8.	-1.
1.73637748	2.	]			
[ 0.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	7.	-1.	-1.
1.82718945	2.	1			
[-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	-1.	-1.	-1.
-1.	-1.	-1.	7.	-1.	9.
1.86717367	2.	1			
[-1.	-1.	-1.	-1.	-1.	5.
-1. -1.	-1.	-1.	-1.	-1.	-1.
	-1.	-1.	-1.	-1.	9.
1.6537441	2.	11			

## Lab Task

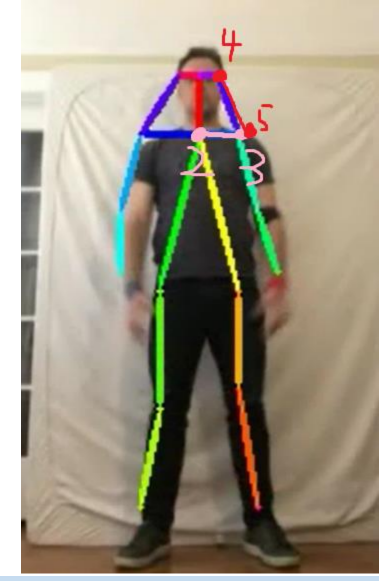


# Some problems you may encounter



• The connections may have misjudgments and may encounter some conflicts.

 Remove some conflicting connections when encountering conflicts



#### Demo



- 1. Revise the path in demo\_image.py
  - input\_image = "./images/1.jpg"
  - output\_image = "output.png"
- 2. Open Anaconda Prompt
- 3. Type the following command line below:
  - conda activate poseLab (your own env name)
  - C:\Users\user\Desktop\EAI\_Lab3 (You need to use your own path.)
  - python demo\_image.py

demo\_image.py1. main()

```
from tf netbuilder ext.extensions import register tf netbuilder extensions
     model_weights_path = "./model_weights/openpose"
     input image = "./images/1.jpg"
     output image = "output.png"
     register_tf_netbuilder extensions()
     module = importlib.impo (variable) module: ModuleType
     create model = getattr(module, "create_openpose")
     model = create model()
     model.load weights(model weights path)
     img = cv2.imread(input image) # B,G,R order
     input img = img[np.newaxis, :, :, [2, 1, 0]]
     inputs = tf.convert to tensor(input img)
     outputs = model.predict(inputs)
     pafs = outputs[10][0, ...]
     heatmaps = outputs[11][0, ...]
     cfg = get default configuration()
     coordinates = get coordinates(cfg, heatmaps)
     connections = get_connections(cfg, coordinates, pafs)
     # ↓ You can analsis the time of your algo. ↓ #
    x = time.time()
     skeletons = estimate(cfg, connections)
43 y = time.time()
     print(y-x) # Inference time
     output = draw(cfg, img, coordinates, skeletons, resize_fac=8)
     cv2.imwrite(output image, output)
```



- coordinates.py
  - 1. Finds the Gaussian peaks coordinates(x,y) in the heatmaps.
  - 2. Assign the Gaussian peaks Identity number.

```
param config: pose estimation configuration
:param heatmaps: heatmaps
 param threshold: threshold for the intensity value
                   in the heatmap at the position of a peak
 return: dictionary:
   { body_part_name:
       [(x ,y, score, id),
all peaks = dict()
peak counter = 0
  Find gaussian peak
 or part_meta in config.body_parts.values():
    hmap = heatmaps[:, :, part_meta.heatmap_idx]
   hmap_right = np.zeros(hmap.shape)
   hmap_right[:, 1:] = hmap[:, :-1]
   hmap_left = np.zeros(hmap.shape)
   hmap_left[:, :-1] = hmap[:, 1:]
   hmap_down = np.zeros(hmap.shape)
   hmap_down[1:, :] = hmap[:-1, :]
   hmap up = np.zeros(hmap.shape)
   hmap_up[:-1, :] = hmap[1:, :]
   peaks_binary = np.logical_and.reduce(
        (hmap >= hmap_right,
        hmap >= hmap_left,
        hmap >= hmap down,
        hmap >= hmap up.
        hmap > threshold))
   peaks = list(zip(np.nonzero(peaks_binary)[1],
                     np.nonzero(peaks_binary)[0]))
   # pack and give id
   sequence_numbers = range(peak_counter, peak_counter + len(peaks))
   peaks_with_score_and_id = [peak + (
       hmap[peak[1], peak[0]],
       seq_num) for peak, seq_num in zip(peaks, sequence_numbers)]
   all_peaks[part_meta.body_part.name] = peaks_with_score_and_id
   peak_counter += len(peaks)
```



- connections.py
  - 1. Finds the connection candidates and returns only valid connections.

$$\mathbf{L}_{c,k}^*(\mathbf{p}) = \begin{cases} \mathbf{v} & \text{if } \mathbf{p} \text{ on limb } c, k \\ \mathbf{0} & \text{otherwise.} \end{cases}$$
 (9)

Here,  $\mathbf{v} = (\mathbf{x}_{j_2,k} - \mathbf{x}_{j_1,k})/||\mathbf{x}_{j_2,k} - \mathbf{x}_{j_1,k}||_2$  is the unit vector in the direction of the limb.

```
Finds the connection candidates and returns only valid connections.
:param config: pose estimation configuration.
:param coords: dictionary with coordinates of all body parts.
:param paf: paf maps.
:param threshold: threshold for the intensity value in paf for a given mid point. If value at a mid point
is below the threshold the mid point is not taken into account.
:param mid num: number of mid point for sampling
:param minimum mid_num: minimum number of valid mid points for the connection candidate
:return: list of arrays containing identified connections of a given type :
       array(
         [id1, id2, score1, score2, total_score]
         [id1, id2, score1, score2, total_score]
        array(
all_cand_connections = []
for conn in config.connection_types:
   # select dx and dy PAFs for this connection type
   paf_dx = paf[:, :, conn.paf_dx_idx]
   paf_dy = paf[:, :, conn.paf_dy_idx]
   # get coordinates lists for 2 body part types which belong to the current connection type
   cand a = coords[conn.from body part.name]
   cand_b = coords[conn.to_body_part.name]
   n_a = len(cand_a)
   n_b = len(cand_b)
   max connections = min(n a, n b)
   # lets check each combination of detected 2 body parts - candidate connections
   if n a != 0 and n b != 0:
       # here we will store the connection candidates 5 columns:
       # [ body part id1, body part id2, body part score1, body part score2, total score of connection ]
       connection_candidates = np.zeros((0, 5))
       for i in range(n a):
           for i in range(n b):
                # find the distance between the 2 body parts. The expression cand b[i][:2]
               # returns an 2 element array with coordinates x,y
               vec = np.subtract(cand b[j][:2], cand a[i][:2])
               norm = math.sqrt(vec[0] * vec[0] + vec[1] * vec[1])
               # skip the connection if 2 body parts overlaps
               if norm == 0:
                   continue
               # normalize the vector
                vec = np.divide(vec. norm)
```

- connections.py
  - 1. Finds the connection candidates and returns only valid connections.

$$E = \int_{u=0}^{u=1} \mathbf{L}_c(\mathbf{p}(u)) \cdot \frac{\mathbf{d}_{j_2} - \mathbf{d}_{j_1}}{||\mathbf{d}_{j_2} - \mathbf{d}_{j_1}||_2} du,$$
(11)

```
# get the set midpoints between 2 body parts (their coordinates x,y)
              start_end = list(zip(np.linspace(cand_a[i][0], cand_b[j][0], num=mid_num),
                                   np.linspace(cand_a[i][1], cand_b[j][1], num=mid_num)))
              # having the coord of midpoint we can read the intensity value in paf map at the midpoint
              # for dx component
              vec_x = np.array(
                  [paf_dx[int(round(start_end[i][1])), int(
                      round(start_end[i][0]))] for i in range(mid_num)]
              vec_y = np.array(
                  [paf_dy[int(round(start_end[i][1])), int(
                      round(start_end[i][0]))] for i in range(mid_num)]
              # calculate the score for the connection weighted by the distance between body parts
              score_midpts = np.multiply(vec_x, vec[0]) + np.multiply(vec_y, vec[1])
              # get the total score
              total_score = sum(score_midpts) / len(score_midpts)
              # number of midpoints with intensity above the threshold shouldn't be less than 80% of all midpoints
              criterion1 = len(np.nonzero(score_midpts > threshold)[0]) > minimum mid_num
              criterion2 = total score > 0
              if criterion1 and criterion2:
                  # add this connection to the list [id1, id2, score1, score2, total score]
                  connection_candidates = np.vstack(
                      [connection_candidates,
                       [cand_a[i][3],
                        cand_b[j][3],
                        cand_a[i][2],
                        cand_b[j][2],
                        total score
      # sort the array by the total score - descending. (the sorted array is reversed by the expression [::-1])
      sorted_connections = connection_candidates[connection_candidates[:, 4].argsort()][::-1]
      # make sure we get no more than max connections
      all_cand_connections.append(sorted_connections[:max_connections, :])
      # not found any body parts but we still need to add empty list to preserve the correct indexing in the
      # output array
      all_cand_connections.append([])
eturn all cand connections
```

- estimators.py
  - Convert connections to human subsets

```
param connections: valid connections:
:param min_num_body_parts: minimum number of body parts for a skeleton
:param min_score: minimum score value for the skeleton
:return: list of skeletons. Each skeleton has a list of identifiers of body parts:
        [id1, id2,...,idN, score, parts_num],
        [id1, id2,...,idN, score, parts_num]
# 2 extra slots for number of valid parts and overall score
number_of_slots = config.body_parts_size() + 2
# the connections are solely used to group body parts into separate skeletons. As a result we will
# specific body part belonging to a skeleton (plus 2 extra columns for: num of parts and skeleton total score)
# we will be adding the skeletons to this array:
subset = np.empty((0, number_of_slots))
 for k, conn in enumerate(config.connection_types):
    if len(connections[k]) > 0:
        part_a = connections[k][:, [0, 2]] # idA, scoreA
        part_b = connections[k][:, [1, 3]] # idB, scoreB
        # determine the slot number for 2 body parts types
        slot_idx_a = config.body_parts[conn.from_body_part].slot_idx
        slot_idx_b = config.body_parts[conn.to_body_part].slot_idx
        for i in range(len(connections[k])):
            found = 0
            slot idx = [-1, -1]
            for j in range(len(subset)):
                if subset[j][slot_idx_a] == part_a[i, 0] and \
                        subset[j][slot_idx_b] == part_b[i, 0]:
                    slot_idx[found] = j
                    found += 1
            # if find no partA in the subset, create a new subset
            if not found:
                row = -1 * np.ones(number_of_slots)
                row[slot_idx_a] = part_a[i, 0]
                row[slot_idx_b] = part_b[i, 0]
                row[-1] = 2
                row[-2] = part_a[i, 1] + part_b[i, 1] + connections[k][i][2]
                subset = np.vstack([subset, row])
subset = merge(subset)
return subset
```

merge.py1. Grouping

```
def merge(subset, min_num_body_parts=4, min_score=0.4):
   Estimates the skeletons.
   :param connections: valid connections
   :param min_num_body_parts: minimum number of body parts for a skeleton
   :return: list of skeletons. Each skeleton has a list of identifiers of body parts:
          [id1, id2,...,idN, score, parts_num],
          [id1, id2,...,idN, score, parts_num]
   position meaning:
                                      , right_shoulder , right_elbow
                                                                       , right_wrist , left_shoulder
       [ [nose
                     , neck
           left_elbow , left_wrist
                                     , right_hip
                                                     , right_knee
                                                                       , right_ankle , left_hip
           left_knee , left_ankle
                                     , right_eye
                                                      , left_eye
                                                                       , right_ear , left_ear
           score, parts_num],
   # 2 step :
   #---merge----
   # Merge the limbs in the subset
   # score : score
   # parts num : How many limbs are in the subset
   # after merge
   # Delete the non-compliant subset
   # 1. parts_num < 4
   delete_idx = []
   for i in range(len(subset)):
          delete idx.append(i)
   subset = np.delete(subset, delete_idx, axis=0)
   return subset
```

# Report



1. Complete description of your algorithm

2. The results of output subset

3. The results of test pictures

4. Challenges and Solutions

# Score assignment



1. Code(60%)

2. Report(40%)

#### File submission



Upload platform: NCKU moodle

- Upload compressed .zip file, named StudentID\_Lab3.zip, including:
  - merge.py
  - StudentID\_Lab3\_report.pdf

• Upload deadline: 11/5 23:59



## **END**