# Automatic morphology: Application on biological images

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### **ABSTRACT**

Phenotype of species are characterized by several informations like, age, sex, morphological measures and environment parameters. Biologists are familar to manually getting morphological measures. In the case of analysis at the macro level (tissues, small parts of animal ...) that can be done directly by measuring the element geometry: length, width, diameter, angles .... Another way is to take pictures of these elements and to run image processing algorithms. From a collection of beetle pictures, we have initiated this work to test the feasability of automatically positionning landmarks on biological images. The set of images contains 293 beetles and for all an image of the left and right mandibles. For each mandible, a set of 16 and 18 landmarks (resp. left and right) have been set manually. In a previous work[] based on Palanaswami article [] we have shown that if we consider the centroid measure of the mandible as the parameter to obtain, the probabilistic Hough Transform can provide very interesting results. But in the case of the main goal is to fix landmarks to get property about specific area (defined by biologists) the result is not as precised as we have expected. In this next turn, we have kept the same method to segment and to register scene images with model but we have change the way to set the model landmarks on the scene. This new procedure uses a SIFT algorithm as the last operation of our process. We will compare all the obtained results in order to define the new workflow to label the mandible images.

# **Keywords**

Automatic morphology, landmarks identification, image registration.

## 1 INTRODUCTION

In biology, morphology analysis is widely used to keep the changing information of the organism or detecting the difference information between the organisms. From the result of morphology analysis, we can conclude the evolution of an organism family, or we may classify the organisms. Especially in agriculture, morphology is one of best ways to learn about the variations of the insect on crops. The morphology methods may be divided into the groups by the features which are used by the methods such as shape, structure, color, pattern or size of the object. In the aim to study the potential links between these variations and agricultual ecosystems, a set of 291 beetles has been collected with all the information about the sex, place where they are found and agricultural practices in each field were recorded. For each beetle, the morphometric landmarks

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has been defined on each part (each insect includes five parts: head, pronotum, body, left and right mandible) of the insect by the biologies. In this context, we try to indicate the landmarks on two parts of beetle (left and right mandible (see figure 1)). Morphometric landmarks are points that can be defined in all speciemns and located precisely. Landmarks are widely used in many biological studies and they are currently included into the classification procedures.





(a) Left mandible

(b) Right mandible

Figure 1: The mandibles of beetle

In this paper, we focus on a method that can automatic identification of landmarks on 2D images of beetle, specify the mandibles of beetle. Through whole the article, we use two images to automatic extraction the landmarks: model image and scene image. The

method mainly includes four stages: firstly, we extract the features of the object in the image; secondly, generalizing Hough transform is used to detect the presence between two images; thirdly, the translation and rotation is identified by image registration; finally, a refinement of the estimated landmarks is done by comparing the descriptors between model landmarks and estimated landmarks.

In section 2, the steps of our methods will be discussed. All experiments and evaluation are described in section 3. Finally, we have some conclusion in the section 4.

#### 2 **METHOD**

For each image, a set of manual landmarks have been set by biologists corresponding to the morphological points of interest (18 landmarks for each right mandible and 16 landmarks for each left mandible). The automatic procedure to determine the landmarks is based on the segmentation of the image. The presence between two images are indicated by generalizing Hough transform, along with theirs translation and rotation are determine by applying registration. The last step is using descriptors measure (L2 distance) to verify the location of estimated landmarks.

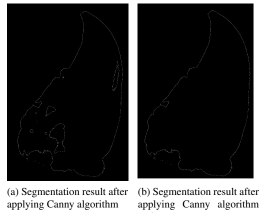
#### **Image segmentation** 2.1

In the methods of image processing, feature extraction is always a important stage. Besides, depending on the nature of the method, the methods are applied before (pre-processing) or after (post-process) extracting the features. In our method, the original Canny alogrithm[1] is ideal for detect the curves on the image. The ratio between lower threshold and upper threshold which are used in Canny algorithm is 1:3 (lower threshold approximated to 1 times threshold value and upper threshold set to 3 times threshold value). This value is evaluated by the experiments. The threshold value has been determined by analyzing the image histogram. During appling the Canny alogrithm to detect the curves of object, the gradient direction of each pixel which belongs to the curves is kept for the next step of the method.

In this study, the aim of segmentation stage is determined the outer border of the object which can be used to reconstruct the shape of the object as well as provide the best data for next step. The curves from Canny algorithm will be post-processed to remove the unnecessary curves i.e hole inside the border (see figure 2).

# **Generalizing Hough transform**

The generalizing Hought transform (GHT)[2] is one of the keys of this study. With two input images, GHT is used to recognize the similar between two images and estimate the landmarks of an image based on the manual landmarks of other image. The GHT includes two



applying Canny algorithm and post-process

Figure 2: The segmentation results of the image

phases: training and recognition.

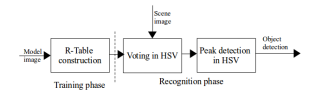


Figure 3: Block diagram of GHT

In the training phase of GHT, model image is used to construct R-table. This is table that contains the polar value of each point on model's curves (the result of segmentation). A polar coordinate system is initialized in the model image by fixing a reference point and using it as origin. The R-table records the polar coordinate values of all boundary points. The rows of R-table are indexed by the gradient directions of the points on the curves. It means that with a gradient direction can be exists many polar coordinate values.

Index	Polar values
gradient 1	(r1,l1), (r2,l2),(r3,l3)
gradient 2	(r2,l2),(r4,l4),(r1,l5),
gradient n	(r1,l1),(r4,l5),(r5,l5),

Table 1: An example of R-Table

During the recognition phase, a 2D accumulator is used, called Hough Space Voting (HSV). The axes of HSV express for the information of polar coordinate system. For each point on scene's curves, we try to find a record in R-table that corresponding with the gradient direction of each scene point. The voting will be carried out at all location in HSV that found in R-table. At the end of voting process, if the scene image is identical to the model image, then the cell where have the highest number of votes corresponds to the reference point of the model in the scene image. Besides, the peak value would be equal to the number of curve points of the scene image when the model and scene image match perfectly.

# 2.3 Image registration

Our study is working on the 2D image, and the transformation between two images are inevitable. Though, GHT is just hired to detect the presence of the model image in the scene image. Translation and rotation are determined by applying an alignment method based on the set of boundary points of two images. In this case, we use Principal Component Analysis Iteration (PCAI) to finish this work. This method is improved from PCA method[5]. Firstly, the centroid point of each image is defined by getting mean coordinate of all boundary points. Secondly, a principal axis is indicated for the images, this is a connected line from centroid point to a point in list of curve points. For each point in curve, we create a line from centroid point to it, and assume it as an axis. The mean perpendicular distance from the remaining points to axis are calculated. This work is finished for all points on curve, and the principal axis is line that has minimum mean perpendicular distance to other points. The translation is indicated by translation between two centroid points. The rotation is the angle between the principal axis of two images. As well as, the rotation direction is determined by checking each direction. HHowever, in some case, the translation and rotation between two images are wrong indicated. To make sure that we have obtained the best match between the images. We applied an iteration to get the best register of them. And instead consider all the points in the curve, we just need to consider a subset of it. The curve points are sorted by y-coordinate first. Then a half points are extracted to check. The PCAI is applied on two new subsets until obtaining best matching of two images.



Figure 4: A registration result between two images.

At the end of step, the landmarks of scene image are estimated by the manual landmarks of model image. Hence, before estimating the landmarks, a translation and rotation are applied on scene image to match the pose between the scene and the model. The obtained landmarks is computed from the distance from the centroid point to each manual landmarks of the model image.

#### 2.4 Refinement the estimated landmarks

The GHT and registration step gave us the estimated coordinate of scene's landmarks. But sometimes the landmarks can stay at the incorrect location. For this reason, at the end of the method, we apply descriptors measure[6] to verify the location of estimated landmarks. The location of manual and estimated landmarks are presented as the descriptors. And L2 distance is applied to measure the distance between them. The equation of L2-distance is given by equation (1).

$$L(x,y) = \sum_{x_i, y_i} \sqrt{(x_i - y_i)^2}$$
 (1)

Where  $x_i, y_i$  are the corresponding location in the descriptors.

For each manual landmark in the model, a sub-image I with the size  $s_1$  is created; and the corresponding estimated landmark on the scene, a template T with the size  $s_2$  is created ( $s_1 < s_2$ ). For each pixel in I, the orientation and gradient magnitude are calculated. The descriptor is created for image I. This is a histogram with 8-bins of orientation and the length is the sum of the gradient magnitude. For each pixel in template T, a sub-template T' is extracted with the same size with image I and its descriptor is calculated with the same way with image I. Then the distance between image Iand sub-template T' is calculated. The last location of estimated landmarks is the location that has the smallest measure distance value. Finally, the automatic landmarks are inverted to match with the real location in the scene image (by inverting the transformation).



Figure 5: Automatic landmarks after refinement

Figure 5 show a complete result on one scene mandible with the manual landmarks (red points) and estimated landmarks (yellow points).

# 3 EXPERIMENTS AND RESULT

All the steps in our method are implemented in MAE-Lab<sup>1</sup>. Two sets of beetle have been analyzed, right and left mandible. After verifying the quality of the image, it remains 288 usable images for right mandible and 285 images for left mandible. The removed images include the images that do not contain the mandible and the mandible is broken.

In all valid images, a set of 18 manual landmarks of right mandible (16 landmarks for left mandible) are indicated by biologists. Along with choosing the centroid size to measure the mandible. This size is obtained by sum of all square distance from each landmarks to the centroid point (see [3]).

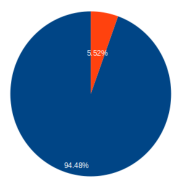


Figure 6: The percentage of correct proportions on right mandibles

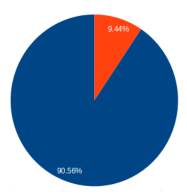


Figure 7: The percentage of correct proportions on left mandibles

By this way, we have compared the centroid size between manual landmarks and estimated landmarks. We can see in figure 6 and figure 7 for all images. The correct proportions of estimated landmarks are 94,48% for the right mandible and 90,56% for the left mandible. In

fact, this rate is evaluated by choosing an arbitrary image as the model. The object in the image is presented with difference size (scale problem) that our method just consider on translation and rotation between two images.

Besides, using the centroid size to evaluate the method. We also consider the position of the estimated landmarks. In this experiment way, we calculate the distance between each manual landmark and corresponding automatic landmark. The error rate of each landmarks are shown in the table. From two experiment ways, we can see that the method is success in indicating all landmarks for each image; and the location of the landmarks is considered near with the manual landmarks in some aspect.

In a different side, when comparing with our previous study (see in [4]). This method has more exactly about the position of automatic landmarks as well as the advantages for the implementation process. The memory to detecting the landmarks, along with the times to execute the process are decreased dramatically.

## 4 CONCLUSION

Morphometric analysis is a powerful tool in biology in classification the species. Automatic identification the characteristics biology of the organism is a difficult problem. In the content of this paper, we have begun to design a method to segment the beetle mandibles and to indicate automatically landmarks which have been determined by biologists. Each mandible is segmented by applying the Canny algorithm. Using GHT and image registration to estimate the landmarks. Finally, cross-correlation will be applied to refine the location of the estimated landmarks. The first version of this method has been implemented. From now, the next stage of our method is to add the features to have the position of landmarks more precisely.

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<sup>&</sup>lt;sup>1</sup> MAELab is a free software in C++. It can be directly obtained by request the authors.

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