

DETERMINING THE ABNORMALITY OF BULL SPERM TAIL MORPHOLOGY USING SUPPORT VECTOR

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Abstrak

Penilaian atas ketidaknormalan spermatozoa bisa dilakukan dari sisi motilitas maupun morfologi (kepala dan ekor). Penelitian ini mengevaluasi ketidaknormalan spermatozoa dari sisi morfologi bagian ekor spermatozoa sapi. Data berupa 50 citra mikroskopis spermatozoa yang diperoleh dari Loka Penelitian Sapi Potong Grati, Pasuruan digunakan dalam penelitian ini. Prosedur yang ditetapkan terdiri atas beberapa tahap. Tahap pertama adalah melakukan segmentasi spermatozoa untuk memisahkan spermatozoa dari latar belakang dan memisahkan bagian ekor spermatozoa dari bagian yang lain. Selanjutnya dari hasil segmentasi dicari garis tengah ekor (*skeleton*) menggunakan metode medial axis transform. Berdasarkan garis tengah yang dihasilkan, dilakukan prosedur ekstraksi fitur menggunakan metode *polynomial curve fitting*. Kemudian, metode *Support Vector Machine* (SVM) digunakan untuk menentukan ketidaknormalan bentuk ekor spermatozoa. Untuk pembelajaran digunakan 25 data spermatozoa normal dan 10 data spermatozoa tidak normal. *Testing* kemudian dilakukan atas 15 data spermatozoa tersisa. Ketelitian SVM dalam menentukan ketidaknormalan bentuk ekor spermatozoa mencapai 73.33%. Dengan demikian ketidaknormalan bentuk ekor spermatozoa dapat ditentukan dengan menggunakan SVM.

Kata kunci: Ekor Sperma sapi, *Morphology*, *Polynomial Curve Fitting*, SVM.

Abstract

Determinining the abnormality of spermatozoa can be done by inspecting its motility or morphology (head or tail). This study examined 50 data of sperm microscopic images. The semen was obtained from Loka Penelitian Sapi Potong Grati, Pasuruan. A sequence of procedure consist of several steps were then carried out. The first step was to obtain sperm tails by segmenting the sperms from its background and removing the heads and the necks parts. The skeletons of the tails were then obtained using a method of medial axis transform. The features of the tails were then extracted using polynomial curve fitting. Then, Support Vector Machine (SVM) was used as a classifier. In the training phase, 25 normal sperm and 10 abnormal sperm were utilized. Afterward, the remaining 15 data were used in the testing phase. The accuracy of SVM was 73.33%. Hence, the abnormality of spermatozoa based on the shape of sperm tail can be determined using SVM.

Key words: Bull Sperm Tail, Morphology, Polynomial Curve Fitting, SVM.

INTRODUCTION

Analysis of sperm shape has been done on either on human or animal [1-3]. However, assessing the abnormality of bull sperm especially based on its morphological condition is still crucial issue since morphological condition related to the quality of bull sperm especially for artificial insemination program for both beef and dairy industry [4,5]. The quality of sperm could be determined by several criterions, such as the size and the form (morphology) of the head, the length and the form of the tail and its movement behaviour (motility) [6,7]. Sperm or also called Spermatozoa are cells of the male reproductive system. Spermatozoa are highly specialized cells and dense that no longer have cleavage or growth, derived from the gonosit into spermatogonia, primary and secondary spermatocytes, subsequently changed to be spermatids and finally turn into spermatozoa. Spermatozoa consists of two functionally important parts, the head and tail [8]. In the normal adult male, spermatogenesis continues throughout life, although the quality and quantity decline with age.

The head of spermatozoa is oval with approximately 5 microns in length, 3 microns in diameter, 2 microns in thickness and mainly formed by the nucleus. Spermatozoa tail section consists of neck, middle, principal, and the end piece. The tail length is about 55 microns and 1 micron thickness. The length of principal piece 45 microns, 0.5 microns thick and the tip of 4-5 microns long, 0.3 microns thick. The tail could not be observed using light microscope but electron microscope should be used [9]. Sperm that has abnormal shape loss its ability to fertilize ovum in the fallopian tubes.

Figure 1 shows the normal bull sperm morphology that has a normal head, midpiece and tail. The figure was derived from one of the images used in this study.

Previous studies in evaluating bull sperm morphology have been done by some researchers in the past using two main methods [10], the first is bright-field (BF) microscopy method and differential interference phase contrast (DIC) microscopy method. It was reported that there was no difference in percentage of normal sperm that

can be seen between each method, however DIC was more effective in visualizing main defect. Moreover, BF is considered to make the sperm looks smaller. From the result point of view, DIC is recognized as a better viewing tool in assessment of bull sperm morphology compared to BF method [11].

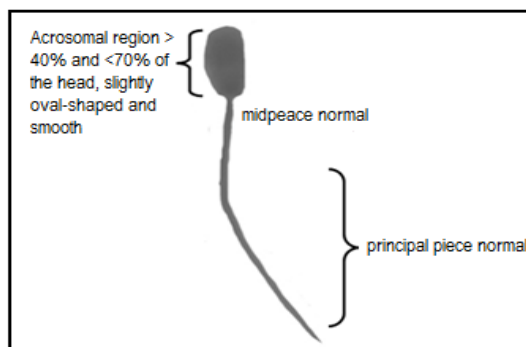


Figure 1. Normal Sperm Morphology.

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In another research, the identification and classification of spermatozoa has been performed by experts on different sperm samples such as human, horses, pigs and bull spermatozoa [12]. Ahmad proposed in their study that the identification of human sperm tail can be done through detecting shapes using Structural Similarity Index (SSIM) and Local Entropy. The advantage of this method was being able to detect the sperm tail at low contrast but the disadvantage was it required a long execution time [13].

This study proposed automatic SVM classification of bull sperm tail abnormalities based on polynomial curve fitting feature extraction.

MATERIAL AND METHODS

Data Acquisition

CASA (Computer Assisted Semen Analysis) as one of the key player device in measuring morphological condition of sperm morphologically. Casa provides many advantages to the researchers but also presents some drawbacks such as high device installation cost and need to be stationed at a fix position like in laboratory. Developing a low cost device and mobile could be a breakthrough in solving some limitations of CASA such as for gathering and analysing data from many beef or dairy industries in remote areas. In this study we implemented our own tool for observing and analysing bull sperm tail based on its morphological condition. The tool was implemented using a microscope equipped with CCD camera (Sony IMX035 CMOS, model: FL3-U3-13S2C-CS) connected to a computer via usb. Using this tool images of sperm samples prepared on slides were taken. The sperm samples were prepared by dropping sperm from straw onto slides, colored using eosin-negrosin agents to distinguish the dead from the live, smear carefully and evenly over the slides. The sperm in straw packaging was obtained from Loka Penelitian Sapi Potong, Grati, Pasuruan, East Java, under the auspices of the Ministry of Agriculture, Republic of Indonesia. Fifty images were obtained in this study. Four out of the 50 were shown in Figure 2 as an example.

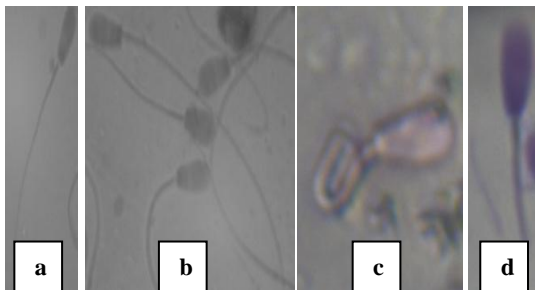


Figure 2. Example of bull sperm images (a) and (b) normal spermatozoa , (c) and (d) abnormal spermatozoa.

Support Vector Machine

Support Vector Machine (SVM) is a famous method for classification since it can find an exact classification boundary using a so-

called support vectors, hyperplanes and kernels. Numerous applications of SVM have been performed by several researchers such as in remote sensing application [14] and medical application [15]. In this research we apply one type of SVM, linear SVM [16], to classify the sperm abnormality based on the shape of sperm's tail.

Work flow

The fifty images obtained were processed. The process followed the steps as depicted in Figure 3. Manual part of Figure 3 were to separate sperm from the background (as in example of Figure 4) and to crop the tail (as in Figure 5).

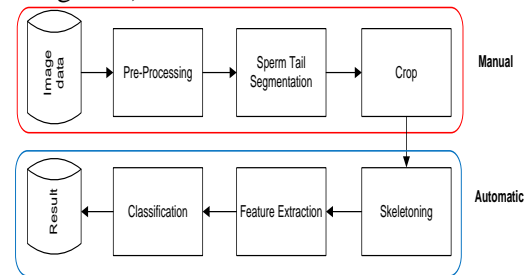


Figure 3. Flow Diagram of the Research Methodology.

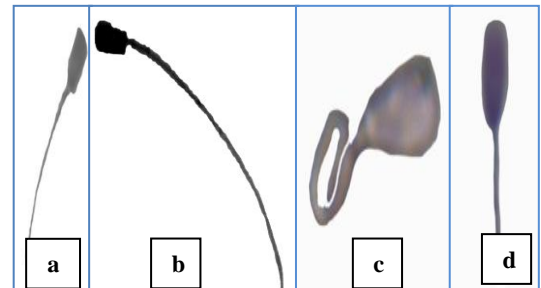


Figure 4: Example of Bull Sperm Segmentation Image; (a) and (b) Normal Spermatozoa; (c) and (d) Abnormal Spermatozoa.

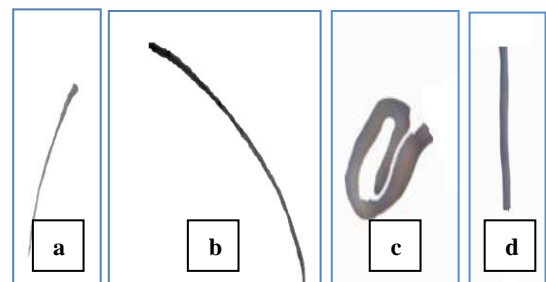


Figure 5: Example of Bull Sperm Tail; (a) and (b) Normal Spermatozoa, (c) and (d) Abnormal Spermatozoa.

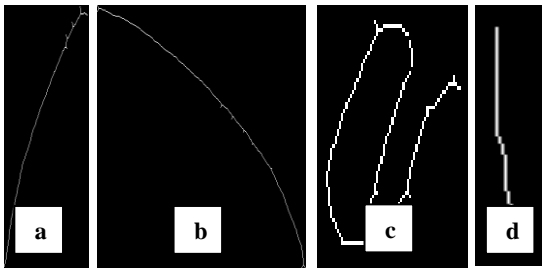


Figure 6: Example of Bull Sperm Tail Skeleton Image; (a) and (b) Normal Spermatozoa; (c) and (d) Abnormal Spermatozoa.

The automatic part of Figure 3 starts with using medial axis transform method to get the skeleton of the segmented tails as shown in Figure 6. Features were extracted from the skeletons using polynomial curve fitting. The skeleton were fit using a polynomial of Equation (1) with $n=4$:

$$p(x) = p_1x^n + p_2x^{n-1} + \dots + p_nx + p_{n-1} \quad (1)$$

Then the sum of the distance between each pixel of the skeleton to the corresponding pixel of the obtained polynomial, known as norm of residual, were calculated using Equation (2).

$$norm(d,2) = \sum(abs(d).^2)^{\frac{1}{2}} = \sqrt{\sum_{i=1}^n d_i^2} \quad (2)$$

The norm of residual together with the curve length constitutes the features extracted from the data in this study.

The next step were classification of the features obtained above using support vector machine. SVM implementation in this study uses linier function. For example: $\{x_1, \dots, x_n\}$ is dataset and $y_i \in \{+1, -1\}$ is class label from x_i data. The two classes are separated by pair of field barrier. Field barrier of class 1 is shown in Equation (3) and class 2 in Equation (4).

$$x_i.w + b \leq \pm 1 \text{ for } y_i = -1 \quad (3)$$

$$x_i.w + b \geq \pm 1 \text{ for } y_i = +1 \quad (4)$$

$$\frac{2}{w} = \frac{1-b - (-1-b)}{w} \quad (5)$$

where w is normal field and b is relative position to the centre of coordinate. Whereas margin value is distance between field barrier of class 1 and class 2 that is shown in Equation (5).

RESULT AND DISCUSSION

For the purpose of training of the linear SVM classifier, 35 data were selected. Twenty five out of the 35 known to be normal data (shown in Table 1) while 10 out of the 35 known to be abnormal data (shown in Tabel 2).

Table 1. Twenty five Training Normal Sperm Data (in pixel x 10^2).

#	Curve path length	Norm of residual
1	5.50	1.81
2	4.79	1.68
3	5.59	1.49
4	3.90	2.67
5	3.91	4.41
6	4.63	1.02
7	2.46	1.05
8	4.31	1.18
9	3.32	2.66
10	3.58	5.51
11	3.35	6.22
12	3.86	5.33
13	5.50	1.82
14	2.89	1.64
15	4.19	4.28
16	2.76	1.30
17	5.69	9.72
18	4.55	1.36
19	3.19	2.12
20	4.63	1.25
21	2.43	1.23
22	4.57	4.61
23	3.62	6.65
24	4.90	5.15
25	3.57	1.67

Table 2. Ten training Abnormal sperm data (in pixel x 10^2).

#	Curve Path Length	Norm Residual
1	6.30	4.80
2	2.61	3.25
3	1.85	8.80
4	4.14	9.06
5	9.30	7.09
6	2.89	7.13
7	9.00	1.25
8	1.37	0.90
9	8.50	2.95
10	6.30	7.34

The rest 15 data, used as the test data set, were then classified using the support vector to be normal (+1) or abnormal (-1). The result of SVM classification is shown in column 4 of Table 3. Visual classification is also shown in column 5 for comparison.

Table 3. Comparison of SVM and Visual Classification.

#	Curve Path Length	Norm residual	SVM classification	Visual classification
1	98	107	-1	-1
2	496	546	1	1
3	519	102	1	1
4	217	273	-1*	1
5	595	35.8	1	1
6	346	402	1	1
7	328	899	-1	-1
8	255	417	-1*	1
9	368	984	-1	-1
10	404	711	1*	-1
11	313	667	-1	-1
12	103	27.1	-1	-1
13	111	19.2	-1	-1
14	247	183	1*	-1
15	84.0	73.1	-1	-1

From Table 3, four out of 15 samples were classified incorrectly. Those are samples number 4, 8, 10, and 14. This gave accuracy of 73.33 %.

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CONCLUSSION

This classification result were confirmed by researchers at Loka Penelitian Sapi Potong Grati, Pasuruan.

In conclusion, applying SVM for determining the abnormalities of bull sperm based on tail morphology in this study has shown good results. Simplicity of the classification process both training data and testing data with SVM provided some advantages in terms of design and speed.

However, evaluation and improvement of the method need to be carried out immediately for future work to get optimum results such as considering to use more training data. In addition, the use of non-linear kernel SVM needs to be tested as a comparison method of linear SVM. Further research can be performed by adding more classes in the classification process rather than only two classes, normal and abnormal

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