Measurements of Body Movement in Chick Embryos During Early Stages

Kenji Moriya* and Yuya Chiba

* Department of Production Systems Engineering, National Institute of Technology, Hakodate College.
Tokura-cho 14-1, Hakodate City, Hokkaido, Japan 042-8501
b Support Center for Engineering Education, National Institute of Technology, Hakodate College.
Tokura-cho 14-1, Hakodate City, Hokkaido, Japan 042-8501

* Corresponding author: moriya@hakodate-ct.ac.jp

Abstract

In this study, we investigate developmental patterns of body movements that may play roles in normal embryonic growth using direct recordings of chick embryos during the early stages of development (72–125 h of incubation). In three individuals, embryonic body movements occurred intermittently and irregularly after 72-h incubation, and frequencies and sizes of movements increased with embryonic growth. Additionally, patterns of the body movements became periodic at 85 h in all individuals, and cyclic periods of body movements were then shortened and periodicities of movements gradually disappeared with embryonic development. If cyclic body movements are a common requirement during normal chick embryo growth, continuous monitoring of body movements could be used to predict abnormal embryonic growth.

Future studies are required to investigate distinctive body movements and developmental patterns during embryonic growth in disease models and under various environmental conditions, such as under hypoxia.

Keywords: Early stages embryogenesis, Embryonic body movement, Template matching.

1. Introduction

1.1 Chick embryo as models of fetal development

Because chick embryos have all the nutrition necessary for growth inside their shells, the external conditions required for development are merely temperature and oxygen, leading to multiple advantages for studies of fetal development. Among these, embryonic physiological parameters that are not influenced by maternal health conditions can be measured, and assessments of responses to changes in temperature, humidity, and oxygen concentrations can be made, as shown previously (1-3). Fertile chicken eggs can also be easily obtained from chicken farmers and physiological parameters in chick embryos have been considered suitable and independent indicators of embryonic development at all incubation periods, particularly in studies of cardiovascular system development (4-9). Although measurements of bio-signals during the early stages of pregnancy in mammals are extremely difficult, those of early stages (until day-7) in chick embryos can be achieved with relative ease. Hence, as models of fetal development, chick embryos offer significant advantages.

1.2 Investigation of body movement patterns in chick embryos during the early stages of development

It is well established that heart tissues in chick embryos are formed after about 30 h incubation, and that other organs, such as chorioallantoic membranes, are under development at this early stage. However, no continuous and direct in ovo video recordings of chick embryos have been published previously. In a previous study, we developed a system for continuously recording chick embryo developments over long periods using a small Charge Coupled Device (CCD) camera mounted on the egg, and investigated physiological developments of chick embryos, including those of heart structures and blood vessels, with minimal or no sacrifice. Although cardiac signals are barely detectable using electrocardiograms during early stages of chick embryogenesis, we determined instantaneous heart rates during the early stages using video images that were directly recorded using our system (10-12).
A recent study indicated that mortality rates of human fetuses are greater among those with significantly reduced body movements (13). Therefore, we developed a system for assessing embryonic body movements from continuous and direct recordings of chick embryos, and investigated developments of body movement patterns during the early stages of chick embryogenesis.

2. Experimental methods

2.1 Direct recording of embryonic body movements

A schematic and a photograph of the body movement recording system are presented in Fig. 1. Because the details of the direct recording system are described in a previous study (10–12), we describe the system only briefly herein. To mimic general incubation conditions of eggs, incubation and recording procedures were performed in a constant temperature incubation at approximately 38 °C with 50% humidity. Holes of approximately 1.5 cm in diameter were made on the tops of egg shells (air-cell side), and a CCD camera (Type MTV-5366ND, Akiduki Denshi Tsusho Co. Inc., effective pixels: 352 × 240) was attached with a connecting plastic tube. Because defective seals between egg shells and the spacer for the CCD camera might cause significant physiological and/or biological adversities during embryonic development, sealing was performed very carefully. Additionally, adequate circulation of air was required to promote heat distribution to the CCD camera, and this was achieved using forced-draft incubation during experiments. Because previous studies report body movements of chick embryos starting from around 80 h of incubation, we started continuous monitoring experiments at 72 h (day-3) incubation. Video images were captured at 30 frames/s (i.e., with a sampling frequency of 30 Hz) using a CCD camera with a video capture board attached to a computer via a coaxial cable. Images were saved for analyses in MPEG format. Although these procedures were invasive, previous experiments showed successful hatching at normal incubation times after closure of the hole in the top of the egg using a nylon-wrap to prevent evaporation (unpublish data).

2.2 Quantitative determinations of body movements

We developed a system for measuring sizes and frequencies of body movements using image processing tools. During the early stages of embryogenesis, whole embryos did not move, and embryonic movements were initially observed as those of the embryo's head. Thus, we calculated distances between embryo head positions as an indicator of body movements. Because body movement patterns develop with embryonic growth, those in the first 5 min of every 1 h of incubation were investigated. In the first 5 min of 1-h incubation time points, images were extracted after continuous video recording and were divided into 1-h segments. Locations of embryo heads were calculated using the template-matching method in the Open Source Computer Vision library (Open CV), which was provided by Intel Corporation. Differences in embryo locations between one frame and the next (distances of movements) were calculated as body movements per frame. This procedure was repeated for 5 min (9000 frames) and data strings of the coordinates of embryos were recorded.

Figure 2 shows an example of the recorded image and the results of template matching following use of the head area for the template image at 85 h of embryo incubation. Whole embryos and blood vessels are indicated by the yellow dotted lines and black arrows in Fig. 2A, respectively. The red rectangles indicate areas of the template image in which correct and incorrect matching was achieved. The embryo's head position was defined as the embryo's position, and was successfully detected (Fig. 2B). However, because template matching occasionally failed (Fig. 2C), incorrect detection data were removed using a digital low-pass filter (FIR filter, cut-off frequency of 2 Hz), as described previously (14). Calculation procedures for distances of body movements are summarized in Fig. 3.
To investigate periodicity of body movements, power spectrum analyses were performed using discrete Fourier transform after applying a Hamming window \(^{15, 16}\). Analyses were performed over 5 min and the data were filtered with a cut-off frequency of 2 Hz, corresponding with an effective frequency component from $3.3 \times 10^{-3}$ to 2.0 Hz.

### 3. Results

#### 3.1 Development of embryonic body movements

Although images were recorded for eight embryos, only those for three individuals were suitable for continuous analyses of locations, angles, and measurement foci, and body movements of these three embryos were calculated using the abovementioned image processing method.

Figures 4, 5, and 6 show recorded embryos and indicate body movements over the first 5 min of each incubation time point. These data were used to characterize changes in body movement patterns. Images on the left side show chick embryos at each incubation time, and middle and right panels show relative distances (in pixels) of movements on X- and Y-axes and two-dimensional orbits of body movements during 5-min periods at each incubation time point. The scales of vertical axes varied according to magnitudes of body movements. Because embryos were rotated to adjust the focus of the camera, the X–Y direction is given as a relative coordinate, and hence, X and Y axes in the middle panels show directional characteristics of body movements.

As shown in Fig. 4, body movements of embryo A were not observed in X- or Y-axes at 72 h after the start of incubation, but these movements gradually began at 85 h after the start of incubation, albeit with limited sizes of movements. In addition, an in-phase periodicity was observed in movements on both X- and Y-axes. Frequencies of body movements did not change significantly at 100 h after the start of incubation, whereas cycles per body movement were approximately three times shorter.
Furthermore, at 110 h after the start of incubation, sizes of body movements decreased and clear periodicity disappeared. However, at 125 h after the start of incubation, movements became energetic and complex, and based on X–Y trajectories, these movements produced a horizontal circular pattern.

In embryo B (Fig. 5), no body movements were observed at 72 h after the start of incubation. Moreover, periodic movements began at 85 h after the start of incubation, as in Embryo A. However, frequencies of body movements in embryo B gradually increased thereafter and became complex with incubation time and embryonic development.

Body movements of embryo C had already started at 72 h after the start of incubation, and both periodicities and sizes of body movements were increased by 85 h after the start of incubation (Fig. 6). At this time, movements of embryo C showed significant periodicity and at around 100 h of incubation, body movements had no apparent directions and became complex and large. Although the

Fig. 4. Recorded images (left panels) and sizes of body movements (pixels) on X- and Y-axes (middle panel), and in the X–Y (right panel) direction during 5-min time points at 72, 85, 100, 110, and 125 h, respectively, in embryo A. Total relative body movements were determined and are expressed in pixels.

Fig. 5. Images of embryo B (left panels) and volumes of body movements on X- and Y-axes (middle panels), and in the X–Y (right panels) direction.

Fig. 6. The recorded images (left panels) and body movements of embryo B on X- and Y-axes (middle panels), and in the X–Y (right panels) direction. Body movements at 125 h of incubation (*1) could not be measured because embryo B was obscured by a newly developed chorioallantoic membrane.
frequencies of movements of embryo C were augmented at 125 h of incubation, determinations of the embryo positions using template matching failed thereafter because a chorioallantoic membrane grew between the CCD camera and the embryo. These observations indicate that the present template-matching method is suitable for embryos until around the 125th hour.

3.2 Periodicity of body movements

Although body movements of embryos A and B at around 72 h were transient and irregular, periodic movements were observed at around 85 h in all individuals, and these were pronounced on the Y-axis (Fig. 6) and subsequently became continuous and random. Thus, periodicity of body movements in these individuals was investigated using discrete Fourier transform (DFT) analyses (Fig. 7). At 85 h after the start of incubation, periodic peaks were approximately 148 (Embryo A), 99 (Embryo B), and 37 s (Embryo C) apart, respectively. Although the frequency of movements was shorter for embryo C than for the other individuals at the same incubation time, peaks of movements were approximately 74 s apart at 72 h, similar to those of embryo B at 85 h after the start of incubation. All embryos had cyclical body movements at the 100th hour of incubation. However, these periods were shorter than those at 85 h, and ranged from 148 to 8 s in embryo A, from 99 to 15 s in embryo B, and from 37 to 6 s in embryo C, respectively. At the 110 h of incubation, movements followed a periodicity of approximately 17 s in embryo B and 5.4 s in embryo C, and had disappeared in embryo A. At the 125th hour of incubation, periodicity of body movements remained only in embryo B, although body movements of embryo C were not visible at this time point. These results (Fig. 7) indicate that cyclic periods between body movements gradually become shorter, and that periodicity is gradually replaced by irregular body movements.

4. Discussions

4.1 Direct recording system and calculations of embryonic body movements

Herein, we directly recorded embryonic development using a CCD camera, and calculated frequencies and periodicities of chick embryo body movements during the early stages of development from 72 to 125 h of incubation. The present direct recording system was developed and calculations were performed using template matching to determine distances of embryonic body movements. Although we tried to record body movements and calculate distances for eight individuals, data were only successfully generated for three embryos. Whereas embryo positions were detected incorrectly on occasion due to miss-template matching, these data were excluded using a low-pass filter during subsequent digital processing procedures (Fig. 2C). Among difficulties faced during these experiments, embryos are relatively transparent apart from the heart and blood vessels (left-images in Fig. 4−6). Thus, the edges of embryo heads that moved well were distinguished using a template from background images of the amnion. Moreover, to reduce failures of template matching, an image processing program was used to distinguish embryos and to emphasize their edges. In particular, we distinguished the head and/or eyes using a differential filter. Focus adjustments of the CCD camera were required to improve image sharpness.

This system for measuring embryonic body movements was effective until about 125 h (5 days) of incubation. In contrast, measurements of body movements
were previously achieved using a laser or a record needle in contact with the egg shell, and this system detected body movements from around 7 days of incubation \(^{(17, 18)}\). Thus, with the use of the present and previously described measurement systems, body movements can be investigated during all incubation periods.

4.2 Body movement patterns in chick embryos at early stages of development

Embryos commonly began moving with floating fluctuations after 72 h of incubation, and we assumed that this is common to all normally growing embryos. Subsequently, intermittent and irregular body movement patterns became cyclic and had circular orbits after 85 h of incubation, but then became irregular with increases in volumes of movement. Although the observed cyclic body movements had periods of 100–150 s at the 85 h incubation time point, these cyclic periods became shorter and then disappeared during further embryonic development until the 125th hour. Although the causes of cyclic movements remain speculative in the absence of developed physiological functions, they may relate to the development of blood vessels, the lack of oxygen, or forces from the heart beat. Because these circular movements were observed in all of the present embryos, they may be necessary for normal growth and could be a significant milestone during embryonic development. However, further investigations are required to determine the presence of these cyclic body movements in abnormal embryos, and to confirm that they are common to all normally growing embryos.

Finally, the cyclic periods (approximately 74 s) of body movements in embryo C at 72 h corresponded with those of embryo B at 85 h (approximately 99 s). Moreover, cyclic periods at other incubation time points were shorter for embryo C than for embryo B, and the body size of embryo C at 72 h was similar to that of embryo B at 85 h. These observations suggest that that embryo C developed earlier than embryo B, and that developmental patterns of body movement could be used to identify normal growth or delayed development.

Mortality rates of human fetuses are reportedly increased among embryos with limited body movements \(^{(13)}\). Hence, the present system for continuous monitoring of body movements in chick embryos may offer a tool for diagnosing abnormal growth patterns. However, future studies are required to correlate distinctive body movement patterns and developmental patterns with embryonic growth in disease models and under conditions of privation, such as hypoxia.

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**Additional statement**

This research was approved by the Life Ethics Committee of NIT(KOSEN), Hakodate College.

**References**


