Culture and Identification of Goat Synovial Membrane Cells for Caprine Arthritis Encephalitis Virus Diagnosis

Niron Ratanapob and Theera Rukkwamsuk
Department of Large Animal and Wildlife Clinical Sciences,
Faculty of Veterinary Medicine, Kasetsart University, Kamphaengsaen,
73140 Nakhonpathom, Thailand

Abstract: Caprine Arthritis Encephalitis Virus (CAEV) is an important pathogen in goats. Definitive diagnosis depends on viral isolation and this method is beneficial for studying CAEV infection. Synovial fibroblast is the most appropriate cell for CAEV isolation. The objectives of this study were to culture synovial membrane cells and to identify whether the cultured cells were fibroblast. Synovial membrane tissue was collected from a CAEV-seronegative, 5 months old, male goat. The tissue was digested by collagenase. Synovial cells were cultured in RPMI 1640 at 37°C and 5% carbon dioxide. The cultured synovial cells were detected for vimentin, a marker for fibroblastic cell type, using immunocytochemistry. Phagocytic activity of the synovial cells was also tested by adding carbon particle in culture media to confirm whether or not the cells were synovial macrophages. Results revealed that the cultured synovial cells were spindle-shaped and were attached to the bottom of the flasks. These primary cultured cells could be subcultured >9 passages. Vimentin could be detected but phagocytic activity could not be detected in these cultured synovial cells indicating that these cultured cells were fibroblast.

Key words: Caprine arthritis encephalitis virus, cell culture, goat, synovial fibroblast, vimentin

INTRODUCTION

Caprine Arthritis Encephalitis Virus (CAEV), a member of retrovirus family causes a life-long infection and economical losses in goat industry (Smith and Cutlip, 1988). Infected goats frequently show asymptomatic but they can transmit the virus for life via colostrum and milk including secretion containing white blood cells (Ellis et al., 1983). Recent study in the central and Western part of Thailand found that the individual and herd seroprevalence of CAEV infection in goats were 12.40 and 47%, respectively. Therefore prevention and control measures should be taken to inhibit spreading of infection in Thailand (Ratanapob et al., 2009). Serological methods are convenient for CAEV diagnosis however, viral isolation is a definitive diagnosis of CAEV infection (Narayan and Cork, 1990). In addition, viral isolation is beneficial for further studies on CAEV pathogenesis providing more essential details and information to proper control strategy of the infection. The virus can be isolated by infected tissue explantation (Coakley et al., 1981) or co-cultivation of virus infected cells with susceptible cells (Narayan et al., 1983). Infected culture cells show syncytial formation which is typically cytopathic effect due to CAEV infection (Coakley et al., 1981). In laboratory, fibroblastic type of synovial cell is the most appropriate cell for CAEV isolation (Knowles et al., 1992). Preparation of the cells from synovial tissue and identification of the fibroblastic cell type are essential parts of the synovial membrane cell culture before the cultured cells could be used further. Therefore, the objectives of this study were to culture synovial membrane cells and to identify whether the cultured cells were fibroblast.

MATERIALS AND METHODS

Synovial membrane cell culture: Method for cell culture was described by Upagarin (2005). Basically, synovial tissues were obtained from elbow and stifle joints of a 5 months old, male goat. The goat serum was tested negative for antibodies against CAEV infection by ELISA (CHEKIT-CAEV/MVV, IDEXX Laboratories Inc., Maine, USA). The obtained synovial tissues were digested by collagenase type 1 (Gibco™ BRL, GIBCO Company, Ltd. Bangkok, Thailand) in RPMI 1640 (Gibco™ BRL) with 3% Fetal Calf Serum (FCS) at 37°C for 1 h. The digested product was centrifuged at 1,500 rpm for 5 min at 4°C. The

Corresponding Author: Theera Rukkwamsuk, Department of Large Animal and Wildlife Clinical Sciences,
Faculty of Veterinary Medicine, Kasetsart University, Kamphaengsaen, 73140 Nakhonpathom, Thailand
1827
sediment was washed with phosphate buffer saline prior to be centrifuged at 1,500 rpm for 5 min at 4°C. The synovial cells were cultured in RPMI 1640 with glutamine (2.05 mM), penicillin (100 unit mL⁻¹), streptomycin (100 µg mL⁻¹), 2 mM sodium pyruvate, 50 µM 2-mercaptoethanol and 10% FCS at 37°C and 5% carbon dioxide. The cells were looked over under light microscope and when reached confluence were subcultured approximately twice weekly.

**Cell type identification**

**Fibroblastic cell:** The ninth passage of the cultured synovial cells was fixed on slides with 10% formalin and was incubated at 60°C for 1 h. Vimentin, an intermediate filament family of protein in fibroblastic cells was detected by immunocytochemistry with anti-vimentin antibody at The Laboratory of Anatomical Pathology, Bangkok Metropolitan Administration Medical College and Vajira Hospital. Cancer cells positive to anti-vimentin were used as positive control. The slides were examined under light microscope.

**Phagocytic activity:** The synovial cells were tested for phagocytic activity as described by Upragaran 2005. Carbon particle was added in the culture media at a rate of 50 µg mL⁻¹ and was incubated with the cultured cells at 40°C for 30 min. Then the media was discarded and the cells were washed 5 times with phosphate buffer saline solution. The cultured cell was looked over under light microscope.

**RESULTS AND DISCUSSION**

A number of synovial cells obtained from digestion of synovial tissues with collagenase. The cells were spindle-shaped and were able to attach to the bottom of the flasks as monolayer (Fig. 1). The cultured synovial cells reached the confluence every 7-10 days and could be subcultured >9 times. In present study, some steps for synovial cells culture were different from other studies. Some researchers used explantation of synovial tissues in which the cells would grow out from synovial tissues until monolayer was formed (Crawford et al., 1980; Narayan et al., 1980; Klevjer-Anderson and Cheevers, 1981; Rolland et al., 2004). While enzymatic digestion was used in present study and the cultured cells could grow as well. Moreover, the study used the synovial cells harvested from a 5 months old male goat instead of embryo, fetus or colostrums-deprived kid which ensured that they were free from CAEV infection (Crawford et al., 1980; Narayan et al., 1980; Klevjer-Anderson and Cheevers, 1981; Gjerset et al., 2007). However, the goat used for cells donor was tested seronegative for CAEV infection before the cells were further processed. In general, cells from embryo, fetus or young animals have higher survival and growing rate than cells from adult animals because they have low specificity and high proliferation rate (Freshney, 2000). Nevertheless, the synovial cells from the adult goat in the present study could also grow well. An important consideration in synovial cell culture was contamination which might be occurred during tissue collection. The contaminating microbes could not be sensitive to antimicrobial drugs used in culture media. One of the most common contaminating microbes is **Mycoplasma** sp. which can cause arthritis in goats and can be isolated from synovial fluid (Singh et al., 2004). Therefore, aseptic technique and antimicrobial agent are required for tissue collection and culture.

Immunocytochemistry for detection of vimentin was applied to scrutinize the fibroblastic type of the cultured cells in this study. When vimentin is presented, the cells were brown-stained in the cytoplasm (the positive control) whereas the negative control cells were not stained (Fig. 2). This result confirmed that the cultured synovial cells were actually fibroblasts. Besides vimentin, propyl 4-hydroxylase, vascular cell adhesion molecule-1, CD44, CD55, CD90 and cadherin-11 can be used as synovial fibroblast markers (Toke et al., 2002; Rosengren et al., 2007).

In general, synovial membrane cells consisted of synovial macrophages and synovial fibroblasts. In this study, phagocytotic activity, detected by measuring an ability of the cultured cell to ingest carbon particles was used to clarify whether or not cultured cells were synovial macrophages. The result revealed that the cultured synovial cells did not have phagocytotic activity. Therefore, synovial cells from the present study are fibroblast because they were positive for vimentin detection and did
ACKNOWLEDGEMENTS

This study was financially supported by the National Research Council of Thailand and Kasetsart University Research and Development Institute. The researchers owed their deepest gratitude to expert technicians of Department of Anatomical Pathology, Bangkok Metropolitan Administration Medical College and Vajira Hospital for their assistance on immunocytochemistry for vimentin detection. Assistant Prof. Dr. Narin Upragarin and Mrs. Wilairat Chansing are thanked for their technical assistance in culture of the synovial cells.

REFERENCES


CONCLUSION

Synovial membrane cells could be obtained from digestion of synovial tissue from a CAEV-seronegative goat with collagenase. The cells could grow in culture media at 37°C and 5% carbon dioxide >9 passages and were confirmed as fibroblastic type by determination of vimentin and phagocytic activity.

Fig. 2: Results of immunocytochemistry for vimentin detection; a) positive control (cancer cells) with brown-stained (100x); b) negative control (goat synovial cells were tested without anti-vimentin) (200x) and c) goat synovial cells positive to anti-vimentin (200x) with brown-stained not have phagocytic activity. Prior to further use of these cultured synovial cells, susceptibility of the cultured cells to CAEV must be tested. Furthermore, other viruses causing pathological changes in the joints should also be used to test susceptibility of these cultured synovial cells.


