EVALUATION OF A SIMPLE *Theileria annulata* CULTURE PROTOCOL FROM EXPERIMENTALLY INFECTED BOVINE WHOLE BLOOD

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**Summary:**
We have evaluated a new simple technique using whole blood from experimentally infected cattle for the isolation and cultivation of *Theileria annulata*. The study was carried out on 20 Holstein-Frisian bovines that had been experimentally infected with a virulent lethal dose of *Theileria annulata*. This technique has been compared to the classical peripheral blood monocyte isolation with Ficoll carried out on 22 experimentally infected Holstein-Friesian calves. The effectiveness of the reference technique was estimated to 86.4 %, whilst the effectiveness of the new technique was 100 %. Moreover, this new technique leads to time and money saving estimated to € 3.06 per sample. It decreases the contamination risks by reducing the steps of sample manipulation.

**KEY WORDS:** *Theileria annulata*, cell culture, blood, Ficoll.

**Material and Methods**

In order to compare the two techniques we used two groups of animals. The animals were purchased from a tropical theileriosis and *Hyalomma detritum detritum* free farm and they were kept in a tick-free isolation unit at the national veterinary school at Sidi Thabet:

- Group 1: 20 Holstein-Friesian bovines aged between five and ten months.
- Group 2: 22 Holstein Friesian calves aged between three and six months.

The animals were infected by subcutaneous injection of a lethal dose of 1.8 and 0.25 equivalent ticks using ground up tick supernates (GUTS) of the Jdaida 4 (Ta3/2 isolate), respectively.

Ficoll (Histopaque© 1077, Sigma) was used for the isolation of peripheral blood monocytes (PBM), as recommended by the supplier. This technique is considered as the reference technique. Culture media were prepared according to the protocol described in Brown (1981). Briefly, sterile RPMI-1640 (Gibco Invitrogen Australia Ltd, Victoria, Australia) was mixed with Heparin (Sigma) at the final concentration of 100 IU/ml, 2-Mercaptoethanol (Sigma) (10⁻⁵ M) and penicillin-streptomycin at the concentrations of 10⁵ IU/ml and 0.1 g/ml, respectively. The media was
filtrated through a 0.2 μm Millipore® filters and mixed with 10 % foetal calf serum (Gibco Invitrogen Australia Ltd, Victoria, Australia) at the concentration and 100 μM/ml L-Glutamine (Gibco Invitrogen Australia Ltd, Victoria, Australia).

Around day 12 post-infection, the animals of both groups presented typical clinical signs of tropical theileriosis. Blood samples were taken at the onset of clinical signs consistent with the onset of an episode of acute tropical theileriosis. Whole blood was aseptically collected in heparinised tubes (Vacutainer®).

For Ficoll (Histopaque® 1077, Sigma) PBMC isolation, ten millilitres of blood were aseptically collected in heparinised sterile tubes (Vacutainer®), peripheral blood monocytes were isolated as recommended by the supplier. For the whole blood technique only one millilitre of blood was mixed with complete RPMI-1640 (Gibco Invitrogen Australia Ltd, Victoria, Australia) in a 25 cm² cell culture vial in vertical position. All cell cultures were incubated at 37 °C, 90 % humidity and 5 % CO₂ for two weeks.

The cell cultures were monitored by making cytospin smears three times a week. The Giemsa stained cell culture smears were examined microscopically for T. annulata schizont infected cells. To compare the two techniques, six indicators were estimated:
1. Cell culture effectiveness is the number of positive cell cultures at the end of the survey divided by the number of starting cells.
2. Range of first day of passage is the minimum delay to obtain a schizont index (SI) superior to 50 %. SI is the ratio of infected WBC/examined WBC.
3. Number of contaminated vials.
4. Schizont index at day 9.
5. First-day of schizont detection.
6. Costs difference = cost of reference technique - cost of the rapid method (the labour costs and laboratory equipments were ignored).

RESULTS AND DISCUSSION

Compared to the reference technique, the rapid protocol is more effective, saving both time and money (Table 1).

<table>
<thead>
<tr>
<th>Indicator</th>
<th>Ficoll extraction</th>
<th>Whole blood</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell culture sensitivity</td>
<td>19/22</td>
<td>20/20</td>
</tr>
<tr>
<td>Range of first day of passage</td>
<td>11 to 21</td>
<td>6 to 9</td>
</tr>
<tr>
<td>Number of contaminated vials</td>
<td>3/22</td>
<td>0/20</td>
</tr>
<tr>
<td>Schizont index at day 9 (in %)</td>
<td>ND</td>
<td>100</td>
</tr>
<tr>
<td>First day of schizont detection</td>
<td>ND</td>
<td>6</td>
</tr>
<tr>
<td>Costs difference ($)</td>
<td>Reference</td>
<td>-3.06</td>
</tr>
</tbody>
</table>

ND: not done.

Table 1 – Comparison of two Theileria annulata isolation techniques: culture of whole blood and following Ficoll purification.

can be attributed to the small quantity of stained samples when using the cytospin® (30 μl per sample).

At day 6, the cultures showed a high proliferation rate, at day 9, the schizont index increases dramatically for all the animals, and reached 80 to 90 % at day 15. The index schizont progresses similarly to the conventional cell culture methods.

The parasite cultures isolated by Ficoll were successful for 19 out of 22 samples. Pipano et al. (1990) estimated the effectiveness of Ficoll isolation to 50 %. However, they examined less severe clinical cases than those used in the present study. Indeed, they worked on animals challenged with T. annulata contaminated blood, whereas in our study the animals were challenged with the Jed 4 GUTS at the lethal dose of 1.8 to 2.5 tick equivalents. It has been established that infection with sporozoites induces a more severe infection than inoculation of T. annulata present in blood (Darghouth, 2000).

When considering the new rapid protocol, the day of first passage (first day when the schizont index is equal or higher than 50 %) was estimated to either day 6 or day 9, whilst the first passage is realised from day 11 to 21 (median: day 15) for Ficoll. Cell culture from total blood allows a more rapid development of infected cells, since they are suspended in whole blood containing several growth factors leading a rapid growth of the cells and good output of cultures. Indeed, the T. annulata-infected cells can activate and induce non-specifically the proliferation of non-infected autologous T cells (Glass & Spooner, 1990), known as Theileria mixed lymphocyte reaction (Theileria MLR). In MLR, the non-specifically recruited lymphocytes produce cytokines that enhance the expansion of infected cells with a dramatic increase of INFγ (gamma interferon) and TNFα (tumour necrosis factor alpha) concentrations (Glass, 2001). These cytokines increases the proliferation of young schizonts and induce the production of activated macrophages that will secrete IL-2. During the early infection stage, exogenous IL-2 secreted by non-infected lymphocytes contributes to the initiation of infected cells transformation (Dobbelaere et al., 1990).
The present technique offers a good environment to the cells, since they are not washed and centrifuged. Cell washing can lead to the loss of infected cells and decrease the output of cell cultures with an increase risk of contamination. Indeed, three out of 22 Ficoll treated samples were contaminated. One contamination was even observed before the first passage, whilst the two others were observed later.

The new technique leads to a dramatic saving in time, since it can be performed in few minutes, whereas the conventional technique requires two to three hours. In addition, the protocol saves €3.06 per sample when compared to the conventional technique. Combined, they could encourage the use of this new technique for the isolation of *T. annulata* from infected animals and its validation under field conditions, where *T. annulata*-infected cattle are in a carrier state.

**ACKNOWLEDGEMENTS**

This study received financial support from the "Laboratoire d’Épidémiologie des Infections Enzootiques des Herbivores en Tunisie (Ministère de la Recherche Scientifique et de la Promotion des Compétences, Tunisie)."

**REFERENCES**


Received on December 19th, 2011. Accepted on February 23rd, 2012.