CASE REPORT

Rapid detection of Candida parapsilosis contamination in the infusion fluid

Shigeharu Oie*1,2, Kyoji Kouda2, Hiroyuki Furukawa2, Akira Kamiya3

1 Laboratory of Hospital Hygiene and Infection Control, Japan
2 Yamaguchi University Hospital, Japan
3 Sanyo-onoda City University, Japan

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ABSTRACT

We encountered a case of microbial contamination of infusion fluid detected based on the discoloration of an inline filter (membrane filter) in the main route. A cause of this contamination may be the use of an extension tube connected to the main route for more than a month. In the infusion fluid above the inline filter (on the high calorie end), \(2.4 \times 10^5\) colony forming units (cfu/ml) of Candida parapsilosis was detected. “Adenosine triphosphate (ATP) + adenosine monophosphate (AMP)” was determined in addition. We conclude that microbial contamination of this infusion fluid can be rapidly estimated by measuring the amounts of “ATP + AMP”.

Key Words: Contamination, Infusion, Fentanyl, Candida parapsilosis, ATP

1. INTRODUCTION

Microbial contamination during injections leads to serious consequences such as sepsis.1–8 However, due to difficulties of visually confirming the contamination by the naked eye and added time required to plate and grow the contaminating species, the infusion fluid does not currently receive rigorous testing as a possible source of microbes causing sepsis in a patient. In clinical practice, we encountered a case in which microbial contamination of an injection was detected based on the discoloration of an inline filter (membrane filter) of the central venous line. Although discoloration of the filter was noticed in the filter, the contamination of the filter was not initially suspected as the cause of the contamination. Therefore, a method for rapidly detecting the presence of microbial contamination is urgently required. In this study, the availability of “adenosine triphosphate (ATP) + adenosine monophosphate (AMP)” measurement as the rapid estimation of microbial contamination in the infusion fluid was evaluated.

2. MATERIALS AND METHODS

2.1 Background

The Department of Pharmacy received an inquiry from a nurse in charge of a 62-year-old male receiving intravenous hyperalimentation (IVH) in the inpatients’ hospital ward after surgery for esophageal cancer. The nurse had noticed that the inline filter of the drip infusion set had become brown in color. The nurse initially had suspected drug incompatibility. A high calorie infusion fluid (Elneopa® No.1, Terumo Co., Japan) was being administered through the main route while fentanyl (Janssen Pharmaceutical Japan Co., Japan) was being administered from a side tube connected to the

*Correspondence: Shigeharu Oie; Email: oie@frontier-u.jp; Address: 4-2-15 Astopia, Ube 755-0152, Japan.
main route (see Figure 1). Both the high calorie infusion fluid bag and the fentanyl injection together with the syringe were replaced at 24-hour intervals.

2.2 Investigation of microbial contamination and measurement of “ATP + AMP”

The residual fluid in the main route was diluted with normal sterile saline. Pipettes were used to transfer 1 ml of undiluted or diluted samples to grow any bacteria on trypticase soy agar containing blood and to grow any fungi on Sabouraud dextrose agar. The plates were streaked with a glass “hockey stick” and incubated at 35°C for 24-72 h. Colonies were counted, and organisms were identified by Gram staining, morphological examination, the oxidation fermentation test, cytochrome-oxidase test, and the API system (bio-Mérieux SA, L’Etoile, France).

For the measurement of “ATP + AMP” (relative light units: RLU), sampling was performed using LuciPac™ Pen-AQUA (Kikkoman Biochemifa Co., Japan), and RLU was measured using Lumitester™ PD-20 (Kikkoman Biochemifa Co.).

The inner walls of the main route and extension tube were observed using a field emission scanning electron microscope (JSM-7000F, JEOL Co., Japan) after gold evaporation at an accelerating voltage of 5 kV.

3. RESULTS

Figure 1 shows viable counts of *C. parapsilosis* and the amounts of “ATP + AMP” (RLU). This microorganism was not detected in the fentanyl injection in the syringe, inside the high calorie infusion fluid bag, in the high calorie infusion fluid in the drip tube connected to the bag, or in the fluid in the main route from the inline filter to the patient. The RLU was 0-93. However, in the other areas, this microorganism was detected, and the RLU was 2,200-2,745. High RLU strongly correlated with the high colony forming units (cfu) (2.2-2.4 × 10⁵).

Figure 2 shows a micrograph (× 4,000) of the inner wall of the extension tube. Attachment of a large amount of *C. parapsilosis* was observed on the inner wall of the extension tube. However, *C. parapsilosis* was not confirmed on the inner wall of the main route.

![Diagram of infusion system](image)

**Figure 1.** *C. parapsilosis* count and “ATP + AMP” (RLU) in a case of infusion fluid contamination
4. DISCUSSION

The observation of microbial contamination in medical drugs by the naked eye is rare. Since many drugs do not have adequate nutrients for microorganisms, their microbial contamination at a level allowing visual confirmation (about $10^7$ cfu/ml for bacteria and about $10^6$ cfu/ml for yeast-like fungi) does not normally occur. However, in this case, since an inline filter was used in the main route of high calorie infusion fluid, microbial contamination was detected based on the color of the inline filter. *C. parapsilosis* is normally white, but may have changed to light brown after turning yellow due to vitamin C as a component of the high calorie infusion fluid.

Microscopy of the inner wall of the used extension tube revealed attachment of a large amount of *C. parapsilosis* to the inner wall of the extension tube used for fentanyl injection administration (see Figure 2). Fentanyl injections have been reported to be susceptible to microbial contamination. In this case, colonization on the inner wall of the extension tube during fentanyl administration was considered. We hypothesize that the contamination may have occurred by not exchanging the extension tube for more than 30 days. Although the hospital was required to exchange the extension tubes and main route twice every week, the extension tube was not exchanged due to exemption warranted by the protocol for having opioid fentanyl in the extension tube. *C. parapsilosis* is part of normal skin flora and therefore it is possible that the extension tube was contaminated due to improper handling.

ATP measurement for the rapid estimation of microbial contamination has been evaluated in in vitro studies. However, ATP is known to be degraded to ADP and subsequently to AMP by the metabolism. Moreover, previous study showed that AMP may predominate over ATP according to microbial species and cell conditions. Therefore, measuring ATP and its metabolites, rather than ATP alone, seems to be useful to detect microbes and contaminants.

Since the rapid “ATP + AMP” monitoring test kit is commercially available, “ATP + AMP” was measured in order to confirm the microbial contamination. Basically, this test kit is not for the specific detection of microorganism because the “ATP + AMP” increase can also be caused by organic substances except for bacteria, such as body fluid. However, as shown in Figure 1, the high correlation between “ATP + AMP” and cfu was observed, and bacteria in the infusion fluid could be monitored using the “ATP + AMP” test.
a slight contamination is happened, bacteria seem to grow and produce the large amount of “ATP + AMP”. The results of those studies suggest that “ATP + AMP” measurement is applicable and useful in the clinical setting. In the future, we intend to further investigate the microbial contamination during injections with the aim to establishing a rapid detection method of microbial contamination.

5. CONCLUSIONS

“ATP + AMP” measurement may be applicable and useful for rapid detection of microbial contamination of infusion systems in the clinical setting.

CONFLICTS OF INTEREST DISCLOSURE

The authors declare they have no conflicts of interest.

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