A Simple Method for Determining Antitubulinic Activities of Ansamitocins and Related Compounds Using a Cilia Regeneration System with Deciliated *Tetrahymena*

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Cilia regeneration with deciliated *Tetrahymena pyriformis* W was tested to determine the antitubulinic activities of ansamitocins and related compounds in a microbial system. Various factors interfered with the regeneration process, i.e. excess shearing force in deciliation procedure, high temperature (32°C or above), high or low pH (pH 9 or 5), and exogenous divalent cations, such as Zn²⁺, Mn²⁺, Cu²⁺ and Co²⁺. Under suitable conditions, cilia regeneration was completed in about 60 min of incubation, and a number of newly formed cilia were observed around the cell surface. When ansamitocins were added to the recovery solution, cilia regeneration was completely suppressed without alterations in cell shape or the surface structure of deciliated *Tetrahymena*. In this assay system, the inhibitory activity of ansamitocins was slightly stronger than that of maytansine.

The cilia and flagella in eukaryotic microorganisms contain highly organized microtubule systems.¹ The formation process of these organelles depends on the assembly of tubulin to ciliary or flagellar microtubules, and is affected by antimitotic agents.²⁻⁶) Rosenbaum and Carlson²) developed the model system for cilia formation of *Tetrahymena* using deciliated cells in which regeneration of cilia was determined as recovery of motility from nonmotile cells. We have found that ansamitocins, maytansinoid antitumor antibiotics,⁷⁻⁹) have strong activity to inhibit cilia regeneration of *Tetrahymena*. Part of this finding has been reported and we suggested the possibility that the regeneration system is a useful tool for determining the antitubulinic properties of ansamitocins and related compounds.⁶)

This paper deals with the factors affecting cilia regeneration of deciliated *Tetrahymena* and the inhibitory effect of ansamitocins and maytansine¹⁰) on regeneration.

**MATERIALS AND METHODS**

*Media and culture.* *T. pyriformis* W was used in all experiments. The growth medium (PPYG) contained proteose-peptone (Difco), 2%; yeast extract (Difco), 0.1%; glucose, 0.2%. The cells were grown for 2 days at 28°C without shaking. The cell density at this time was about 5 x 10⁵ /ml. The organism was maintained in PYPG at room temperature and transplanted to fresh PYPG at two-week intervals.

*Assay for cilia regeneration.* A 0.5ml portion of the stock culture was introduced into 20ml PYPG in a 250ml Erlenmeyer flask and incubated for 2 days at 28°C. The cells were harvested at room temperature by centrifugation at 200 × g for 3 min. After removal of the supernatant by aspiration, the pellet was resuspended in 5ml cold PYPG. Deciliated cells were obtained by the method of Rosenbaum and Carlson¹¹) with some modifications. A 2ml portion of the suspension was added to 4ml cold EDTA-acetate buffer (10mM EDTA·2Na; 50mM sodium acetate, pH 6.0) in a 50ml centrifuge tube with a screw cap. After addition of 2ml of cold distilled water, followed by the addition of 0.2ml of 0.2m CaCl₂, the tube was inverted five times. If necessary, the suspension was subjected to shearing with a 5ml glass syringe with a needle. To 1ml of a recovery solution (mainly 0.02m potassium phosphate buffer, pH 7.0), 0.05ml of the deciliated cell suspension was added and

The cells grown at 28°C for 2 days were harvested by centrifugation at 200 × g for 3 min. Deciliated cells were obtained by the method of Rosenbaum and Carlson2) with the modifications at the final step.

A, a cell before deciliation; B, magnification of A; C, a deciliated cell without final shearings with a needle; D, a deciliated cell with five times shearings; E, magnification of C; F, a cell capable of swimming slowly which was partially recovered in 0.02 M potassium phosphate buffer, pH 7.0; G, magnification of F; H, a cell with full motility in the recovery solution; I, magnification of H. (A, C, D, F and H, ×1000; B, E, G and I, ×3500.)
incubated at 28°C. Ansamitocins and maytansine were added to the recovery solution prior to the addition of the cell suspension. The regeneration of cilia was assayed by observing the recovery of motility in deciliated cells with a microscope (× 40 magnification).

Scanning electron microscopy. Tetrahymena cells fixed with 0.1% glutaraldehyde were dehydrated through ethyl alcohol series. For scanning electron microscopy the cells were critical-point dried, coated with gold and viewed on a MSM4C-101 scanning electron microscope.

Chemicals. Ansamitocin P-3 and P-4, purified from a culture broth of Nocardia sp. C-15003 (N-1),8,9) were dissolved in methanol to a final concentration of 1,000 µg/ml and stored at -20°C. Under these conditions the biological activity of ansamitocins is stable for more than one month. Maytansine used in this experiment was prepared in our research division.

RESULTS AND DISCUSSION

Influence of shearing force on deciliation of Tetrahymena

The condition of deciliation is traumatic to the maintenance of viability in Tetrahymena. In order to obtain deciliated cells with minor loss of viability, deciliation was carried out according to the procedure of Rosenbaum and Carlson2) with the modification of shearing at the final step of this treatment. Scanning electron microscopic observation shows that all of the deciliated cells (Fig. 1C, D, E) lost most of their cilia and ceased swimming, and that the cells were also damaged by this treatment resulting in a rough surface (Fig. 1E) as compared with that of the normal cell (Fig. 1A, B). The deciliated cells prepared by inverting the tube with the cell suspension followed by shearing five times with a needle lost some of oral and all of somatic cilia (Fig. 1D), whereas the cells without the final shearings still had oral and a few somatic cilia (Fig. 1C). Figure 2 shows the kinetics of recovery of motility of the deciliated cells in 0.02 M potassium phosphate buffer, pH 7.0 as the recovery solution. After 60 min of incubation, more than 90% of the deciliated cells without final shearings were fully motile. As shown in Fig. 1H and I, cells recovering their motility had numerous newly formed cilia on their surface. In the early stage of cilia regeneration, many of the slow swimming cells had many immature short cilia (Fig. 1F, G). On the other hand, only half of the deciliated

![FIG. 2. Recovery of Motility of Deciliated Tetrahymena. Deciliated cells were prepared as in Fig. 1. To 1 ml of 0.02 M potassium phosphate buffer, pH 7.0, 0.05 ml of deciliated cell suspension was added and incubated at 28°C without shaking. Recovery of motility was assayed by scoring of motile cells with a phase contrast microscope. ●-○, inverted the tube; ●-●, inverted the tube followed by shearing five times with a needle; ○-○, inverted the tube followed by shearing once; ○-○, only shearing five times; ■-■, no physical force added.](image-url)

![FIG. 3. Effect of Recovery Solutions on Cilia Regeneration in Deciliated Tetrahymena. Deciliated cells were prepared by only inverting the tube for detachment of cilia. Procedures are the same as in Fig. 2. ●-○, in 0.02 M potassium phosphate buffer, pH 7.0; ○-○, in 0.05 M Tris-HCl buffer, pH 7.5; ○-○, in PPYG.](image-url)
cells with once or five times shearings recovered their motility because of the loss of viability of many cells during this treatment. When a cell suspension after addition of CaCl₂ was immediately subjected to shearing five times, the recovery pattern of these cells was similar to that of the cells deciliated without final shearing (Fig. 2). Although shearing with a needle at the final step in this procedure was not necessary to obtain deciliated cells, physical force was essential for the detachment of cilia. Most of the cells after addition of CaCl₂ were capable of swimming in the recovery solution unless the tube was inverted or shearing with a needle was followed (Fig. 2). From these results, deciliated *Tetrahymena* was prepared without shearing at the final step of this procedure.

**Effect of environmental factors on cilia regeneration in *Tetrahymena***

In order to determine the antitubulinic activity of ansamitocins in culture broths on the cilia regeneration system in deciliated *Tetrahymena*, effect of various environmental factors on the regeneration process were examined. Figure 3 shows the recovery kinetics of deciliated cells in various recovery solutions. No significant difference was observed among the recovery patterns. The incubation temperature and pH of recovery solutions affected the cilia regeneration. After 90 min of incubation, normal recovery of motility was obtained in the variation between 24 to 28°C, and between pH 6 to 8 (Fig. 4A, B). At 32°C or above, recovery of motility was reduced. Similarly, reduction in the recovery of motility was observed at pH 5 or 9. The effect of temperature and pH on cilia regeneration was correlated with that on growth of *Tetrahymena*¹¹ and therefore, the interfering effect was due to nonspecific actions to the organism.

Ansamitocins are produced by an actinomycete, *Nocardia* sp. No. C-15003 (N-1)⁷,⁸ in liquid media. In order to determine the amount of ansamitocins in culture broths of the *Nocardia*, effect of compounds commonly used as components of the culture media for fermentative production were tested with 0.05 M Tris-HCl buffer, pH 7.5 as the recovery solution. As shown in Fig. 5, raising the concentration of KCl, NaCl or potassium phosphate buffer caused marked reduction of cilia regeneration. The suppression by potassium phosphate buffer at 0.1 M or above may be mainly due to the function of the K⁺ ion and not of the phosphate ion. Some of the divalent cations also affected regeneration. As shown in Fig. 6, ZnCl₂ at 0.5 mM, MnCl₂ at...
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No. C-15003 (N-1) was examined. Since cations in the fermentation media for production of ansamitocins are less than levels that affect cilia regeneration, the amount of the antibiotics in the culture was able to be determined by the assay system with 0.05 M Tris-HCl buffer, pH 7.5 containing up to 50% (v/v) of the supernatant of the culture broth which was prepared by centrifugation at 3000 × g for 10 min at room temperature. The inhibitory activity depends on the amount of ansamitocins because the antibiotics are only products with antitubulinic properties of the Nocardia. As compared with the in vitro polymerization system for tubulin to microtubules, the cilia regeneration system was relatively tolerant to various environmental factors. Therefore the regeneration system is suitable in the determination of the antitubulinic properties of fermentation products.

Effect of ansamitocins and maytansine on cilia regeneration in Tetrahymena

Ansamitocins at low concentrations prevent cilia regeneration in deciliated Tetrahymena. When deciliated cells were exposed to ansamitocin P-3 at 1 µg/ml, no cilia regeneration occurred and the cell shape remained unaffected (Fig. 7). But the surface structure of the cells recovered from the damage caused by the deciliation treatment. In this experiment, no significant difference was observed between ansamitocins P-3 and P-4. This indicates that ansamitocins have specific actions against cilia formation process in this system, probably due to interference with the assembly of tubulin to ciliary microtubules. On the basis of structural similarity, effect of maytansine, originated from plants with antitumor and antimitotic properties, originated from plants with antitumor and antimitotic properties, on cilia regeneration of Tetrahymena was tested. As shown in Fig. 8, maytansine interfered with cilia regeneration of deciliated Tetrahymena at low concentrations. The inhibitory activity was slightly weaker than that of ansamitocins. The action of maytansine, like ansamitocins, may be based on its antitubulinic properties.

0.5 mM or CuSO₄ at 1 mM completely suppressed cilia regeneration, and CoCl₂ at 0.5 to 1 mM showed partial inhibition. Although CaCl₂ is known to cause interference with the polymerization of tubulin to microtubules in vitro, cilia regeneration was not affected by CaCl₂ at levels up to 1 mM. On the basis of these results, determination of the amount of ansamitocins in culture broths of Nocardia sp.
FIG. 7. Scanning Electron Micrographs of Inhibition of Cilia Regeneration in *Tetrahymena* by Ansamitocin P-3.

Ansamitocin P-3 was added at 1 µg/ml to 0.02 M potassium phosphate buffer before inoculation of the deciliated cell suspension. After 90 min of incubation, the cells were fixed with glutaraldehyde and were observed by a scanning electron microscope.

A, ansamitocin P-3 added; B, magnification of A. (A, ×1000; B, ×3500.)

FIG. 8. Effect of Ansamitocins and Maytansine on Cilia Regeneration in *Tetrahymena*.

Ansamitocin P-3, P-4 or maytansine, in varying concentrations, was added to the recovery solution before inoculation of the deciliated cell suspension. After 90 min of incubation, motile cells were scored.

- - - - - , ansamitocin P-3; O—O, ansamitocin P-4; O---O, maytansine.

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