Biological activities and chemical composition of a cytotoxin of *Klebsiella oxytoca*

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A low-molecular-mass cytotoxin produced by *Klebsiella oxytoca* isolated previously from patients with antibiotic-associated haemorrhagic enterocolitis was purified, and its biological and chemical properties were elucidated. The toxin inhibited the syntheses of DNA and RNA by HEp-2 cells dose-dependently, whereas protein synthesis was only slightly inhibited, as measured by the incorporation of radioactive precursors. When synchronously cultured HEp-2 cells were examined in the presence of cytotoxin, inhibition of DNA synthesis occurred promptly within 5 h, but cell-rounding, the earliest visible morphological change, was not observed until 6 h after exposure. The intracellular levels of ATP decreased with an approximately similar time course. These results suggest that cytotoxicity toward HEp-2 cells is primarily due to the inhibitory effect of the cytotoxin on nucleic acid synthesis, possibly on DNA synthesis. Cell rounding and cell death were induced even in the absence of the cytotoxin after incubation with the cytotoxin for 6 h. The cytotoxin was heat-labile, cytotoxic activity decreasing to 50% of the initial level on heating at 70 °C for 20 min. Plasmids were extracted from three strains of *K. oxytoca* producing the cytotoxin and analysed by agarose gel electrophoresis. Two strains possessed plasmids of different sizes, but one strain possessed no plasmid, indicating that the cytotoxin is probably chromosomally encoded. Analysis by NMR and FAB-mass-spectrometry revealed that the molecular mass of the cytotoxin should be 217.1062 Da (exact mass), its molecular formula being C8H15O4N3.

Introduction

*Klebsiella oxytoca* isolated from patients with antibiotic-associated haemorrhagic enterocolitis have been shown to produce a cytotoxin (Minami et al., 1989). This cytotoxin exerts a cytotoxic effect, e.g. cell-rounding followed by cell death, on many tissue culture cells, such as HEp-2, CHO, HeLa and Vero cells, and hence can be regarded as a factor responsible for the enterocolitis caused by this organism (Minami et al., 1989). The most outstanding feature of the cytotoxin is its low molecular mass. Cytotoxins have been isolated from many bacteria, such as *Shigella* sp. (O'Brien & Holmes, 1987), *Escherichia coli* (O'Brien & Holmes, 1987), *Clostridium difficile* (Donta & Shaffer, 1980, Lima et al., 1988), *Campylobacter jejuni* (Guerrant et al., 1987) and *Salmonella* sp. (Ketyi et al., 1979; O'Brien et al., 1982; Baloda et al., 1983; Koo & Peterson, 1983; Ashkenazi et al., 1988), but they are protein or proteinaceous toxins with high molecular masses (more than 10 kDa) in contrast to the *K. oxytoca* cytotoxin (less than 1 kDa).

Recently, a cytotoxin of low molecular mass from *Bordetella pertussis* was purified and characterized (Rosenthal et al., 1987; Cookson et al., 1989). The *B. pertussis* tracheal cytotoxin is a disaccharide tetrapeptide subunit of peptidoglycan, molecular mass approx. 1 kDa, causing a ciliated-cell-specific respiratory tract pathology. This cytotoxin, together with that of *K. oxytoca*, suggests that cytotoxins of low molecular mass can be produced by some bacteria and can contribute to the virulence of highly or even potentially pathogenic bacteria. Elucidation of the molecular nature of the *K. oxytoca* cytotoxin, and the mechanism underlying cell-rounding and cell death induced by the cytotoxin would provide useful information for understanding the pathogenicity of this bacterium and related Enterobacteria-
ceae. In this study we examine the effects of the cytotoxin on macromolecular synthesis by HEp-2 cells and present evidence that it preferentially inhibits nucleic acid synthesis.

In a previous paper (Minami et al., 1989), we described a procedure for the purification of the cytotoxin, consisting of gel filtration on Sephadex G-25 and Bio-gel P2, and reversed-phase HPLC. In this study, the cytotoxin has been further purified by extraction with chloroform after the above procedure, and has been analysed by NMR and FAB-mass spectrometry to determine its chemical composition.

Methods

Organisms. Three clinically isolated strains of K. oxytoca (OK-1, KA-1 and KA-2) and the type strain of K. oxytoca, ATCC 13182, were used in this study. The three strains were isolated from patients with hemorrhagic diarrhoea after the administration of penicillin derivatives as chemotherapy for an upper respiratory infection, and were characterized previously (Minami et al., 1989).

Purification of K. oxytoca cytotoxin. K. oxytoca OK-1 was grown in Trypto-Soya broth (Nissui Pharmaceutical, Tokyo) at 37 °C for 12 h, and the K. oxytoca cytotoxin was purified from the culture supernatant by gel filtration on Sephadex G-25 and Bio-Gel P-2, and reversed-phase HPLC, as described previously (Minami et al., 1989).

Measurement of ATP in HEp-2 cells. A 24-well plate was used to culture HEp-2 cells. Each well was seeded with 1 × 10^5 HEp-2 cells. The cells were cultured in 1 ml of Eagle's minimum essential medium with Earle's salts, 100 μg streptomycin ml⁻¹ and 100 U penicillin ml⁻¹ (MEM medium) supplemented with 10% (v/v) foetal bovine serum in the presence of 5% (v/v) CO₂ at 37 °C for 24 h to establish monolayers. The medium was replaced by fresh MEM medium supplemented with 5% (v/v) foetal bovine serum. After incubation at 37 °C for 1 h, the cytotoxin was added to a final concentration of 8 μg ml⁻¹. ATP in HEp-2 cells was then solubilized in 200 μl 0.5 M-perchloric acid. The extract was neutralized with 5 M-potassium hydroxide and the precipitate removed by centrifugation. The extracted ATP was determined by HPLC on an anion exchange column (4.6 x 250 mm) of DAEAE-25W (Toyo Soda Manufacturing, Tokyo) equilibrated with 360 mM-sodium phosphate buffer, pH 6.0, as described by Watanabe et al. (1985). After solubilization of HEp-2 cells with 1 M-NaOH, protein was measured by the Lowry method with bovine serum albumin as standard.

Macromolecular synthesis. The effects of K. oxytoca cytotoxin on macromolecular synthesis were examined using HEp-2 cells. The synthesis of DNA, RNA and protein by HEp-2 cells in a 96-well plate were monitored by measuring the incorporation into the TCA-precipitable fraction of [3H]thymidine, [3H]uridine and [3H]leucine, respectively. HEp-2 cells were maintained in Eagle's minimum essential medium containing Earle's salts supplemented with 10% (v/v) foetal bovine serum, 100 μg streptomycin ml⁻¹ and 100 U penicillin ml⁻¹ (basal medium). Each well was seeded with 1 × 10^4 HEp-2 cells, followed by incubation in the presence of 5% (v/v) CO₂ at 37 °C for 24 h to establish monolayers. Nonsynchronous cultures of HEp-2 cells were used for the experiments on protein and RNA syntheses. Cultures synchronized at the S phase were used for the experiment on DNA synthesis. Synchronization was performed as follows. Monolayers of cells in the wells of a microculture plate were incubated in the basal medium supplemented with 2.5 mM-thymidine for 24 h. The medium was then removed and the plate was washed once with Dulbecco's phosphate-buffered saline (PBS). After the addition of fresh basal medium without thymidine, the plate was incubated for 10 h, followed by the addition of hydroxyurea to a final concentration of 1 mM. After incubation for 14 to 16 h, the medium was removed. The plate was washed three times with PBS, and then 0.2 ml of incorporation medium [Earle's minimum essential medium containing Earle's salts supplemented with 5% (v/v) foetal bovine serum 100 μg streptomycin ml⁻¹ and 100 U penicillin ml⁻¹] was added. Leucine-deficient incorporation medium was used for the experiment on protein synthesis. Prewarmed washing solutions and media were used throughout these procedures.

Experiments on DNA and protein syntheses were done as follows. The basal medium in each well with a monolayer of cells was replaced with 0.2 ml incorporation medium, followed by the addition of 50 μl of medium containing cytotoxin and radioactive precursors to initiate cell labelling. One micromole of [methyl-3H]thymidine (25 Ci mmol⁻¹; 925 GBq mmol⁻¹; Amersham) and 0.5 μCi of [1-4,5-3H]leucine (120–190 Ci mmol⁻¹; 4·44–7·03 TBq mmol⁻¹; Amersham) were added to each well to monitor the syntheses of DNA and protein, respectively, and the plate incubated at 37 °C. At the indicated times, the medium was removed and the plate washed three times with PBS. Cells were solubilized by the addition of 37 μl 0·5 M-KOH to a well, and incubation on a vibrating platform at room temperature for 30 min. Bovine serum albumin, 7·5 μl of a 0·15% (w/v) solution, was added as a carrier protein and the solubilized cells were then precipitated with 270 μl cold 10% (w/v) TCA. TCA-precipitable materials were collected at 4 °C on a glass fibre filter using a TiterTek cell harvester (Flow Laboratories) and then washed with cold 5% (w/v) TCA. The precipitate on the filter disk was treated with NCS solubilizer (Amersham). Radioactivity was measured with a liquid scintillation counter, using ACS II scintillation cocktail (Amersham).

For the experiment on RNA synthesis, cells were treated as described above, except that 1 μCi per well of [5-3H]uridine (29 Ci mmol⁻¹; 1·1 TBq mmol⁻¹; Amersham) was added and the labelled cells were solubilized in a different manner. After the indicated incubation times, the cells were washed, and detached by trypsinization [0·25% (w/v) trypsin and 0·02% EDTA at 37 °C for 5 min], collected on a glass fibre filter and washed with PBS. After precipitation and washing with cold 5% (w/v) TCA, the filter was treated with NCS solubilizer and its radioactivity was measured as described above.

Assay for cytotoxic activity. HEp-2 cells were monolayered and cultured in a 96-well plate, and then the cytotoxic activity (CD₅₀) of K. oxytoca cytotoxin was measured using HEp-2 cells as described in the previous paper (Minami et al., 1989). CD₅₀ was expressed as the dry weight of the cytotoxin (in 250 μl medium) required to cause the rounding of 50% of the HEp-2 cells after incubation at 37 °C for 48 h. The CD₅₀ of the purified cytotoxin was 0·15 μg per well (Minami et al., 1989).

Plasmid analysis. Plasmids were extracted from K. oxytoca strains by the method of Kado & Liu (1981) and then electrophoresed on a 0·7% agarose gel at 30 V with the electrophoresis buffer, i.e. 2 mm-EDTA and 40 mm-Tris acetate, pH 7·9. After electrophoresis, the gel was stained with ethidium bromide (0·5 μg ml⁻¹) and then photographed under a UV illuminator.

Chemical analysis. 1H- and 13C-NMR spectra were recorded on a Varian VXR-500 instrument (500 MHz for 1H-NMR and 126 MHz for 13C-NMR). 31P-NMR spectra were determined on a Varian VXR-200 instrument. Deuteriochloroform was used as a solvent. Chloroform soluble cytotoxin (several mg) was purified on a short-path column packed with 2 g silica gel (Merck 60-7734) using chloroform ethanol (24:1) as the eluent. Fractions containing the cytotoxin, as judged by analytical TLC, were concentrated under reduced pressure at room
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I80 temperature to give an almost colorless viscous oil, which was dissolved in deuteriochloroform (0.7 ml) for NMR analysis. All NMR measurements were carried out in the standard manner at 20°C (probe temperature). FAB mass spectra were obtained on a VG Analytical VG-70SE double focusing mass spectrometer, employing 3-nitrobenzyl alcohol matrix, in the standard manner. Analytical TLC was performed on pre-coated Merck silica gel 60FZs4 (0.25 mm thickness).

Results

**Effects of *K. oxytoca* cytotoxin on HEP-2 cells**

A characteristic feature of the cytotoxicity exerted by the *K. oxytoca* cytotoxin is cell-rounding. Fig. 1(a) shows the time course of cell-rounding and mortality of HEP-2 cells during incubation in the presence of the cytotoxin (2 µg per well). In the early incubation period, up to 6 h, cell-rounding occurred only slightly (less than 13%), but thereafter rounded cells increased in number, accounting for more than 90% of the total cells after 24 h incubation. On the other hand, cell death was observed in a significant number of cells only after 8 h incubation and proceeded similarly to cell-rounding except for a time lag of 2–3 h. These results indicate that the cell-rounding does not result from cell death and that heterogeneity exists in susceptibility of individual HEP-2 cells to the cytotoxin.

An irreversible change might have occurred in an early incubation period when no morphological change of the HEP-2 cells was observed. We examined irreversibility of the cytotoxicity to HEP-2 cells. At various time intervals after addition of the cytotoxin, the cells were washed and cultured in fresh medium without the cytotoxin. As shown in Fig. 1(b) the cytotoxic effect was reversible when cells were exposed to the cytotoxin only for 1 h. However, an irreversible effect was exerted on a substantial fraction of the cells when they were exposed for more than 2 h. Mortality increased proportionally to length of exposure time, which may also reflect the various periods of time. HEP-2 cells were cultured in 250 µl MEM medium supplemented with 5% (v/v) foetal bovine serum in a 96-well plate. At various times during incubation in the presence of 2 µg cytotoxin per well, the medium was removed and HEP-2 cells were washed. The cells were cultured in fresh medium without the cytotoxin for 24 h. Mortality of HEP-2 cells was determined by counting cells taking up trypan blue under a microscope. A total of 600 cells were examined. Data are means of three determinations.

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Fig. 1. (a) Time courses of cytotoxin-induced cell-rounding, and mortality of HEP-2 cells. HEP-2 cells were cultured in 250 µl MEM medium supplemented with 5% (v/v) foetal bovine serum in the presence and absence of the cytotoxin in a 96-well plate. Mortality was determined by trypan blue exclusion. Numbers of rounded cells and trypan-blue-uptaking cells were determined under a microscope. A total of 600 cells were examined. Data are means of three determinations. ○ Rounded cells in the absence of cytotoxin; ● rounded cells in the presence of 2 µg (13-3 CD₅₀) cytotoxin per well; △, mortality in the absence of cytotoxin; ◇, mortality in the presence of 2 µg cytotoxin per well. (b) Mortality of HEP-2 cells exposed to *K. oxytoca* cytotoxin for various periods of time. HEP-2 cells were cultured in 250 µl MEM medium supplemented with 5% (v/v) foetal bovine serum in a 96-well plate. At various times during incubation in the presence of 2 µg cytotoxin per well, the medium was removed and HEP-2 cells were washed. The cells were cultured in fresh medium without the cytotoxin for 24 h. Mortality of HEP-2 cells was determined by counting cells taking up trypan blue under a microscope. A total of 600 cells were examined. Data are means of three determinations. (c) Changes in ATP levels in HEP-2 cells during incubation with *K. oxytoca* cytotoxin. HEP-2 cells were cultured in 1 ml MEM medium supplemented with 5% (v/v) foetal bovine serum in the presence and absence of the cytotoxin (8 µg ml⁻¹) in a 24-well plate. Data are means of three determinations ± standard deviation. ATP levels in HEP-2 cells were measured as described in Methods. ○, ATP in HEP-2 cells cultured in the absence of cytotoxin; ●, ATP in HEP-2 cells cultured in the presence of cytotoxin.
heterogeneity of HEp-2 cells in susceptibility to the cytotoxin.

Cell-rounding seems to be a critical morphological change leading to cell death. However, it did not occur so rapidly and cells were irreversibly affected before it. This suggests that specific changes in cell metabolism might induce the early period of incubation. To examine the effects on cellular energy metabolism, intracellular ATP levels were determined during the course of incubation with the cytotoxin (Fig. 1c). The level of ATP increased slightly in the early period, but started to decrease at 6 h incubation. It decreased to 15% of the ATP level in a control culture after 24 h. Its decrease is similar to appearance of round cells both in initiation time and rate. Therefore, the decrease in the ATP level could be associated with the morphological change but could not be a primary effect of the cytotoxin.

Inhibition of macromolecular synthesis

The effects of the cytotoxin on macromolecular synthesis by HEp-2 cells were examined. Monolayers of nonsynchronized HEp-2 cells in a 96-well plate were used to assay the incorporation of [3H]uridine and [3H]leucine, while cells synchronized at the S phase were used to assay the incorporation of [3H]thymidine with efficient labeling of DNA on the microculture plate. Fig. 2 shows the dose-dependent inhibition of macromolecular synthesis in HEp-2 cells by the cytotoxin. Protein synthesis remained virtually unaffected with concentrations of up to as much as 5 µg per well and was inhibited only slightly with 10 µg per well. In contrast, both DNA and RNA syntheses were significantly inhibited with 5 µg or less per well. The inhibition of DNA synthesis was especially marked, the inhibitory effect of 1 µg of the cytotoxin on DNA synthesis being comparable to that of 10 µg of the cytotoxin on RNA synthesis. This suggests that the cytotoxin primarily inhibits DNA synthesis, and the inhibitions of RNA and protein syntheses are secondary effects.

Time course of DNA synthesis inhibition by the cytotoxin

Synchronized HEp-2 cells at the S phase in a microculture plate were incubated in the presence and absence of 2 µg cytotoxin per well, examining inhibition of DNA synthesis and cell-rounding at various times. No significant difference in the time course of cell-rounding was observed between synchronous and nonsynchronous HEp-2 cells (data not shown). The incorporation of [3H]thymidine in the presence of the cytotoxin was inhibited to approximately 45% of the level in the absence of the cytotoxin as early as 1 h after incubation was started (Fig. 3). The inhibition continued during
subsequent incubation, the incorporation gradually decreasing and finally stopping, indicating that the cytotoxin exerts its inhibitory effect on DNA synthesis within 1 h. On the other hand, cell-rounding was significantly observed only after 6 h incubation, as with nonsynchronous cells (Fig. 1a).

**Analysis of K. oxytoca plasmids**

Many virulence factors of bacteria are encoded by plasmids. Therefore, we extracted and analysed plasmids from the three clinically isolated strains of K. oxytoca produced the cytotoxin. Fig. 4 shows the plasmid profiles for strains OK-1, KA-1 and KA-2 on agarose gel. Strain OK-1 carried 34 kb and 3.5 kb plasmids, and strain KA-2 carried a 3.5 kb plasmid, whilst strain KA-1 carried no plasmids. Therefore, it can be concluded that the cytotoxin is likely not encoded by plasmids, but is probably chromosomally determined.

**Heat stability of the purified cytotoxin**

There was a possibility that the cytotoxicity might be due to the presence of a trace amount of heat-stable endotoxin in the purified cytotoxin. In order to rule out this possibility, the sensitivity of the purified cytotoxin to heat was examined. The cytotoxic activity decreased to 50% of the initial level on heating at 70°C for 20 min. Heating at 90°C for 20 min completely abolished the cytotoxicity (Fig. 5). This clearly indicates that the cytotoxin is heat-labile, eliminating the above possibility.

**Chemical analysis of the cytotoxin**

It was also possible that low-molecular-mass substances derived from peptidoglycan, such as the tracheal cytotoxin of B. pertussis, were responsible for the cytotoxic activity of the purified cytotoxin. Chemical analysis, however, revealed that neither amino acid nor amino sugar was a component of the cytotoxin. In a previous study (Minami et al., 1989), we analysed the cytotoxin purified by HPLC and tentatively estimated its molecular mass to be at most 651 Da on the basis of mass spectrometry results. However, this preparation was not suitable for accurate and precise chemical analysis by NMR spectroscopy. In order to remove possible impurities which could not be separated from the cytotoxin by the column chromatography routinely used, the cytotoxin fraction obtained on HPLC was separated into chloroform-soluble and chloroform-insoluble fractions. When the soluble fraction was developed on a silica gel thin layer plate with a solvent system of chloroform

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*Fig. 4. Analysis of plasmids in K. oxytoca strains by agarose gel electrophoresis. Lanes: 1, K. oxytoca ATCC 13182; 2, K. oxytoca OK-1; 3, K. oxytoca KA-1; 4, K. oxytoca KA-2. Molecular sizes are indicated on the left side of the gel. Reference plasmids: pUC19 (2.7 kb), pBR322 (4.4 kb), pSK201 (15.0 kb, Katayama et al., 1990), RP4 (54.5 kb), and R100 (106 kb).*

*Fig. 5. Heat lability of the K. oxytoca cytotoxin. Lyophilized cytotoxin was dissolved in Dulbecco’s PBS without Mg²⁺ and Ca²⁺, pH 7.4, and then heated at various temperatures for 20 min. The untreated control had a cytotoxin titre of 20 CD₅₀ per well. Cytotoxic activity was measured as described in the text.*
ethanol (24:1), a single spot, of which the $R_f$ value was 0.25, was detected. This soluble fraction showed cytotoxicity toward HEp-2 cells, while the insoluble fraction showed neither cytotoxicity when used alone, nor an additive effect when used together with the soluble fraction. The chloroform-soluble fraction was subjected to NMR and FAB-mass-spectroscopy to gain more insight into the chemical structure and molecular mass of the cytotoxin. Before subjecting the cytotoxin to $^1$H and $^{13}$C-NMR spectroscopies, $^{31}$P-NMR analysis was performed to confirm that no phosphorus atoms were present in the cytotoxin. The $^1$H-NMR spectrum of the cytotoxin indicated the presence of eleven protons ($H_{11}$), i.e. four olefinic ($4 \times -CH=,$), three heteroatom-linked aliphatic ($CH+CH_2$), and four normal aliphatic ($2 \times CH_2$) and rather intense exchangeable protons. The $^{13}$C-NMR spectrum of the cytotoxin indicated the presence of eight carbons, i.e. three CH$_2$s, one CH, and four =CH$_s$. Two-dimensional heteronuclear correlation analysis of the cytotoxin indicated that all of these carbon signals were reasonably correlated with the observed proton signals, except for the exchangeable proton signals, thereby establishing the C$_8$H$_{11}$ unit. Thus, at the present time, the number of carbon atoms could be unambiguously determined by NMR, whereas the exact numbers of protons and heteroatoms could not, because of the presence of exchangeable OH and/or NH protons. However, the FAB mass spectrum of the cytotoxin indicated that $M^+$ was 217; six molecular formulae satisfied $m/z = 217$, among which only one, C$_8$H$_{15}$O$_4$N$_3$ (exact mass = 217-1062), contained the eight carbons determined by NMR spectroscopy. The chemical structure of the cytotoxin, therefore, should comprise the combination of C$_8$H$_{11}$ and H$_2$O$_{18}$N$_3$ (= C$_8$H$_{15}$O$_{18}$N$_3$-C$_8$H$_{11}$) units, but a final conclusion must await future studies involving, e.g. X-ray crystallographic analysis or chemical synthesis.

**Discussion**

Previously, we showed that *K. oxytoca* cytotoxin causes cell-rounding of tissue culture cells such as HEp-2, CHO, HeLa, and Vero cells. The results presented in this paper indicate that cell-rounding is a process leading to cell-death with energy metabolism suppressed. That HEp-2 cells were irreversibly damaged before the manifestation of the morphological change implies that cell metabolism should be critically affected at an early stage. Although energy metabolism and macromolecular synthesis were all inhibited by the cytotoxin, DNA synthesis was inhibited most prominently. Furthermore, inhibition of DNA synthesis occurred in the early period when the irreversibility of the cytotoxicity was established. Therefore, the cytotoxic effect on HEp-2 cells seems to be primarily the consequence of the inhibitory effect of the cytotoxin on DNA synthesis.

The cytotoxin is not produced by *K. oxytoca* type strain ATCC 13182 (Minami et al., 1989), suggesting that a gene encoding enzymes involved in its synthesis might be on a plasmid. In *K. oxytoca* OK-1 there were two plasmids, 34 and 3-5 kb in size. This strain produced the cytotoxin even after curing of the smaller plasmid by treatment with ethidium bromide. In addition, a plasmid-free strain, KA-1, produced the cytotoxin similarly to the other strains. Therefore, the cytotoxin is probably chromosomally encoded, although it is still uncertain why some strains can but others cannot produce it.

Analysis of the purified cytotoxin by NMR and FAB-mass-spectrometry allowed the determination of its molecular mass and formula. The extremely low molecular mass and simple molecular formula of the cytotoxin are unexpected considering the sizes and structures of bacterial cytoxins reported previously. We speculate that such a low-molecular-mass toxin or related compounds could be produced by many other enteric bacteria, and be responsible for their pathogenicity in the intestinal tract. A further detailed study on the chemical structure of the *K. oxytoca* cytotoxin might be very useful for understanding the pathogenesis of potentially pathogenic bacteria causing opportunistic infections as well as the virulence of *K. oxytoca*. The examination of chemical groups by means of NMR spectroscopy and crystallographic analysis is now in progress.

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**References**


