Scanning electron microscopy and antibiotic sensitivity of the actinobacterium, Kocuria sediminis DDK6

Ashraf Y. Z. Khalifa1,2*
1Biological Science Department, Faculty of Sciences, King Faisal University, Saudi Arabia.
2Botany and Microbiology Department, Faculty of Sciences, Beni-Suef University, Beni-Suef 65211, Egypt.

1. INTRODUCTION

Kocuria is a heterogeneous bacterial genus and widely distributed in diverse habitats [1]. Members of Kocuria are spherical-shaped cells that form colonies with different pigmentation such as red, orange, pink, yellow or cream. Kocuria comprises 20 species and is taxonomically classified as Actinobacteria. Generally, kocurial species are harmless bacteria that can be found in human [2], sediments [3], soil [4], water, [5] and food [6] or associated with plants [7]. Significant ecological and environmental roles of Kocuria spp. have been documented in many previous studies. Cleaning up environments from hazardous aliphatic and aromatic compounds has been reported [8-9]. Furthermore, production of probiotics, bioactive and plant-growth promoting compounds are examples of the beneficial roles of Kocuria spp. [10-13]. Nonetheless, certain members of the genus Kocuria are human pathogens such as K. kristinae [14], K. palustris [15], K. marina [16], K. rosea [17], indicating the spectrum of human infections is expanded.

Like many other bacterial diseases, antibiotics are powerful means in fighting infections caused by virulent Kocuria spp. Killing or preventing reproduction are the two strategies that antibiotics exert their actions against bacterial pathogens. Yet, emergence of antibiotic resistance has become a catastrophic crisis that threatens the outstanding benefits that gained from wise-use of antibiotics worldwide. Misuse and overdose are the two main causes that lead to curbing the effectiveness of antibiotics [18]. Resistance to antibiotic could emerge from acquiring susceptible strains to resistance genes via lateral gene transfer. Additionally, bacterial cells could exhibit tolerance to antibiotic agents because they already have intrinsic resistance genes [19]. Antibiotic categories, mechanisms of action and resistance in both Gram-positive and Gram-negative bacteria have recently been reviewed [20]. The scanning electron microscope is an easy and suitable tool to study the external features of microbial cells in their natural habitats at high magnification and resolution. Many studies have been conducted on known Kocuria spp. to investigate cell shape and arrangement using light microscopy and SEM, and ultrastructure using transmission electron microscopy, [21-23].
2. MATERIALS AND METHODS

2.1. The bacterial strain

The DDK6 strain was previously isolated from a diesel-contaminated soil; details about isolation are previously described [9]. Briefly, 0.5 gram of soil sample was enriched with diesel oil (1% v/v) as a single energy and carbon source, in a 200 ml conical flask containing 50 ml mineral salts (MS) medium with the composition: 1 g (NH₄)₂SO₄, 0.8 g K₂HPO₄, 0.2 g KH₂PO₄, 0.2 g MgSO₄·7H₂O, 0.1 g CaCl₂·2H₂O, and 5 mg FeSO₄·7H₂O in 1 L distilled water, pH 7). Flasks were incubated at 30ºC with shaking for one week. After incubation, subsequent transfer of inoculum (3%) into a fresh MS medium was performed and single colonies were obtained upon streaking on MS plates.

2.2. Scanning electron microscopic examination

For scanning electron microscope examination, a single colony of DDK6 was fixed in a 2–3% Glutaraldehyde solution for 3–4 h at room temperature, and then washed three times in 0.1 M phosphate buffer for 15 min per wash. Next, the bacterial cells were dehydrated in a graded ethanol series as follows: 35%, 50%, 70%, 80%, 95%; three changes at 100% for 15 min each, and a final wash in acetone for 5 min. Cells were then washed in distilled water, pH 7. Cells were dried in CO₂, mounted on aluminum stubs, and sputter-coated with gold. Cells were examined and photographed using a scanning electron microscope (Joel) attached to a computer.

2.3. Intrinsic antibiotic resistance of the strain DDK6

DDK6 was further tested for its susceptibility to 10 different antibiotics. Fresh bacterial culture of DDK6 was spread homogenously onto nutrient agar plates. Commercially available antibiotic discs of piperacillin (PRL 100µg), clindamycin (DA 2µg), nitrofurantoin (F 300µg) amikacin (AK 30µg), imipenem (IPM 10µg), ampicillin/sulbactam (SAM, 20µg) cephradine (CE, 30µg), chloramphenicol (C, 30µg), erthromycin (E, 15µg) and tetracycline (TE, 30µg) were carefully placed on the inoculated plates under aseptic conditions. Plates were incubated at 30ºC for 24h. Observation of inhibition zone around the antibiotic discs highlighted that the DDK6 is sensitive to the antibiotic tested at the concentration applied. The average diameter of the zone was estimated for each antibiotic disc. The experiment was carried out in triplicate.

3. RESULTS AND DISCUSSION

*K. sediminis* strain DDK6 was isolated from diesel-contaminated soils from a petrol station in Al-Hofuf, Saudi Arabia. The cultural, biochemical and molecular characteristics were recently described [9].

3.1. Scanning electron microscope

Application of the scanning electron microscope is microbiology provides deep insights and understanding of many aspects of microbes such as cell shape, arrangements, cell-cell communication and division. SEM is a feasible and powerful tool for investigation of the morphological characteristics of microbial cells in their natural habitats at high magnification and resolution. Taking advantage of the SEM, DDK was investigated in the present study. Cells of the strain DDK6 appeared as typical cocci with a diameter of (0.7 - 0.9 µm) and arranged in pairs (Fig.1 A and B), tetrads (Fig.1 C and D) and grapelike clusters (Fig.2 A - D). These diverse arrangements were consistent with that expected to different species of *Koreura* such as *K. palustris* and *K. rhizophila* [24], *K. marina* [25], *K. aegyptia* [26], *K. sediminis* FCS-11T [3]. The surface of the strain DDK6 (Figs 1 and 2) seemed to be relatively smooth in texture. This finding confirmed the characteristic feature of Gram-positive bacteria; they have smooth surfaces. DDK6 is a Gram-positive bacteria and not an exception. Occasionally, some cells appeared with protrusions of the surface this could be artifact occurred during fixation and dehydration processes of the sample. Another characteristic feature that was observed on the surface of the DDK6 is the appearance of ridges (arrowed in Fig. 1 B, Fig.2B), that run along the circumference of the cell. As can be seen in Figure ridges could divide the cell surface into two parts. Such ridges were observed in Gram-positive bacteria such as *Bacillus subtilis* and *Staphylococcus haemolyticus* but not in Gram-negative bacteria (*Escherichia coli*) [27], [28]. Such differences in the ridges appearance are attributed to the chemical composition of the cell walls of Gram-positive and Gram-negative bacteria. The outer membrane is a unique feature of the later. The area separated by the ridge formation is likely to be the surfaces of the old and the newly formed cells indicating a peculiar mechanism of cell wall formation during division of the DDK6. Viewing cells of strain DDK6 over a small scanning area showed that rings with two perpendicular lines in tetrads indicating division septum (Fig. 1C). The observation of the circumferential rings in different angles indicates different planes of divisions. This ring is likely to be the Z ring which is formed by polymerization of a highly protein called FtsZ, at the cell division site. Thus, the main function of Z ring is to mark site of the cell division [29]. The existence of a diverse cell arrangement of the strain DDK6 is likely to be an advantage to best suit its lifestyle and occurrence in divers environmental niches.

Recently, it has been reported that spatial regulation of bacterial cell division is species-specific. To ensure the proper and equal partitioning of DNA into daughter cells, bacterial species adjust the position of the cell division site to cope with the mode of living and the habitat in which they live [30]. Additionally, the depth of the circumferential rings could differentiate between bacterial strains based on the antibiotic susceptibility in certain bacterial species such as *Staphylococcus aureus* exposed to the vancomycin [31]. Whether or not this happens to the DDK6 needs further investigation.
Fig. 1: Scanning electron micrographs showing the arrangements of the DDK6 cells: A, cells in pairs, dividing cells with shallow groove (long arrow) cell connections (short arrow), B, cells in pairs with deeper groove (short arrow), C, cells in tetrads and D, cells in cubes. Magnification and scale bar are written at the bottom of each image.

Fig. 2: Scanning electron micrographs showing the grape-like arrangements of the DDK6 cells: A, cells in clusters with relatively smooth surfaces, B, cells within circumferential ring indicating the division plane, pairs with deeper groove (short arrow), C, cells in tetrads and D, cells in cubes. Magnification and scale bar are written at the bottom of each image.
3.2. Intrinsic antibiotic resistance

In general, DDK6 strain was sensitive to antibiotic tested at the applied concentration, except for the nitrofurantoin (F 300 µg). The diameter of the inhibition zone ranged between 0.85-1.3 cm highlighting variations in susceptibility levels to the antibiotic tested. The sensitivity of the strain DDK6 to the antibiotics used in the current study is accordant with those obtained by Ma et al. (2005) [31] who reported that K. kristinae was sensitive to clindamycin, erythromycin cloxacinil, linezolid, penicillin, trimethoprim/sulfamethoxazole, vancomycin and levofloxacin. Additionally, the majority of 219 strains belonged to Kocuria spp. and K. marina. Unlike what expected, no β-lactamase was observed by these strains [32] indicating an alternative mechanism of ampicillin resistance other than production of β-lactamase in Gram-positive bacteria. Resistance to nitrofurantoin could be used, in addition, to many other phenotypic traits for identification of Kocuria spp.

The antibiotic mechanism of actions varied in targeting vital components in the bacterial cells (Table 1). Certain antibiotics such as cephradine could inhibit cell wall synthesis via inactivation of penicillin-binding proteins and cross-linking of the peptidoglycan. Consequently, bacterial cell lysis occurred. A combination of two different antibiotics such as ampicillin/ sulbactam (SAM) has a synergistic effect in bacterial growth inhibition. In this combination, the role of sulbactam is to decrease bacterial resistance mechanisms to ampicillin. Additionally, key molecules that are involved in the protein synthesis, such as 30S ribosome and 30S ribosome, could be targets of aminoglycosides (e.g., amikacin) 30S ribosome Lincosamides (e.g., clindamycin), respectively [20].

Nitrofurantoin exerts its action against susceptible bacterial strain via interfering with DNA synthesis [20]. Nevertheless, resistant strains develop chromosomal or plasmid-interceded mechanisms that involves inactivation of the of the nitrofuran reductase [20], [36], preventing the subsequent damage of molecules involved in DNA synthesis. Whether or not such mechanism is mediated by the strain DDK6 to nitrofurantoin requires further investigations.

In conclusion, DDK6 could adjust the position of the cell division site to cope with the habitat in which it lives, and resistance to the nitrofurantoin could be used as a rapid identification of Kocuria spp, in addition, to many other phenotypic traits.

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6. CONFLICT OF INTERESTS

The author declares that there is no conflict of interest.

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