The Prophylactic Role of Lipopolysaccharide of *Pseudomonas* aeruginosa Against Corneal Infection Pseudomonas aeruginosa الدور الوقائي لمتعدد السكريد الشحمي لـ ضد خمج قرنية العين

May T. FlayyihRasmyia A. Abu-rishaJinan M. HasanCollege of Science/ University of Baghdadمي طالب فليحكلية العلوم / جامعة بغداد

Abstract

The lipopolysaccharide (LPS) was extracted by using digestive enzyme and hot phenol water method from Pseudomonas aeruginosa P (10) a local isolate from patient suffering from eye infection-and partially purified by gel-filtration chromatography. The sera were prepared in rabbits, they were injected with five doses over 70 days of different concentrations of partially purified LPS. The pathological infection was done by using two groups of female mice; the first group represents the non-immunized group while the second group represents the immunized group. The left and right eyes of the two groups were scratched then the left eyes were infected with P. aeruginosa P (10) while the right eyes serve as a control scratched eye. In the second group that represents the immunized group, the mice were immunized intraperitoneally (i.p.) with sera after and before scratching and infection with P. aeruginosa. The results have shown that the typical pathological scores were observed in the left eyes of nonimmunized mice group 72hr. after infection with P. aeruginosa while no pathological scores were observed in the left eyes of immunized mice. This indicated that the immunization with LPS gave a protection against P.aeruginosa corneal infection. These results indicate that immunization with LPS reduces the corneal infection caused by P. aeruginosa.

المستخلص

تم استخلاص متعدد السكريد الشحمي م ن عزلة محلية لبكتريا Pseudomonas aeruginosa من مريض يعاني من التهاب العين بواسطة استخدام الأنزيمات الهاضمة و طريقة الفينول الساخن وتم تنقيته جزئيا بواسطة كروماتوكرافيا الترشيح الهلامي . حضرت المصول في الأرانب بخمس حقنات على مدى 70 يوما و بتراكيز مختلفة من متعدد السكريد الشحمي المستخلص والمنقى جزئيا . تم أحداث الإصابة المرضية باستخدام مجموعتين من إناث الفنران مثلت المجموعة الأولى مجموعة الفئران الغير ممنعة في حين مثلت المجموعة الثانية مجموعة الفئران الممنعة حيث تم تخديش العيون اليسرى والعيون اليمنى لكلا المجموعتين ثم إصابة الثانية مجموعة الفئران الممنعة حيث تم تخديش العيون اليسرى والعيون اليمنى لكلا المجموعتين ثم إصابة العيون اليسرى فقط ببكتريا *Aeruginosa العزل*ة (10) في حين مثلت العيون اليمنى سيطرة للعيون المخدشة فقط . المجموعة الثانية التي مثلت مجموعة الفئران الممنعة تم تمثلت العيون أليمون العيون معلو بعد التخديش و الإصابة . وقد أظهرت النتائج بعد ٢٢ ساعة من الإصابة ببكتريا *P. aeruginosa* بمصل ملاحظة الإصابة المرضية المثلى في العيون اليسرى للفئران الممنعة تم تمنيعها (داخل غشاء الصفاق) بعصل قبل و بعد التخديش و الإصابة . وقد أظهرت النتائج بعد ٢٢ ساعة من الإصابة ببكتريا مناهر ألي وأصابة في ملاحظة الإصابة المرضية المثلى في العيون اليسرى للفئران غير الممنعة فقط في حين لم تظهر أي إصابة في العيون اليسرى للفئران الممنعة وهذا يدل على إن التمنيع باستخدام عديد السكريد الشحمي قد وفر حماية في إصابة قرنيات العيون به *P. aeruginosa* ما يدل على إن التمنيع باستخدام متيات العيون م المرضي المتحمي قد وفر حماية في العيون اليسرى الفئران الممنعة وهذا يدل على إن التمنيع باستخدام متعدد السكريد الشحمي قد وفر حماية في إصابة قرنيات العيون به المتنامي المالي الما لعلى إن التمنيع باستخدام من يد المرمي الما م من إصابة قرنيات العيون به ماليون عالي الما على إن التمنيع باستخدام متعدد السكريد الشحمي قل من

Introduction

Pseudomonas aeruginosa is the most commonly encountered gram-negative species that is not a member of the family *Enterobacteriaceae* [1]. It is widely distributed in nature and it is commonly present in moist environment in hospitals. It can cause disease in humans with abnormal host defenses and colonize normal humans in whom it is saprophyte [2].

The virulence factors of *P. aeroginosa* include: exotoxins, endotoxins, and a variety of cytotoxic substances including proteases, phospholipases, rhamnolipids and the blue-green pigment pyocyanin, and alginate-like exopolysaccharide that is responsible for the mucoid phenotype [3], in addition to pili and intrinsic resistance to many antimicrobial agent [1]. Lipopolysaccharide (LPS) is the major component of the outer membrane of gram-negative bacteria; it protects the pathogenic bactera from host defenses and mediates the entry of the bacteria into eukaryotic cells [4]. Keratitis caused by *P. aeroginosa* is one of the most rapidly developing and destructive diseases of the cornea. Once the bacteria infect the cornea, complex host tissue reactions occur, including inflammation, cellular and humoral immune responses and degradation of stromal proteins [5]. It is very uncommon in a normal eye and usually only develops when the ocular defense has been compromised [6]. *P. aeroginosa* keratitis following trauma to the cornea may result in blindness [7]. During corneal infection LPS has been proposed to be a ligand responsible for the invasion of the cornea [8]. This study aims to:

isolate *P. aeruginosa* from patients suffering from corneal infection, extracting and partially purifying lipopolysaccharide and studying the prophylactic role of LPS of *P. aeruginosa* by preparing antiserum against LPS in rabbits and by active immunization in mice.

Methods

1.Bacterial Isolates

Specimen were collected by cotton swap. The collected specimen were inoculated on MacConkey agar and Cetrimide 0.03% medium then incubated at 37°C for 24 hr. The isolated colonies were identified by biochemical tests [1] and the identification was confirmed by API 20 E system and agglutination sera test for the grouping of *P. aeruginosa* (Sanofi Diagnostic Pasteur).

2. Extraction and Partial Purification of LPS of Pseudomonas aeruginosa

According to [9] the LPS was extracted and partially purified. LPS was extracted by using digestive enzymes and hot water phenol method and the purification by gelfiltration chromatography by using Sephacryl 200S (unpublished study). Protein was determined according to [10], carbohydrates were determined according to [11] and the nucleic acids were determined according to [12].

3. Preparation of antisera

Antisera were prepared in rabbits. Two rabbits (2-2.5) kg were hyperimmunized over a period of 70 days by five injections, the first two injections were given intramuscularly with olive oil as an adjuvant, on day 0, one ml of 10 μ g of LPS in olive oil and on day 14, one ml of 20 μ g of LPS in olive oil, the third, fourth and fifth injections were given intravenously on days 28, 40, 56 respectively, one ml of 50 μ g of LPS. One control rabbit was injected PBS instead of LPS. On day 70, blood was collected from immunized and control rabbits by heart stabbing and the sera was pooled [13]. Indirect passive hemagglutination was used according to [14] in order to measure antibodies titers against LPS antigens of *Pseudomonas aeruginosa* in immunized and control rabbits sera.

4.Corneal infection experiment

Two groups of mice were used, each group consisting of 4 female mice. The first group which was not injected with sera represents a non- immunized group. The second group was injected intraperitoneally (i.p) with 0.2 ml of 2×concentrated rabbit serum [11] 18 hr prior to corneal challenge and again 2 hr after corneal challenge. This group represents the immunized group [15]. Corneal infection was initiated on scratch-injured eyes of mice as described by [16]. Briefly, mice eyes of the two groups were wounded by gently scratching the corneal surface with a sterile 26-gauge needle. 5 µl of bacterial cell suspension 1×10^8 cell/ml viable organisms were delivered onto incise left corneas with a micropipette having a sterile disposable tip, whereas the right cornea of each mouse was served as a control scratching eye. Control mouse for each group has normal eyes without immunization or scratching. Generally, maximal pathology scores were observed 72h after infection [15].

Results and Discussion

-Studying the Prophylactic Role of LPS of *P. aeruginosa* by Corneal Eyes Infection of Non-immunized and immunized mice

The clinical examination show that there were no pathological scores were observed in the left or right eyes of control mice that have not been immunized, scratched or infected with *P. aeruginosa* as shown in Figures (1 A, 1 B).



Figure (1 A): Photograph of the left eve of control mouse



Figure (1 B): Photograph of the right eve of control mouse

Maximal pathological scores were only observed in the left eyes of non-immunized mice that were scratched by 26 gauge sterile needles and challenged with 5μ l of 1×10^8 cell/ml viable organism of *P. aeruginosa* P (10) isolate, 72 hr. after infection, as shown in Figure (2 A).



Figure (2 A): Photograph of the left eye of non-immunized mouse that was scratched by and challenged with *Pseudomonas aeruginosa* P (10) showing the pathological score (\rightarrow)

While no scores were observed in the right eye of the same mice that was only scratched but not infected, as shown in Figure (2 B).



Figure (2 B): Photograph of the right eye of non-immunized mice that was only scratched and not infected by *P. aeruginosa*. No pathological score was observed

No scores were observed either in the left eyes of the immunized mice that were immunized (i.p) with 0.2 ml of 2x of concentrated rabbit anti LPS sera, 18hr. before and 1-2 hr. after infection with *P. aeruginosa*, as shown in Figure (3 A) or in the right eyes that were scratched only but not infected with *P. aeruginosa*, as shown in figure (3 B).



Figure (3 A): Photograph of the left eye of immunized mouse that was immunized and infected by *Pseudomonas aeruginosa* P (10). No pathological score was observed



Figure (3 B): Photograph of the right eye of immunized mouse that was immunized and not infected by *Pseudomonas aeruginosa* P(10). No pathological score was observed

These results show that immunization with LPS against *P. aeruginosa* corneal infection gave a protection against *P. aeruginosa* corneal infections and reduced the infection, as shown in figure (3 A) that demonstrated that no pathological scores were observed in the left eyes of immunized mice while the pathological scores were clearly observed in the left eye of non-immunized mice Figure (2 A). The clinical examination of non-immunized animals challenged with *P. aeruginosa* developed a predominantly edematous response at 24h post-challeng [16], while [15] recorded that the maximal pathology scores were observed 72h after infection. LPS isolated from several strains of *P. aeruginosa* was derived either from the phenol or water phase was found to be highly immunogenic and protective in mice [17].

P. aeruginosa-induced keratitis is a sight-threatening disease, it is in large part a consequence of the inflammatory response invoked by the host [18]. The pathogenesis of *P. aeruginosa* keratitis is a maltifactorial process requiring a combination of different bacterial and host cell factors [19]. Both bacterial (e.g., LPS) and host factors released from infiltrating cells during infection are thought to contribute to a rapidly progressing liquefactive stromal necrosis [20]. The administrating LPS-

specific antibodies, either prophylactically or therapeutically may be of a value in the treatment of *P. aeruginosa* corneal infections; this inconforming with the result recorded in this study [21].

References

- Forbes, B. A.; Sahm, D. F. and Weissfeld, A. S. (2007). Bailey and Scotts Diagnostic Microbiology. 12th ed. Mosby, Inc. China. pp. 340-350.
- Brooks, G. F.; Butel, J. S.; Carroll, K. C. and Morse, S. A. (2007). Jawetz, Melnick and Adelbergs Medical Microbiology. 24thed. The McGraw-Hill Companies, Inc. USA.
- **3.** Govan, J. R. W. (2005). Pseudomonas and non-fermenters. In: Medical Microbiology. 16thed. (Greenwood, D.; Slack, R. C. B. and Peutherer, J. F. (eds)). Churchill Livingstone. China. pp. 282-287.
- **4.** Brandenburg, K. and Wiese, A. (2004). Endotoxins: Relationships between structure, function and activity. Curr. Top. Med. Chem. 4(11): 1127-1146.
- **5.** Hazlett, L. D.; McClellan, S.; Kwon, B. and Barrett, R. (2000). Increased severity of *Pseudomonas aeruginosa* corneal infection in strains of mice designated as Th1 versus Th2 responsive. IOVS. 41(3): 805-810.
- 6. Kanski, J. J. (2007). Clinical Ophthalmology.
- **7.** Gerke, J. R. and Nelson, J. L. (1977). Oral 6th ed. Elsevier Limited. China. pp. 250-260. vaccination and multivalent vaccine against
- Van Delden, C. (2004). Virulance factors in *Pseudomonas aeruginosa* keratitis. IOVS. 16(1): 76-80.*Pseudomonas aeruginosa*. In: *Pseudomonas*: Virulance and Gene Regulation. Vol.2. (Ramos, J. L. (ed)). Plenum Publishers, New York. PP. 3-46.
- **9.** Johnson, K. G. and Perry, M. B. (1975). Improved techniques for the preparation of bacterial lipopolysaccarides. CAN. J. Microbiol. 22: 29-34.
- **10.** Lowry, O. H.; Rosebrough, N. J.; Farr, A. L. and Randall, R. J. (1951). Protein measurment with folin phenol reagent. J. Biol. Chem. 193(1): 265-275.
- **11.** Dubois, M.; Gilles, K. A.; Hamilton, J. K.; Rebers, P. A. and Smith, F. (1956). Colorimetric methods for determination of sugars and related substances. Anal. Chem. 28(3): 350-356.
- Ashwell, G. (1957). Colorimetric analysis of sugar. In: Methods in Enzymology.Vol.3. (Colowick, S. P. and Kaplan, N. O. (eds)). Academic press Inc. Publishers, New York. PP.: 73-105.
- **13.** Cryz, S. J.; Fűrer, E. and Germanier, R. (1983). Passive Protection against *Pseudomonas aeruginosa* infection in an experimental leukopenic mouse model. Infect. Immun. 40(2): 659-664.
- Herbert, W. J. (1978). Passive haemagglutination with special reference to the tanned cell technique. In: Handbook of Experimental Immunology. Vol.1. 3 rd ed. (Weir, D. M. (ed)). Blackwell Scientific Publications, Great Britain. pp. 20.1 20.19.
- **15.** Zaidi, T. S.; Priebe, G. P. and pier, G. B. (2006). A live-attenuated *Pseudomonas aeruginosa* vaccine elicits outer membrane protein-specific active and passive protection against corneal infection. Infect. Immun. 74(2): 975-983.

- 16. Thakur, A.; Kyd, J.; Xue, M.; Willcox, M. D. P. and Cripps, A. (2001). Effector mechanisms of protection against *Pseudomonas aeruginosa* keratitis in immunized rats. Infect. Immun. 69(5): 3295-3304.
- **17.** Cryz, S. J.; JR; Pitt, T. L.; Fűrer, E. and Germonier, R. (1984). Protection against fatal *Pseudomonas aeruginosa* burn wound sepsis by immunization with lipopolysaccharide and high-molecular-weight polysaccharide. Infect. Immun. *43(3)*: 795-799.
- **18.** Hazlett, L. D. (2004). Corneal response to *Pseudomonas aeruginosa* infection.Prog. Retin. Eye Res. 23: 1–30.
- **19.** Lee, E. J.; Cowell, B. A.; Evans, D. J. and Fleiszig, S. M. (2003). Contribution of ExsA-regulated factors to corneal infection by cytotoxin and invasive *Pseudomonas aeruginosa* in a murine scarification model. IOVS. 44(9): 3892-3898.
- **20.** Pillar, C. M., and Hobden, J. A. (2002). *Pseudomonas aeruginosa* exotoxin A and keratitis in mice. IOVS. 43: 1437–1444.
- **21.** Preston, M. J.; Gerceker, A. A.; Koles, N. L.; Pollack, M. and Pier, G. B. (1997). Prophylactic and therapeutic efficacy of immunoglobulin G antibodies to *Pseudomonas aeruginosa* lipopolysaccharide against murine experimental corneal infection. IOVS. 38(7): 1418-1425.