Histopathologic Study in lung & kidney of mice post Infection with *Klebsiellae pneumoniae*

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Abstract:

*Klebsiella pneumoniae (k. pneumoniae)* as opportunistic pathogen caused nosocomial infections 5%–20% of cases from gram-negative bacterial sepsis, and multidrug-resistant in clinical cases a serious health. *Klebsiella* has the ability to disseminate and colonize most internal organs and the inflammation was rapidly converted from acute to chronic stages. *K. pneumonia* is a frequent cause of severe pneumonia with extensive lung destruction due to neutrophil infiltration (purulent exudate) in the pulmonary tissue, which is the pathologic lesion of bacterial pneumonia. The present experiment concerned on the histopathologic examination of lung and kidney in mice post- infection intraperitoneal with pathogenic local isolate of *k.pneumoniae*. Fifty Balb /mice as two groups: 1st group (n=20) infected with 0.5 ml (2x10⁹ CFU/ml) I/P and 2nd group (n=20) re-infected as in 1st group, 3rd group (n=10) injected I/P with PBS as negative control group. The microscopic lesions of (lung and kidney) post-infection and re-infected were examined. **Results:** at (4, 8 and 16 hours) the microscopic sections revealed acute inflammatory response, and post 7 days until 14 (re-infection) showed chronic inflammatory response. **Conclusions:** The roles of pathogenic *klebsiella* in induction acute infection post few hours which developed to chronic infection post 7 days re-infection.

**Keywords:** Klebsiella pneumoniae, infection, recurrent infection, pathogenesis of klebsiella pneumonia, histopathology, kidney, lung.
1-Introduction:

*Klebsiella* sp. is gram negative bacteria and important opportunistic pathogens causing life-threatening nosocomial infections (1). The virulence factors of *K. pneumonia* strains include the capsular serotype, lipopolysaccharide L.P.S (O-Ag), capsular polysaccharide C.P.S (K-Ag), iron scavenging systems, and fimbrial and non-fimbrial adhesins and produce protective against a disease via both humoral and cell-mediated immunity (2 & 3). *K. pneumoniae* is responsible for a variety of diseases in humans and animals, is a prominent nosocomial pathogen mainly responsible for urinary tract, respiratory tract or blood infections (sepsis) (4), resist to antibiotics (6) which are related to Biofilm formation in Klebsiella, so scientists have tended to produce vaccines or antigens against *K. pneumoniae* as some researchers in the world (5). WSK Ag was provided a good immune response better than WHKK Ags against infection, also the encapsulated *K.pneumoniae* has the ability to dissemination and colonization of most internal organs and the inflammation was rapidly convert from acute to chronic stages. Lower respiratory tract infections have always remained a cause of concern for medical care in the world associated with both community acquired and nosocomial pneumonia with high mortality rates of chronic pneumonia in untreated cases (7), even the immunocompetent elderly as well as the healthy non-elderly adults are at a risk of developing pneumonia (8 &9). Recently, pyogenic liver abscess (PLA) (16) occurs in diabetic patients with a prevalence rate from 45% to 75%. Sometimes complicated endophthalmitis or meningitis emerged in Taiwan and other Asian countries, and other continents (10 & 11). Rhinoscleroma and atrophic rhinitis (also called ozaena) are two chronic and potentially severely disturbing diseases of the upper respiratory tract, associated respectively with *K. pneumoniae* subsp (12), meningitis, necrotizing fasciitis and prostatic abscess reported as severe cases (13).

Finally, granuloma inguinale (donovanosis) is caused by uncultivated bacteria, which may belong to *K. pneumoniae* (14).

Endogenous Endophthalmitis (EE) in the USA and Asian population (15). Potential explanations for these geographic differences in clinical manifestations include host factors such as rates of diabetes mellitus, alcoholism, access to healthcare, and socioeconomic factors (17).

2-Materials and Methods:

2-1: Bacterial isolate: was obtained from Microbiology department Science college, which isolated from a case of urinary tract infection and the bacterial isolate was maintained on brain-heart infusion agar slant then transferred on MacConkey agar (37°C/24 hrs.) to study the morphology of the bacterial colonies and biochemical tests (18) gram’s stain and capsule stain also.

2-2: Culture media: prepared according to the manufactures company (Oxoid) and sterilized by autoclave in 121°C under 15p/inj for 15 minutes; which are:

1- MacConkey agar.
2- Nutrient agar.
3- Brain-heart infusion broth.
4- Triple sugar iron agar.
5- Simon citrate agar.
6- Indol.
7- Methyl red & Voges proskauer broth.
8- Urea broth.

2-3: [Laboratory animals]: were maintained in the animal house of Veterinary Medicine College for two weeks with a good conditions from nutrition and standard environmental conditions from light and temperature.

2-4: [Infective dose]: the pure culture of bacteria was estimated to prepare the infective dose by serial tenfold dilutions with sterile PBS; it was $2 \times 10^9$ CFU / ml.

2-5: [Histopathological examination]: according to (19) by using fixative formalin 10%, processing of tissue samples was done automatically, embedded in liquid paraffin and sectioned 4-5 µm and stained manually by Hematoxyline and Eosin stain (H&E stain) then examined by light microscope on 10X, 20X and 40X magnification.

2-6: [Experiment design]: Fifty of laboratory mice were divided randomly as following:

First group: (n=20) infected with 0.5 ml I.P ($2 \times 10^9$ CFU/ml)/24 hrs.
Second group: (n=20) re-infected with 0.5 ml I.P ($2 \times 10^9$ CFU/ml)/7-14 days
Third group: (n=10) injected with PBS as control group.

At early to 24 hours post infection and at 7-14 days re-infected, sacrificed the then preserved their lung and kidney tissues in 10% formalin for histopathology preparation.

3-Results:

3-1: [Pathological study]:

3-1-1: [Gross appearance]: during few hours post-bacterial infection showed congestion (red color) of internal organs (lung and kidneys), edema also in mice re-infected but in less amount of edematous fluid.

3-1-2: [Histopathology]: the sections of lung and kidney showed severe acute inflammatory changes at few hours post-infection characterized by; congestion of B.Vs (dilated and filled with blood), presence of edematous fluid in interstitial tissues, fibrin and infiltration of PMNs (neutrophils), (Figure: 1) at 16 hours liquefactive necrosis also noted (figure-2) that agreed with (1) who described acute respiratory infection post-intranasal infection and in glomerular tuft as kidney changes (20) after intra-urethral infection.
Figure 1: Histopathologic section in kidney of mouse at 8 hours post-infection; showed acute cell swelling of tubular-lining epithelia ( ), congestion of glomerular tuft and vacuolar degeneration of their endothelium ( ) (H&E stain, 40X).

Figure 2: Histopathologic section in kidney of mouse at 16 hours post-infection; revealed prominent epithelial cell swelling (star-shaped ( ), liquified necrotic tissue in peritubular and within tubules ( ). (H&E stain, 40X).
At 7 - 14 days from re-infection significant histopathologic lesions were less blood vessels congestion and fibrinous exudate with PMNs infiltrated and few lymphocytes also (Figure 3&4).

**Figure-3:** Histopathologic section in kidney of mouse at 7 days re-infection; infiltration of PMNs within glomerular tuft (arrow) (H&E, 40X).

**Figure-4:** Histopathologic section in lung of mouse at 7 days post-re-infection; showed fibrinous exudate perivascular and in interstitial & in alveoli (arrow) (H&E, 40X).
Figure-5: Histopathologic section in lung of mouse at 14 days re-infection; showed lymphocytic peribronchiolitis.

( ) (H&E stain, 10X).

4- Discussion:

The present study appeared predominant infiltration composed from polymorphonuclear cells (neutrophils) in most sections of lung and kidney organs through few hours post bacterial infection and re-infection. The histopathologic examination of lung tissue illustrate that the infection begins as congestion of blood vessels, edema and few neutrophils infiltration during early hours post infection. At few days later (week and more) the lesion progress to lobar pneumonia, consolidation of pulmonary lobules, (21) who noted that neutrophils as phagocytic cells so are critical to inhibit bacterial growth (22) who noted the chronic infection post exposure to bacterial infection again. On day 14 of infection, revealed chronic pulmonary changes in the lungs of infected mice with positive bacterial load, tissue damage (necrosis) of alveoli.

The experiments deals that the central mechanism of Klebsiella pathogenesis is the production of a LPS-rich cell surface protected germ cells against immune response (23). That evident in the present study from occurrence of bacterial infection in early time and sustained days later in re-infected mice.

However, (24) who explain that non-immunized animals may indicated a wide spectrum of severe pathological lesions post infection which upheld by (25) who demonstrated that K. pneumoniae was accompanied with massive pathological changes such as suppurative infections in lung, liver, urinary tract and brain, disseminated all over the world. Neutrophilic cells mainly in examined organs may indicated K. pneumoniae infection activated the cytokines production IL-6, IL-8, TNF-α and IL-1β which have prime role in inflammatory cells attraction (26, 27) revealed that the Enterobacteriaceae bacteria showed proliferation of mononuclear cells aggregation around congested veins and other blood vessels which also contained the inflammatory cells in their lumen.
Conclusions:

1. The capacity of *klebsiella pneumoniae* to induce infection due to had virulence pathologic agent in acute inflammatory changes post few hours intraperitoneal infection.

2. The lesions characterized by PMNs infiltration, edema admixed with fibrinous exudate later in the Lung and kidney tissues.

References:


