Investigation of Sewage and Drinking Water in Major Healthcare Centres for Bacterial and Viral Pathogens

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Abstract

Water is a major source of microbes, including pathogens that can cause critical pathological conditions and outbreak of epidemics. Due to lack of proper medical waste-management system in Peshawar, most of the waste is disposed near sewage lines which run parallel to drinking water supply increasing the chances of water contamination. This study was undertaken to examine bacterial and viral pathogens in fresh and waste water in major Health care units. Conventional culturing techniques were used to identify bacterial pathogens followed by biochemical analysis, whereas viral pathogens were detected by Polymerase Chain Reaction (PCR). Analysis of sewage and drinking water supply in major health care facilities of Peshawar city indicated that Klebsiella pneumoniae and Staphylococcus aureus were found in all water samples whereas serious health risk causing bacteria including Mycobacterium tuberculosis were also detected in some regions. Two viral pathogens, Hepatitis C virus (HCV) and Hepatitis B virus (HBV) were found in open sewage water of Khyber Teaching Hospital and Dabgari Garden (G). The presence of these pathogens in water is a serious threat to public health and the environment and calls for immediate action to enforce proper medical waste-management to eliminate the risks to human health.

Keywords: Water; Major healthcare centers; Pathogens; Public health

Introduction

Water pollution is one of the most pervasive problem afflicting people throughout the world. Waterborne illness and multiple epidemics are related to the consumption of contaminated or inadequately treated water is a global public health concern. As a developing country, Pakistan has poor water treatment system and ranked 80th among 122 nations in terms of providing good quality drinking water [1]. However, scenario is even worst in many cities of Pakistan where drinking water is unsafe for direct human consumption and severely contaminated with bacterial and viral pathogens. In Pakistan, waterborne diseases and parasitic infections due to contaminated water accounts for nearly 60% and 80% of children’s death respectively. Every year approximately 250,000 children die due to water-borne diarrhea solely and 1.2 million people get affected by waterborne pathogens in Pakistan [2]. Furthermore, scientific data and evidence considering role of waterborne pathogens in the epidemiology of hospital-acquired infections are insufficient [3]. Recent reports on pathological conditions of identified waterborne pathogens have provided novel insights into the understanding of pathology and effect of diseases [4-6], which persist in numerous aquatic systems due to the advantage of resistance to various environmental factors [7].

Health facilities, mainly health care centres, hospitals, clinics and laboratories pose higher risk of water contamination since these are more likely to be the sources of viral and bacterial pathogens [8-11]. Although numerous research has been conducted to address the detection and origination of pathogens in both drinking water and wastewater [12,13], however, insufficient studies have done explicitly about tracing the occurrence of pathogens in water sources near healthcare facilities [14,15].

Our current study was conducted to detect the presence of pathogenic microbes both in drinking and waste water samples collected from Khyber Teaching Hospital (KTH), Hayatabad Medical Complex (HMC) and Dabgari Garden (DBG/DG), the major healthcare facilities of Peshawar, capital city of province Khyber Pakhtunkhwa (KP), Pakistan. Everyday these hospitals provide health facilities to thousands of local people, patients coming from far flung areas of KP, and from Afghanistan as well. During personal visits to these hospitals for sample collection improper sewage systems allowing stagnant water retention for several days and mass of untreated disposed materials were observed, which may stir up risk of contamination in drinking water. In addition, the disposal of waste materials from diagnostic laboratories and pharmaceutical centres poses a significant threat to public health.

Inadequate information is available about sewage and drinking water quality near major health care units of Peshawar, KP, Pakistan and no investigation for pathogens has been done particularly considering water sources of the healthcare centers. Additionally, no pre-defined rules and laws presented by WHO are set and applied by healthcare management and higher authorities for such investigation and providing good quality treated water. Therefore, the focus of current study is determination of viral and
bacterial pathogens in drinking and sewage water of major health care units of Peshawar to highlight critical role contaminated water plays in waterborne diseases. Culture techniques and Polymerase Chain Reaction (PCR) as the most commonly used methods for monitoring and detection of bacterial and viral pathogens [16-19], are applied in this research.

Materials and Methods

Study site description and sampling

Khyber Pakhtunkhwa (KP) province - with population of 26.9 million and area of 74,521 square kilometer - is located in the north-western region of Pakistan and by the size of population is the 3rd biggest province of Pakistan. Peshawar (33° 99' 16' N, 71° 51' 36' E) is its provincial capital and largest city and hub of hospitals where patients come not only from all around the KP but also from neighbor country Afghanistan. It is crucial that hospitals providing health facility to thousands of patient’s everyday have accessibility to pathogen free drinking water. In this study, total of 252 drinking and sewage water samples were examined over a period of one year from January 2013 to December 2013, samples were collected 3 times a year in 2013, in January, June and October respectively. First session of sampling was completed during last two weeks of January, second in first two weeks of June and third time samplings were done during the last two weeks of October. Totally, 126 drinking and 126 sewage water samples were collected from three major hospitals of Peshawar i.e., KTH, HMC and DG, with the interval of 4 months except the rainy days. All samples were collected in sterile bottles from the premises of these health care sites (42 different sites in total, 6 samples were collected from each site, 3-drinking, 3-sewage water samples).

Filtration and DNA/RNA isolation

Water samples were taken to lab soon after collection in ice containers and filtered through sterile filter membranes, 0.22 μm, (Science laboratory, Islamabad, Pakistan) to concentrate the samples for investigation of bacterial pathogens. The whole water samples were processed through DNA/RNA isolation kit (Norgen Biotek, Canada) for the detection of viral pathogens.

Screening and selective media preparation for incubation

Screening was done for the samples collected (nutrients media) and positive samples were further tested through selective media. Selective media (Merck, Rawalpindi, Pakistan) were prepared (according to the prescription provided by Merck, Rawalpindi, PK) for the culturing of bacterial colonies (selective media shown in Table 1). Sterilized media were poured into Petri plates, followed by spreading of concentrated water samples with the help of a sterilized loop. Petri plates were incubated at 35°C for 24 h (Also 48 hours to get the correct density) and 5 weeks for MTB, followed by sub-culturing of colonies on fresh selective media at 34°C and 36°C [14]. This step of colony sub-culturing was repeated 3 times for confirmation of the resultant colonies. The colonies which showed consistent growth were noted and non-consistent growing bacterial pathogens were neglected to avoid false positive results.

Biochemical analysis

Biochemical tests conducted for identification of bacterial species grown previously on selective media were Catalase, Oxidase, Tube coagulase, Alkaline phosphatase, Motility, Arginine, 

| Bacterial Pathogens | Catalase test | Oxidase test | Citrate utilization | Lactose utilization | H2S production | Voges-Proskauer test | Nitrate test | Indole test | Urea test | Motility test | Glucose test | Malto test | Mannose test | Inositol test | Tellurite test | Sucrose test | Triple Sugar iron test | Lysine Iron agar test | Gelatin hydrolysis test | Lysozyme-hydrolysis test | Catalase test | Oxidase, Tube coagulase, Alkaline phosphatase, Motility, Arginine, |
|---------------------|---------------|--------------|---------------------|--------------------|----------------|---------------------|--------------|-------------|-----------|-------------|-------------|-----------|--------------|---------------|----------------|---------------|-------------------|------------------------|------------------------|------------------------|
| E. coli             | +             | +            | +                   | +                  | +             | +                   | +            | +           | +         |             |             | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| E. aerogenes        | +             | -            | +                   | +                  | +             | +                   | +            | +           | +         |             | +           | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| H. influenzae       | +             | +            | +                   | +                  | +             | +                   | +            | +           | +         |             |             | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| K. pneumoniae       | +             | -            | +                   | +                  | -             | -                   | +            | +           | -         |             | +           | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| S. marcescens       | +             | -            | +                   | +                  | -             | +                   | +            | +           | +         |             | +           | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| H. alvei            | +             | +            | +                   | +                  | +             | +                   | +            | -           | +         |             |             | -         | +            | +             | +              | -               | +                 | +                        | +                       | +                       |
| P. vulgaris         | -             | +            | +                   | +                  | +             | +                   | -            | -           | +         |             |             | -         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| P. mirabilis        | +             | +            | +                   | +                  | -             | +                   | +            | -           | +         |             |             | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| S. enterica         | +             | +            | +                   | +                  | +             | +                   | -            | +           | -         |             |             | -         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| S. dysentariae      | -             | +            | +                   | +                  | +             | +                   | -            | +           | -         |             |             | +         | +            | +             | +              | -               | +                 | +                        | +                       | +                       |
| P. stuartii         | +             | +            | +                   | +                  | +             | +                   | -            | +           | +         |             |             | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| P. aeruginosa       | +             | +            | +                   | +                  | +             | +                   | -            | +           | +         |             |             | -         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| S. typhimurium       | +             | -            | +                   | -                  | +             | +                   | -            | +           | +         |             |             | -         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| S. aureus           | -             | +            | -                   | -                  | -             | +                   | -            | -           | +         |             |             | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| E. faecalis         | -             | +            | +                   | +                  | +             | +                   | -            | +           | +         |             |             | -         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| A. hydrophila       | +             | -            | +                   | +                  | +             | +                   | -            | +           | +         |             |             | -         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| A. sobria           | +             | +            | +                   | +                  | +             | -                   | -            | -           | +         |             |             | -         | -            | -             | -              | -               | -                 | -                        | -                       | -                       |
| S. epidermidis      | +             | +            | +                   | +                  | +             | -                   | +            | +           | +         |             |             | +         | +            | +             | +              | +               | +                 | +                        | +                       | +                       |
| M. tuberculosis     | -             | +            | +                   | -                  | +             | -                   | -            | +           | -         |             |             | -         | -            | +             | +              | -               | +                 | +                        | +                       | +                       |

Table 1: Biochemical tests used for identification of the microorganisms at specie level.
Pyruvate, Mannitol, Sucrose and Ornithine, Esculin, fermentation of Sucrose and Lysine decarboxylase (Table 1). Sterile loop was used to pick the bacterial colony from the selective media and tested for biochemical test. The result was noted and process was repeated for three times. Only those bacterial pathogens were noted which gave same result every time.

The instructions provided by Merck science lab Rawalpindi were followed to get the results. For Lactose test color change was noted as positive after broth culture. Indole test was noted as positive by appearance of pink red layer. Red color formation after addition of alpha-naphthol+sodium hydroxide while shaking the tube for 10 minutes was an indication of positive result for Voges-Proskauer test. Green color change to blue confirmed the positive result for Citrate test. For Nitrate test the color changed into dark red within 5-10 minutes. This test was carried with the addition of N,N-dimethyl-1-naphthylamine and sulphuric acid. The Oxidase test gave positive by appearance of purple color after applying 1% tetrathyl-p-phenylenediamine dihydrochloride on filter paper. For Catalase test the oxygen bubbles demonstrated positive result. Black precipitates affirmed positive test for H.S. Appearance of reddish color during Methyl Red test confirmed positive results for presence of E. coli. The yellow color was commuted to red for urease test indicating positive result. Visualizing under microscope, a hazy zone (irregular movement) formation confirmed the positive result for motile bacteria and a single line of growth formation indicated presence of non-motile bacteria. Regain of purple color from yellow after 48 hours’ incubation confirmed positive result for Ornithine test. Maltose test showed positive result after conversion of red color to yellow color. Here phenol red was used as PH indicator. In case of Mannose test the normal red color (phenol red indicator) commuted to yellow or pink, an indication of positive result. Similarly, Inositol test was noted positive by color transformation from red (phenol red indicator) to yellow or pink. For the Trehalose test the transformation of red color to yellow affirmed positive result. For sucrose test the color change from red to yellow was observed as an indication of positive result. For acetate test the clear zone formation was an indication of the acetic acid producing bacteria so it was considered positive for Acetobacter. For Triple Sugar Iron and Lysine Iron Agar tests the color change, butt and gas production was noted to the slants and compared the information available in the list provided by science lab Rawalpindi Pakistan. Gelatine hydrolysis test was performed and the starch hydrolyzation by making clear zones in surrounding was noted for positive results after addition of iodine. Lysine decarboxylase test was noted as positive by the color change to purple. A small amount of oil was added to prevent oxygen from moving out. In Coagulase test the clot formation indicated positive result. For Pyruvate test the change of blue green color to yellow was taken as confirmation of positive result. In Arginine test the purple color was changed to yellow, which is acquired as an indication of positive result. Tellurite test was confirmed as positive by the appearance of grey color on the growing colonies. In mannitol test the red color change to yellow confirmed the positive result. Furazolidone test was performed and the resistance or sensitivity was observed and compared to the list of information provided by science lab Rawalpindi Pakistan.

RT-PCR and gel electrophoresis

HCV RNA isolation from concentrated water samples were carried out using the Water RNA/DNA purification kit (NORGEN Biotek, Canada), according to manufacturer’s instructions. The extracted RNA was reverse transcribed into cDNA. The amplified cDNA/DNA was subjected to PCR amplification. The PCR product was run on gel electrophoresis followed by observation of the obtained bands through gel documentation.

Results

Identification of bacterial pathogens

Determination of pathogenic bacteria and viruses by conventional culturing and molecular techniques, respectively, is a reliable approach for assessment of water quality. Only those pathogens were included in final results which were found throughout the year in collected samples to make sure the presence of microbes regardless of physiological effect and environmental changes. Therefore, it is concluded that these pathogens made the studied sites their permanent habitat. Out of all the samples analysed, 42% pathogens were identified in drinking water and 58% pathogens in sewage water samples. Considering overall results (drinking and sewage water samples), KTH samples were highly contaminated (40%), followed by DG (31%) and HMC (29%) with little difference in results. Common bacterial pathogens traced in drinking water samples collected from all sites indicated that KTH water being highly contaminated had 10 different pathogenic species, HMC had 6, whereas, 4 different pathogenic species were detected in DG water. However, in case of sewage water, high species diversity was observed in DG samples that was contaminated with 14 different pathogenic species, as compared to 13 and 11 different pathogenic species investigated from HMC and KTH respectively. Besides common bacterial pathogens, some other important but seldom bacterial (Mycobacterium tuberculosis) pathogens were also identified in sewage water samples. Among the identified pathogens, Klebsiella pneumoniae and S. aureus were detected frequently, as compared to Proteus mirabilis, Pseudomonas aeruginosa and Enterococcus faecalis, which were least common observed pathogens in all samples. Paradoxically, fresh water samples collected from DG had shown presence of Proteus vulgaris, and M. tuberculosis in sewage water that was present in almost 80% of all the samples collected from different locations of DG. The largest number of pathogenic bacterial species in fresh water systems was found in KTH samples, while the lowest number of pathogenic bacteria species in fresh water sources was found at HMC. However, in sewage water systems the largest numbers of bacterial species were observed at DG and the lowest numbers of bacterial pathogens were detected at HMC. A detailed list of bacterial pathogens identified in each sampling site is given in Table 2.

Identification of viral pathogens

Water samples collected from multiple sites of DG, KTH and HMC was further investigated for the presence of viral pathogens i.e., HCV and HBV. Sewage water samples collected from KTH and DG determined presence of HBV, whereas, HCV was only detected in the sewage water samples collected from KTH. However, no viral pathogens were detected in fresh water samples collected from any studied area (Table 2; Figure 1).

Comparative analysis for pathogens identified in healthcare centres

Based on type of species, comparatively more pathogens were detected in sewage water, that is, total 17 different types of pathogens were ascertained in sewage water and 11 in fresh water systems (Figure 2).
Table 2: Identification of pathogenic microorganisms in fresh water (F.W) and sewage water (S.W) samples collected from different healthcare centers. The table also illustrates the health hazards caused by the detected microorganisms and the detection methods and media used. The +, - sings indicate the presence and absence of microorganism in the specified region respectively.

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Health hazard caused</th>
<th>Detection Method</th>
<th>Selective media</th>
<th>Colony appearance</th>
<th>KTH F.W</th>
<th>KTH S.W</th>
<th>DG F.W</th>
<th>DG S.W</th>
<th>HMC F.W</th>
<th>HMC S.W</th>
</tr>
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</table>
Both HCV and HBV targeted amplicons of size 170 and 230 bp respectively, were isolated from sewage water of major healthcare centers in Peshawar. HCV was detected in water samples collected from KTH, whereas HBV was detected in sewage water samples collected from KTH and DG.

DG sewage water contains the most diverse species of pathogens while its fresh water sources contain the least pathogenic species, as shown in Figure 3.

Most frequently observed pathogens in either fresh water or sewage water samples from all sample collecting sites were klebsiella and Staphylococcus epidermidis, whilst the least common pathogens were Proteus vulgaris, Providencia, Enterobacter facalis and Mycobacterium tuberculosis (Figure 4).

The overall result of both fresh and sewage water sources confirmed that KTH samples were comparatively more contaminated than DG/DBG and HMC (Figure 5). The least number of the bacterial species in DBG makes it safer than others and it might be due the privatized sector is taking better care to dispose the materials. Although there was no cleaning and burning systems but the lower bacterial burdens in water samples indicated the better treatment of wastes comparatively the other two sectors. KTH water indicated the most risk posing among all the investigated healthcare centers. It shows that lesser attention is provided to the treatment.

Considering samples (both fresh and waste water) collected from each sample collecting site/health care units, the highest numbers of pathogens were observed in sewage water i.e., 11.77 ± 0.57. Besides in sewage water the maximum numbers of pathogens were found in DG i.e., 13.33 ± 0.66 as compared to HMC i.e., 10.33 ± 0.66 where the lowest numbers of pathogens were detected. However, in case of fresh water, the maximum numbers of pathogens were found in KTH, whilst the minimum numbers of pathogens were identified in DG (Table 2). From overall result, it is evident that KTH water is highly contaminated and inadequate for consumption having highest number of pathogenic species i.e., 10.66 ± 0.61, whilst the lowest number of pathogens identified in HMC i.e., 8.00 ± 1.09 is also not safe to use.

Discussion

For all living organisms water is the most vital and important factor of survival. Inadequate access to clean water, inappropriate water treatment and bad sanitation systems is one of the most pervasive issues distressing people throughout the globe, causing waterborne infectious diseases, cause approximately 10 million deaths per year [3,20,21]. Human health is prone to microbial risks caused by enteric viruses and bacteria [22]. Studies have shown that contaminated drinking water has been source of several critical diseases, for instance, diarrhea, nausea, Cholera, typhoid, dysentery, abdominal pain and food poisoning. Situation is even worst at health care centres, where drinking water is source of pathogens transmission showing negligence of managerial authority towards supplying properly treated water. Variant pathogens are observed in ground and surface water, flood and dam water [23-25]. Furthermore, presence of bacterial pathogens is associated with physiochemical characters and location of drinking water sites [26].

To the best of our knowledge, the present study is the first systematic analysis on water sources of healthcare centres of Peshawar, KPK, Pakistan highlighting the presence of multiple substantial bacterial pathogens in hospital’s drinking and sewage water [27]. List of variant infectious bacterial and viral pathogens identified in water samples that were present consistently throughout the year at KTH, DB and HMC are given in Table 1. The abundance of these pathogens in water sources calls out for appropriate initiatives to be taken to curb outbreak of waterborne epidemics associated with contaminated water consumption [28]. In different sites the variation is characterized by physiochemical differences of the water sources [26]. Presence of Hepatitis B and C viruses in open water sources causes death of 60% of the affected people if persists for a longer time and proliferate continuously [29,30], are associated with serious public health issues [29,31]. Most frequently reported pathogenic species considering all water samples are E. coli, S. auerus, K. pneumonia, S. typhi and P. aeruginosa [2,26,32], on the basis of current study it is suggested that consumption of such water is threat to public health.

In our analysis the pathogens investigated can cause severe health problems in humans [9,33]. Most of the bacterial pathogens detected have been reported previously to be present in common water sources or home based drinking water sources [23,26] but their presence in the water sources of healthcare centers was not considered to be investigated. Furthermore, DG fresh water sources were contaminated with one third of pathogens number to that of KTH. We suggested that this high number of pathogens might be because of the improper water supply sources where sewage water can get entered into drinking water sources because of leakages in pipelines. Interestingly P. vulgaris was only found in fresh water of DG but it was not detected in sewage water sources or other fresh water sources. Analysis of sewage water allowed us to detect diverse numbers of pathogens, the highest number in DG. The presence of K. pneumonia and S. auerus in all...
sites regardless of the water type is an indication that these are the permanent species as these were also reported in daily used water sources in surrounding regions [26]. *P. mirabilis, P. aeruginosa* and *E. faecalis* were unexpectedly found in the least sites as these are generally found in water sources of diverse locations [23,27]. In addition *P. stuartii* was interestingly found in almost all water sources, which was considered to be present at most in sewage water sources only. *A. sobria* presence was detected in fresh water of KTH too which is an unexpected result. Other bacterial pathogens (except *M. tuberculosis*) were found in diverse sites as they are generally considered to be found in water sources [26,28]. Surprisingly the viral species were detected in KTH and DG sewage water which is a threat to treatment seekers and patients care takers. We investigated that some highly pathogenic bacteria including *M. tuberculosis* were present persistently throughout the study period. Furthermore, the viral pathogens were also detection continuously throughout the year, which indicates that no proper treatment is carried to the water sources.

To our knowledge, in hospitals, fluids from diagnostic tests and laboratories are improperly disposed of allowing pathogenic bacteria and viruses to contaminate water that runs off to the tap water and sewage systems, subsequently contaminating drinking water. The investigated pathogens, however, may be present in fresh water due to the lack of management interest in providing properly treated water wiped off from the pathogenic bacterial species; and treatment of hospital wastes accurately before disposing it. At the same time the laboratories owners are not admonished to throw the wastes in open places. Other sources of bacterial contamination of fresh water are surface runoff through hospitals and urban areas, pastures and agricultural lands, leakage of sewage disposal systems and septic tanks, overloaded sewage treatment plants, disposal systems and raw sewage deep well injection [2,9,34]. Similarly, we propose that contamination of drinking water observed during the present study involves factors like cross-connections, broken or leaking pipes, back-siphonage (backflow of polluted or contaminated water, from a plumbing fixture or cross-connection into a water supply line, due to a lowering of the pressure in the line) and intermittent water supply [9,34,35], and these pathogens have made the studied sites as their permanent habitats.

Our approach offers an unbiased identification of those bacterial and viral pathogens which can lead to serious human health problems. The overall investigated pathogens in hospital’s water samples are similar to the investigations of other water sources either drinking water, dam water, flood water or sewage water in KPK [26,27], somehow, our results differ in a way that through our investigation few uncommon bacterial pathogens like *M. tuberculosis*, and exceptional viral species are also identified. The reason might be the investigation site (hospitals) and consideration of only those pathogens in the results which were found in all sample collection sites and present throughout the year.

**Conclusions**

We came up with a conclusion that if current condition continued water borne illnesses will pose serious threat to public health. Addressing existence of disease causing pathogens in water sources for instance, *E. coli, S. aureus, P. stuartii, K. pneumonia, H. influenzae*, and *P. sobira*, calls out for a tremendous amount of research to be conducted to identify robust new water purifying techniques at lower
cost, with minimal use of chemicals. These pathogens can enter into water pipelines through back-siphon age, cross-connections, broken or leaking rusted pipelines, thus intermittent water supply results in contamination of the distribution system. Hospital’s waste and patient’s fluid should be disposed of properly. It is encouraged to drink boiled water and have drinking utensils autoclaved, since most bacterial and viral pathogens cannot survive in boiled water.

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Conflict of Interest

The authors declare that there is no conflict of interests regarding the publication of this paper.

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