Isolation, characterization and efficacy of lytic bacteriophages against pathogenic *Escherichia coli* from hospital liquid waste

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Abstract. Lingga R, Budiarti S, Rusmana I, Wahyu AT. 2020. Isolation, characterization and efficacy of lytic bacteriophages against pathogenic *Escherichia coli* from hospital liquid waste. Biodiversitas 21: 3234-3241. *Escherichia coli* is known as a pathogenic contaminant bacteria in hospital wastewater hazardous to humans and the environment. Concerns about the emergence of chlorine- and antibiotic-resistant bacteria increase the urgency to find an alternative strategy to control pathogenic bacteria in hospital wastewater. One of the alternatives is using lytic bacteriophage. This study aimed to isolate, characterize, and examine the efficacy of lytic bacteriophage against pathogenic *Escherichia coli* from hospital wastewater. It isolated, characterized (plaque morphology, host range, virion electron micrograph, and sensitivity to temperature, pH, and chlorine treatments), and tested the efficacy of lytic bacteriophages in controlling pathogenic *E. coli* isolated from hospital wastewater. Five phages were successfully obtained, all of which had clear plaques (lytic phage character). Based on host range assay, most of the phages could lyse all tested *E. coli* strains but not for other species. Electron micrograph photography revealed that the phages belonged to Myoviridae. The phages showed stability in high temperature, broad-range pH, and high concentrations of chlorine treatments. Assay on phages efficacy suggested that the phages are capable of significantly reducing the *E. coli* population both in sterilized and non-sterilized wastewater. The combination of phage treatment and chlorine was more effective than single phage treatment. The efficacy test revealed that phage application in wastewater had the best result seen from cocktail treatment and a combination of phage treatment and chlorine. These results suggested that the phage can be a potential candidate for disinfection purposes.

Keywords: *Escherichia coli*, lytic bacteriophage, hospital liquid waste, antibiotic-resistant

INTRODUCTION

*Escherichia coli* is a Gram-negative bacterial species belonging to the Enterobacteriaceae family. It is a non-spore-forming rod that may or may not be mobile (some rods are flagellated, and some are not). The organism is a facultative anaerobe and ferments simple sugars such as glucose to form lactic, acetic, and formic acids. *E. coli* is not only known as a commensal of the gastrointestinal tract of vertebrates, including humans (Teinalllon et al. 2010), but also an important zoonotic pathogen with different pathogenic properties. It is characterized by particular subsets of genes associated with the virulence that identify them into distinct groups or pathogroups (Van den Beld and Reusbaet 2012). *E. coli* is often found as the most common hospital waste contaminating group.

Moreover, numerous studies reported the existence of antibiotic-resistant *E. coli* in wastewater treatment plants (Aslan et al. 2018; Budiarti et al. 2018; Diwan et al. 2010). Hu et al. (2011) also reported an increase in the Gram-negative pathogens, especially those of the Enterobacteriaceae family, from hospital waste. Anastasi et al. (2012) reported that some groups of pathogenic *E. coli* survive in all stages of sewage treatment, including the disinfection stage. They found that 56% of environmental strains carry virulence-gene belonging to intestinal pathogenic *E. coli* (IPEC) and uropathogenic *E. coli* (UPEC) pathotypes identical to those found in the final effluent from the sewage treatment plant. The results suggested that *E. coli* strains surviving wastewater treatment processes could also survive in the environment (Anastasi et al. 2012).

On the other hand, there are concerns about the adverse effects of using chemical disinfectants on the environment. The reaction between chlorine and organohalogen compounds can release Trihalomethane compounds that are carcinogenic and mutagenic (Chowdhury 2012). Also, the higher the dose of the chlorine used, the higher the probability of the formation of Trihalomethane (Herawati and Yuntarso 2017).

Nowadays, a significant health issue associated with *E. coli* is its role in the emergence and spread of antimicrobial resistance (Skurnik et al. 2015). The spread is related to the development of specific clones that acquire resistance genes, mostly via mobile genetic elements (Woodford et al. 2011).

All the problems above were the driver to find alternative methods for treating pathogenic bacteria in the
wastewater treatment plant. The application of lytic bacteriophage is an alternative strategy to combat pathogenic bacterial contamination. A bacteriophage is a virus with the ability to infect and kill its host bacteria. Because of its narrow-spectrum infection ability, the bacteriophage is unlikely to trigger resistance in untargeted bacterial strains (Gu et al. 2012). Besides, unlike chemical therapeutic agents, phage resistance may also induce losses to bacteria because phage-resistant bacteria may become less virulent in case of mutations in surface virulence factors such as lipopolysaccharide (LPS) (León et al. 2015). The present study aimed to isolate, characterize, and test the efficacy of lytic bacteriophage in controlling pathogenic E. coli isolated from hospital wastewater. Furthermore, the results of this study are expected to serve as a reference on the potency of lytic bacteriophage in controlling pathogenic bacteria in hospital wastewater.

**MATERIALS AND METHODS**

**Bacterial strains and culture conditions.**

This study used a total of 10 E. coli isolates collected from hospital wastewater. The isolates were identified for their biochemical and hemolytic activities and tested for their chlorine and antibiotic susceptibility. The list of strains used in this study is presented in Table 2. The isolates were grown at 37°C in nutrient broth (NB; Oxoid), on nutrient agar (NA; Oxoid), or NA soft agar overlays (0.75% agar).

**Phage isolation**

The phages were isolated based on the method in a previous study by Budiarti (2011). The samples were obtained from cattle farming sewage and Cisadane River near Bogor Agricultural University. A total of 20 ml of centrifuged effluent was mixed with 20 ml double-strength NB and 50 ml of each E. coli strain that had exponentially grown. The solution was incubated at 37°C, 120 rpm, for 24 h. Next, it was centrifuged (30 min, 6000 rpm), and the supernatant was filtered through a 0.22 mm cellulose acetate membrane (GE Healthcare). Spot assays were performed against bacterial lawns to check the presence of phages. Inhibition haloes were further purified using toothpicks or Pasteur pipettes and paper to isolate all of the different phages. The plaque picking was repeated until single-plaque morphology was observed. The diameters of each plaque of the isolated phages were measured and characterized.

**Phage quantification**

Bacteriophage suspensions were diluted serially in NB media. A total of 100 μl desired dilution of phages was incubated with 100 μl specific strain of E. coli at 37°C for 20 minutes to allow the bacteriophage particles to attach to the host bacterial cells. The suspension was added into a tube containing overlay medium and mixed thoroughly. The mixture was poured over an underlay plate. The plates were incubated at 37°C for 18-24 h. The plaques were counted on plates. Only plates with 30-300 plaques were considered acceptable. The titer of the original phage preparation was determined by following the formula:

\[
\text{Phage concentration (pfu/ml)} = \frac{\text{Number of plaques} \times 10}{\text{reciprocals of counted dilution}}.
\]

**Host range assay**

The phage host range was determined using the double-agar-layer plating method, as described previously by Budiarti (2011). A total of 100 μL cell culture of each strain in exponential-phase was added to produce bacterial lawns. The bacteriophage titer was adjusted to 10^9 PFU ml^-1; its bacterial hosts and 10-fold dilution series were spotted on each bacterial lawn. After 24-h incubation at 37°C, the results were observed and scored.

**Examination of bacteriophage morphology**

A phage stock solution (5 μL) was dropped onto the grid for 30 seconds and dried up using filter paper. 2% uranyl acetate solution (5 μL) was also dropped onto the grid for 1 minute and dried up using filter paper for 60 minutes. The grid was placed in a holder and left for perfect drying. The specimen was observed using a transmission electron microscope (TEM JEOL JEM-1010 model) at 80 kV, and phages were examined at 20,000-80,000 times magnification.

**Sensitivity to temperature, pH, and chlorine treatments**

Phage sensitivity to temperature, pH, and chlorine treatments was examined following the same procedure by Rattanachaikunsopon and Phumkhachorn (2012). The page sensitivity to temperature was monitored by giving various temperature treatments, i.e., 40°C, 50°C, 60°C, 70°C, 80°C, 90°C, and 100°C. A total of 100 μl 10^6 PFU/ml phage liquid stock was added into 900 μl SM buffer then placed in a microtube. The temperature treatment was carried out for 30 minutes. After heating at the assigned temperatures, the solutions were placed in an ice bath. Surviving a bacteriophage titer was assayed using the double-layer method.

Meanwhile, pH treatment was carried out using the SM buffer ranging between 1 to 12. The phage sensitivity was also examined using 1, 10, 100, 500, 5,000, and 50,000 ppm chlorine. A total of 100 μl 10^6 PFU/ml phage liquid stock was added into 900 μl SM buffer at various levels of pH and chlorine. The stock was incubated for 24 hours at 37°C.

**In vitro phage efficacy assay**

A phage efficacy assay was conducted using a single phage for various bacterial hosts. A total of 100 μl 24-h culture of E. coli in OD 1 was infected by 100 μl phage stock (titer 10^5). The plating was conducted during various incubation times, i.e., respectively 0, 2, 4, 8, 12, 16, 18, 20, and 24 hours of incubation.

**Phage efficacy in hospital wastewater**

The efficacy of phage application into hospital wastewater was examined by introducing phage stock and bacteria into the sterilized wastewater. A total of 100 μl 24-h culture of E. coli in OD 1 was infected by 100 μl phage
stock (titer 10⁸). The plating was conducted during various incubation times, i.e., respectively 0, 2, 4, 6, 8, 12, 16, 20, and 24 hours of incubation.

**Efficacy of phage-chlorine combination**

The efficacy of the application of phage-chlorine combination into hospital wastewater was examined by introducing phage stock-chlorine combination into various bacteria into the sterilized and non-sterilized wastewater. A total of 100 μl 24-h culture of *E. coli* in OD 1 was infected by 100 μl phage stock-chlorine combination for sterilized wastewater treatment. For the non-sterilized wastewater treatment, 1% of the total mixing volume of the phage stock-chlorine combination was added. The plating was conducted during various incubation times, i.e., respectively 0, 2, 4, 6, 8, and 12 hours of incubation.

**RESULTS AND DISCUSSION**

**Bacteriophage isolation**

A bacteriophage was isolated from various sources (household waste, river, chicken farm waste, and slaughterhouse waste). A total of ten isolates of *E. coli* were used in this procedure. Plaque formation on the bacterial lawn indicated the presence of phage. A total of five phages were successfully obtained, all of which were lytic phages, as characterized by the presence of clear plaques on the bacterial lawns (Figure 1). Phages were named after their bacterial-host codes, i.e., FE1A, FE1B, FE1C, FK7A, FI21. See Table 1 for the plaque characteristics. The plaque formations were observable due to the inhibition of growth and lyses of the phage-infected cells on the bacterial lawn. They were distinguished into clear and turbid plaques. They also varied in their diameters.

**Bacteriophage host range**

The specificity of bacteriophage against various bacteria was examined using the spot test method. All of the tested *E. coli* isolates were susceptible to the phages. Phages FE1A, FE1B, FE1C, and FK7A were able to lyse all tested *E. coli*. Meanwhile, only phage FI21 was unable to lyse *E. coli* isolate E1 (Table 2). Among ten *E. coli* isolates used as phage hosts in this test, there were different patterns of antibiotic resistance. All of *E. coli* isolates were resistant to Clandamycin and Rifampicin. Meanwhile, only isolates K7, I2, I18, 126, and K11 were resistant to Amoxicillin. Isolates K7, I2, I18, 126 also showed resistance to Trimethoprim-Sulfamethoxazole. All isolates were sensitive to chloramphenicol and levofloxacin (Data not shown).

All 6 phages were unable to lyse non-*E. coli* hosts such as *Bacillus pumilus*, *Citrobacter freundii*, *Photobacterium damselae*, *Salmonella* sp., and *Proteus mirabilis*.

**Bacteriophages virion morphology**

Morphological characteristics of bacteriophages were examined using transmission electron microscopy. Phage FK7A appeared to have a hexagonal head (34.16 ± 1.54 nm in diameter) with a contractile tail of 100.42 ± 1.38 nm long and 7.74 ± 2.5 nm wide. A tail knob was observed (20.03 ± 3.66) as well as a base plate. However, there was no collar nor tail appendage observed (Figure 2).

**Sensitivity to temperature, pH, and chlorine treatments**

A sensitivity test to various temperature, pH, and chlorine were conducted to test the stability of phages to physical and chemical factors. The sensitivity of the isolated phages to temperature was examined by calculating the PFU/ml of the plaque counts after treating with various temperatures for 30 minutes. The results of the temperature sensitivity test showed that some phages titer significantly reduced at 90°C and 100°C. The lowest titer reduction at 90°C and 100°C was found in phages FE1A and FE1B. Meanwhile, phage FI21 was activated at 90°C, and phage FE1C was activated at 100°C. All phage titers were slightly reduced at a lower temperature.

**Table 2. Determination of the host range of the phages**

<table>
<thead>
<tr>
<th>Bacterial Testers</th>
<th>FE1A</th>
<th>FE1B</th>
<th>FE1C</th>
<th>FEK</th>
<th>FK7</th>
<th>FI2</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>E. coli</em> isolate E1</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>E. coli</em> isolate K7</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>E. coli</em> isolate I18</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>E. coli</em> isolate I18</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><em>E. coli</em> isolate K11</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td>+</td>
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<tr>
<td><em>Citrobacter freundii</em></td>
<td>-</td>
<td>-</td>
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<tr>
<td><em>Salmonella</em> sp.</td>
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<td>-</td>
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<tr>
<td><em>Bacillus pumilus</em></td>
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<td>-</td>
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<tr>
<td><em>Proteus mirabilis</em></td>
<td>-</td>
<td>-</td>
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<tr>
<td><em>Photobacterium damselae</em></td>
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</tr>
</tbody>
</table>

**Table 1. Phages’ characteristics**

<table>
<thead>
<tr>
<th>Phages</th>
<th>Morphology of plaque</th>
<th>Diameter of plaque (mm)</th>
<th>Phage titer (pfu/ml⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FE1A</td>
<td>Clear plaque, regular-shaped</td>
<td>4.67±0.32</td>
<td>6.3x10⁷</td>
</tr>
<tr>
<td>FE1B</td>
<td>Clear plaque, regular-shaped</td>
<td>3.13±0.21</td>
<td>4.0x10⁷</td>
</tr>
<tr>
<td>FE1C</td>
<td>Clear plaque, regular shaped, with halo zone at the center of the plaque</td>
<td>3.23±0.55</td>
<td>8.6x10⁷</td>
</tr>
<tr>
<td>FK7A</td>
<td>Clear plaque, irregular shaped</td>
<td>2.77±0.25</td>
<td>1.6x10⁷</td>
</tr>
<tr>
<td>FI21</td>
<td>Clear plaque, irregular shaped</td>
<td>1.47±0.15</td>
<td>1.4x10⁷</td>
</tr>
</tbody>
</table>
In terms of stability to various pH, all phages were inactive at pH 1. Only phage FI21 was inactive at pH 2. On the other hand, all the phage titers significantly decreased at pH 12, and phage FI21 was activated. All the phages’ activities were most stable at pH 7 and pH 8. The present study also found that all phages were stable at low chlorine concentration. Phages FE1A, FE1B, FE1C, and FK7A remained active in all chlorine concentrations (50,000 ppm, 5,000 ppm, and 500 ppm). Only phage FI21 was inactive in 50,000 ppm chlorine.

**Bacteriophage efficacy in bacterial control**

A phage bacteriolytic efficacy test was conducted in vitro and wastewater treatment. For the in vitro test, all treatments were added with 10^5 CFU/ml bacterial-host and 10^6 PFU/ml phage to gained MOI 10^5. The populations of the bacterial host were counted following the Total Plate Count Method. See Figure 4 for the counting result. In this test, there was no bacterial population increase in treatment 0 to 4 hours of incubation (on the contrary, control without phage showed a constant rise in its bacterial population). After 8 to 24 hours, the bacterial population gradually increased. However, the increase was significantly different from (lower than) the control, presumably because of the number of applied phages in the treatment. In wastewater treatment, in general, all the treatments showed an increase in the population of *E. coli* at the initial hour until the 4th incubation time. Next, the bacterial population was seen to decrease during the phage treatment (on the contrary from the control) (Figure 5). This decrease suggested the activity of the phages that were capable of lysing bacterial cells in wastewater.

**Bacteriophage-chlorine efficacy in wastewater**

In the efficacy test on the phage-chlorine combination (both in sterile and non-sterile wastewater), a different population decline was observed between high and low chlorine concentrations (Figures 6 and 7). Moreover, in the treatment with chlorine concentrations of 50,000 ppm and 10,000 ppm, no bacterial population was observed in the wastewater (data not shown) because high chlorine concentration is capable of eliminating all bacterial populations in sewage. There was a significant difference in the population declines at the beginning of the treatment, wherein the sterile wastewater treatment the drop was faster than that in non-sterile one. The difference was correlated with the fact that non-sterile wastewater had higher bacterial density and diversity than the sterile one.
Figure 3. Phage titer decreasing after treated at various temperature for 30 minutes.

Figure 4. The efficacy of phages in vitro to control *E. coli* (AR1: *E. coli* E1 and phage FE1A; AR2: *E. coli* K7 and phage FK7A; AR3: *E. coli* E1 and phage FE1B; AR4: *E. coli* E1 and phage FE1C; AR5: *E. coli* I21 and phage FI21).

Figure 5. The efficacy of phages in hospital wastewater to control *E. coli* (BR1: *E. coli* K7 and phage FE1A, FK7A, FE1C; BR2: *E. coli* I21 and phage FE1A, FK7A, FI21; BR3: *E. coli* E1 and phage FE1A, FK7A, FE1C; BR4: *E. coli* K7 and phage FK7A; BR5: *E. coli* E1 and phage FE1A, BR6: *E. coli* E1, K7, I21, and phage FE1A).
Discussions

In this study, it was observed that all the phages varied in size and shape of plaques. The plaque sizes were affected by several factors, including adsorption rates, lysis time, virion morphology, and diffusion abilities. The adsorption rate negatively affected the plaque size. It was a function of the phage diffusion coefficient, which is a function of medium viscosity and the size of phage virion. The larger the virion size is, the smaller the diameter of the plaque will be. Long phage lysis time will increase the size of the plaque (although at a specific time, the lysis will decrease the plaque size). This result is because longer progeny formation time will reduce the time for phage particles to diffuse between host cells. Short lysis time and lower number of progenies will lead to a smaller plaque size (Gallet et al. 2011). The phages that were successfully isolated appeared to have virulent characteristics or a lytic life cycle, as shown by the clear plaques. Lytic phages are highly appropriate to control bacterial contamination because they will immediately kill their hosts without possibly forming lysogen in the infected cell population (Fernández et al. 2019). Phage is also known only to infect and proliferate in a suitable host cell (Budiarti et al. 2019). In this study, it was observed that all phages were specifically infected and lysed E. coli strains but not for other bacteria. The study revealed that two phages could
not lyse all tested E. coli strains. Lu and Breidt (2015) reported that §241 phage, a specific E. coli O157: H7 phage, just only infect strains that have O157 antigen, and E. coli strains to lack O157 antigen were resistant to the phage infection, regardless of the presence or absence of H7 antigen. In terms of virion trait, the electron micrograph indicated that the virions of the phages belonged to Myoviridae of the Caudovirales order. Ackermann (2009) classified phage with a tail surrounded by sheath into Myoviridae. Family of Myoviridae are virulent phage, so that not integrated their genetic material to the host (lysogeny cycle) but kills the host immediately (lytic cycle) (Suttle 2005). All of these are essential requirements in the use of phage to control pathogenic bacteria.

Temperature plays an essential role in the stability of phage protein (Budiarti et al. 2019). They found that Salmonella-phages are stable at 27°C to 30°C for 30 minutes incubation and become unstable at 45°C, 55°C, and 60°C. Rattanachaikunsonpon and Phumkhachorn (2012) also reported that phages would generally be inactive at 80°C. Furthermore, Basdew and Laing (2014) stated that high-temperature treatment could reduce phage activity. Atamer et al. (2010) reported that electron microscopic analysis of Lactococcal phage revealed that thermal treatments cause morphological changes on phage particles. Nevertheless, their findings also suggested that heat resistance of phage P680 (heat-resistant Lactococcal Phage) is based on both the increased stability of phage DNA packaged in the phage capsids and enhanced stability of phage particles. This mechanism presumably was what happened during the examination of the unusual phage stability to high temperature in this study.

It is crucial to examine the effect of different pH on the viability of the isolated phages to be able to handle the preservation, usage, and application of the phage correctly throughout phage treatment. The effect of pH on reducing phage activity was through the inhibition of the attachment process, formation of the particle aggregates, and attachment of the phage particles to media or phage packaging (Taj et al. 2014). Phage stability at different pH was also determined by chemical composition in the buffer. In this study, sodium magnesium buffer was used (SM buffer). Budiarti et al. 2019 found that phage had the best stability in SM buffer at 27°C or 4°C. Meanwhile, the present study revealed that the stability of phages to various concentrations of chlorine is useful information for phage application in wastewater treatment containing chlorine. Kanna (2015) reported that phage is more resistant to free chlorine than any other group of viruses. Besides, Zhong et al. (2017) suggested that viruses can develop resistance mechanisms against disinfectants that target proteins and nucleic acids.

The results of this study suggested that phage treatment both in sterile and non-sterile wastewater is capable of reducing the bacterial population. The most significant result was the phages that were stable when combined with chlorine. These results could become one of the alternatives to control pathogenic bacteria in wastewater, i.e., by combining the two treatments. By doing so, the efficacy of phage treatment can be improved as long as numerous constraints about the number of applied phage titers and application time are identified. Periasamy and Sundaram (2013) claimed that as the time of the incubation increase, the host population in lactose and sewage water samples would also increase. In contrast, in other treatments, the increase was not that significant. In this study, while the target population increased in phage treatment samples, the increase was not that high, presumably due to the adsorption of the phage particles that might change the metabolic rate of the target pathogens. In un-inoculated treatment, there was a steady increase in the target population, indicating that the phages reduced the number of the target population. The ratio between phages and the number of bacterial target populations also influenced the results. To control the bacterial population, applying more phages will generate a better result. Also, the minimum phage concentration required to achieve antibacterial efficacy changes when the bacterial cell concentration is different, while the phage input concentration is constant (Abedon 2016). The potential application of the phage-chlorine combination in controlling pathogenic bacteria in liquid waste will be enormous because of the concern about bacterial resistance to the chlorine disinfection process in the sewage treatment plant. As reported by Se’ka et al. (2001), bacterial resistance to chlorine in sewage treatment can cause disinfection failure (for example, chlorine-resistant 021N isolates).

In conclusion, a total of five phages were obtained from cattle farming sewage and Cisadane River in Bogor. The host range test showed that all phages were capable of lysing tested E. coli host only. The electron micrograph indicated that the virions of the phages belonged to Myoviridae. Phages were stable in heat, pH, and chlorine treatments and were effective in reducing bacterial growth, both in vitro and in wastewater. Their capability in lysing the bacterial host was the same when combined with chlorine, both in sterile and non-sterile wastewater.

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