REMOVAL OF E. Coli FROM GROUNDWATER AND SURFACE WATER BY USING NYLON MEMBRANE FILTRATION TECHNIQUE

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ABSTRACT
In the present study, a low-cost filter material such as Nylon was used for the disinfection of groundwater poses an excellent bacterial filtration activity against Escherichia coli (gram-negative) cultures. Textile fabrics have gained wide acceptance as a filtration membrane, and various substrates having a pore size in microns such as Nylon, polyester, and cotton have been shown to develop microbial filtration properties. In this study, a simple method to filter bacteria using nylon is presented and the solutions are prepared using Autoclave and Laminar Air Flow Chamber. For checking the bacterial concentration in the underground water, a very small volume of sample was tested by keeping it in shaking Incubator. The results revealed that nylon filters were able to decrease the concentration of E. coli from a groundwater sample, with a higher removal efficiency achieved by using two membranes and a lower efficiency by using a single membrane. This study, therefore, suggests that the Nylon membrane demonstrated antibacterial activity against E.Coli. For the disinfection of groundwater, Nylon membrane can be used as a potential alternative cost-effective filter for production of safe drinking water.

Keywords: E.Coli, Nylon membrane, groundwater, surface water, bacteria

INTRODUCTION
In developing countries, waterborne diseases have a negative impact on public health where the drinking water is of a poor quality. Escherichia coli are present in large numbers in the normal intestinal flora of humans and animals, where it generally causes no harm. However, in other parts of the body, E. coli can cause serious diseases, such as urinary tract infections, bacteremia and meningitis. The silver-impregnated porous pot filter (SIPP) can be recommended for use by rural communities as it consistently produced high-quality water that complied with the SANS 241 turbidity and microbiological limits for drinking water. Various point-of-use (POU) water treatment methods, which include bio-sand and ceramic filtration, appropriate chemical disinfection (e.g. the use of disinfectants such as chlorine and iodine), solar disinfection and natural water purifiers (e.g. Moringa oleifera) have been reported to improve the microbial quality of drinking water as well as to decrease the incidence of endemic diarrhoea caused by waterborne pathogens. The source of the microorganism was traced to the park's water supply, which had been contaminated with raw sewage and Enterotoxigenic E.coli was identified as the microorganism responsible for this outbreak. The functionalized nano fiber structures have an effect on bacterial growth was confirmed by microbiological, physic-chemical and molecular biological analyses, such as the inoculation in a nutrient agar culture medium, flow cytometry, and real-time polymerase chain reaction.

The silver-impregnated porous pot (SIPP) can be an effective and sustainable household water treatment devices/systems (HWTS) for the Southern African Development Community (SADC) rural communities, in the removal of the total concentration of bacteria from test water, manufactured using locally available materials, easy to operate and to maintain.
Decentralized point-of-use (POU) systems are possible short-term to medium-term options for improving water quality for rural communities and could be very beneficial to individuals or families who treat their own water. Waterborne pathogens and related diseases are a major public health concern worldwide. Water purification easier for the rural community, by different techniques and methods involving naturally available resources such as sand, carbon, soil, bricks and so on. Contamination of water by foreign matter such as micro-organisms, chemicals, and industrial or other wastes, or sewage are water pollution. One of the Nation's most important natural resources is groundwater and an important source of water supply for drinking, irrigation, and industrial purposes. Causes greater deterioration of the ecological system is because of pollution level acceleration.

Higher the number of fecal coliforms and fecal streptococci indicates the greater is the possibility of the presence of pathogenic micro-organisms and live in the gastrointestinal tract of humans and warm-blooded animals such as Escherichia coli, Salmonella, Shigella, Vibriones, and Hepatitis A and D. The proven antibacterial activity of synthesized copper nanoparticles against Escherichia coli established and capable of rendering high antibacterial efficacy and hence has a great potential in the preparation of drugs used against bacterial diseases.

### EXPERIMENTAL

All the chemicals were purchased from Delta Laboratories Pt. Ltd. and the Nylon membrane was purchased from PALL Corporation (Life Sciences). The ingredients of different chemicals are tabulated Table-1.

<table>
<thead>
<tr>
<th>Table -1: Illustrate the ingredients used in the preparation of E.coli</th>
<th>Nutrient Agar</th>
<th>Luria Bertani Broth</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ingredients</td>
<td>Quantity(gm/L)</td>
<td>Ingredients</td>
</tr>
<tr>
<td>Beef extract</td>
<td>3.0</td>
<td>Yeast extract</td>
</tr>
<tr>
<td>Peptic digest of animal tissues</td>
<td>5.0</td>
<td>Casein enzymic hydrolysates</td>
</tr>
<tr>
<td>Agar</td>
<td>15.0</td>
<td>Sodium chloride</td>
</tr>
<tr>
<td>Final pH(at 250C)</td>
<td>6.8±0.2</td>
<td>Final pH ( at 25°C)</td>
</tr>
</tbody>
</table>

Luria Bertani Broth, Nutrient Agar, Distilled Water, Agar-Agar type-I, Cotton plug; Micropipette, Ethanol, Loop, etc. were used as per the requirement. Underground Water and E.Coli culture were used accordingly. Specifications of the materials used are Plastic Petri dish (9 cm dia.), conical flask (250 ml), and Test tube (10 ml), Micropipette Tips (1 ml), Nylon 6, 6 Membrane (0.45 micron) (Fig.- 2).

![Plastic Petri dish (9 cm dia.), conical flask (250 ml), and Test tube (10 ml), Micropipette Tips (1 ml), Nylon 6, 6 Membrane (0.45 micron).](image)

Fig.-1: Plastic Petri dish (9 cm dia.), conical flask (250 ml), and Test tube (10 ml), Micropipette Tips (1 ml), Nylon 6, 6 Membrane (0.45 micron).
Plate Dish
Figure-1 shows the details of a plastic Petri dish (9 cm dia.), conical flask (250 ml), and test tube (10 ml), micropipette Tips (1 ml), nylon 6, 6 membranes (0.45 micron). Petri dishes are used to culture cells such as bacteria or small mosses. And it is reused after sterilization (via an autoclave at 121 °C for about 15-20 minutes in the case of moist heat sterilization or one hour's dry-heating in a hot-air oven at 160 °C.

Nylon 6, 6 Membranes
The separation of microorganisms has been done using the pressure-driven membrane processes. In this paper, the preparation and characterization of a novel nylon 6, 6 nanofibres membranes and applied it for filtration of Chlorella vulgaris broth. Phase inverted polyvinylidene fluoride (PVDF) membrane is used to compare its performance. Filterability test of both membranes and their harvesting efficiency were conducted. Results show that nano fiber membrane is more hydrophilic (contact angle of zero), and has 45% higher surface pore size and 20% surface pore population that contribute significantly into its higher clean water permeability (of 1018 and 493 l/m2hbar for nano fiber and PVDF membranes respectively). From the filterability results, the nano fiber membrane has more advantages over the phase inverted one (2-5 times higher in productivity while maintaining similar rejection of 92%). Figure-2 shows the details of the experimental setup.

Silver Activated Carbon
Silver Activated carbon is used to purify liquids and gases in a variety of applications, including municipal drinking water, food and beverage processing, odor removal, industrial pollution control, and point-of-use filters in the home. Also helps in killing bacteria, gas purification, gold purification, metal extraction, water purification, medicine, sewage treatment, air filters, decaffeination, pet litter and odor removal.

Nylon 6,6
Nylon comes from a family of synthetic polymers known as polyamide and was first introduced by Wallace Carothers on 28th February 1935. Nylon 6,6 is a polyamide made by polycondensation of adipic acid methylenediamine, and contains a total of 12 carbon atoms in each repeating unit and make polyamides suitable for plastic applications are resistance to toughness, thermal stability, good appearance, resistance to chemicals etc.

Sediment Cloth Filter
A sediment filter acts as a sieve to remove these particles and don't remove chemicals or heavy metals or make the water taste or smell better. Sediment filters can be made of a variety of materials are used to
make the sediment filters and the most common are wound string or cord, polypropylene, polyester, cellulose, ceramic, glass fiber, and cotton.

**Water flow rate**
The Fig.-3 illustrates that water flow rate of nylon membrane of pore size 0.45 micron is more as compared to the pore size 0.2 micron. Pressure drop of pore size 0.2 micron i.e 3.52 kg/cm² is more than the pressure drop of pore size 0.45 micron i.e. 2.25 kg/cm². The rate of filtration of water purifier is very less, it takes more time to filter the water and remove a high amount of *E.coli* from the groundwater sample i.e. bore well water, tap water and pond water which is passed through a nylon membrane.

![Water Flow Rate](image)

**Procedure**
Set up a water purifier (named Bio filters) and arranged a 1/2 filter in between the filter mechanisms of the filter machine. Checked its efficiency by passing water be it tap water, bore well water and pond water. The filtered water was checked in the lab for the *E.Coli* concentration. Pressure drop of nylon membrane of pore size 0.2 micron and 0.45 micron is 3.52 kg/cm², 2.25 kg/cm² respectively.

**E.Coli Bacterial Analysis Method**
Viable Count Method or Colony Forming Count (CFC) methods are used to measure *E.Coli* bacteria. Viable count method can be accomplished by techniques such as pour plating, spread plating, and most probable number method. Plate count assumes that every colony is founded by a single cell. Based on the known volume of culture/sample that was spread on plate and colonies are counted and the *E.Coli* concentration has been calculated. Prior to plating, samples are heavily diluted in order to obtain single colonies.

![Methodology flow chart](image)
Collection of Sample
Sample collected from tap water Fig.-4a, bore water Fig.-4b and from pond water Fig.-4c n the closed container with minimum air inside it. Refrigerate these samples to prevent the growth of microbes inside the sample. Do not store the sample for more than one day.

Preparation of Media for Bacterial Growth
Apparatus required for preparation of media for bacterial growth is 100mL media, 250 ml conical flask, 2 gm of Luria Broth(LB), 4gm Nutrient Agar, 100 ml distilled water, water bath, etc. Procedure: Take 100 ml distilled water in 250 ml clean conical flask. And take 2 gm of Luria broth on glossy paper. As Luria broth is hygroscopic, transfer the measured quantity immediately in the conical flask. Shake well until it dissolves completely in water. Transfer measured 2 gm of Nutrient Agar in the above solution. Heat the flask on a water bath (80-90°C) until agar dissolves completely. (Even if some lumps are remaining then they will get dissolved in an autoclave). Seal the flask with a cotton plug and cover it with the paper before placing it in an autoclave.

Sterilization of Media Using Autoclave
Apparatus required for sterilization of media is Autoclave instrument (OSWORLD LTD.) Fig.-5a. Set the autoclave to at 121°C, time 15 minute, pressure 15 lbs/cm². Cover everything to be placed in an autoclave with paper. Check the water level in the autoclave (water level must be just above to cover the perforated plate in the autoclave). Do not put glassware with water on the outside surface of it. Tighten the lid of the autoclave with the provided screws on the lid. Tight the two opposite screws at a time and the steam release knob on the lid. After checking the settings, turn on the autoclave. It will take about 1 hr to reach the desired temperature. When the temperature reaches 121°C, the countdown will start. After the end of the cycle, wait until the temperature of autoclave reaches 80-85°C, then open the lid. For that first release the steam valve, then open the opposite screws with help of metal rod. Remove the apparatus inside the autoclave using gloves. Pour the media in Petri plates immediately only inside Laminar Air Flow Chamber after autoclave, otherwise medium will get solidified inside the flask. And the sterile apparatus from autoclave should be opened inside the laminar air flow chamber only.

Laminar Air flow Chamber
Figure-5b shows Laminar Chamber and it consists of a closed chamber with two burners, laminar air flow, UV light and the door which slides up and down vertically. The dilutions, sample transfer, pouring media into petri plates demand sterile conditions. Laminar air flow chamber provides those conditions. About 15 cm diameter sphere around the burner flame is more sterile. So dilutions, transfer of media and sample, spreading are done in the area between two burners. Turn on the light and fan of the Laminar Air Flow Chamber. Clean the hands and laminar with 70% ethanol in water using cotton (the area between two burners should be cleaned properly) Light two burners. Any operation such as transfer of material should be done in between two burners only. Open the Petri plates from autoclave in such a way to minimize air exposure. Open flask between two burners. Hold the neck of flask near the burner every time you open it. Pour the media in the Petri plates. The
media should form a uniform layer inside the plate. Transfer the specific amount of the sample to the Petri plates and spread the sample using a spreader. Before spreading, the spreader should be sterilized by holding it on the burner flame. When the spreader cools then spreading should be done. After these steps, mark the plates according to the sample and do not open the lid of the plates otherwise contamination from the air will take place.

![Image of Autoclave Machine, Laminar Chamber, and Shaking Incubator]

**Fig. 5:** (a) Autoclave Machine (b) Laminar Chamber (c) Shaking Incubator

### Incubation

Figure 5c shows the shaking incubator. While lifting and placing the Petri plates, do not open the lid. Place each Petri plate separately inside the incubator. Set the temperature of the incubator as per the requirement. Let the petri plates remain inside the incubator for minimum 24 hours.

### Preparation of *E. Coli* culture

Take 100 ml of distilled water and add 2.5 g of Nutrient Broth (NB) in 250 ml conical flask (conical flask should be clean and dry). Shake well so that NB gets dissolved properly and heat if necessary. Cover the mouth of the conical flask with cotton and then with paper. Use rubber band to tighten the plug. Place the conical flask solution in Autoclave (Autoclave at temperature 121°C, the pressure of 15 lb/cm², and time 15 minute). After the Autoclave cycle is complete, remove the conical flask from it and allow it to cool down at room temperature only in Laminar Chamber. Before using laminar air flow Chamber, it should be cleaned with 70% ethanol solution. Properly dip the Loop and shake it to *E. Coli* culture, and then dip the *E. Coli* containing Loop to conical flask solution and shake it properly. Place the Conical flask solution to Incubator at 37°C and 120 RPM. Leave it for at least 12 hours to 16 hours so that fresh *E. Coli* can grow in it. After 16 hours the solution has become turbid which indicates that the *E. Coli* has growth has taken place.

### Sub Culturing of *E. Coli*

Take 100 ml of distilled water and add 2.5 g of Luria Broth(LB), 3 g of Agar-Agar Type-I in 250 ml conical flask (conical flask should be clean and dry). Shake well so that Luria Broth and Agar get dissolved properly and heat if necessary. Cover the mouth of the conical flask with cotton and then with paper. Use a rubber band to tighten the plug. Place the conical flask solution in Autoclave (Autoclave at temperature 121°C, the pressure of 15 lb/cm², and time 15 minute). After the Autoclave cycle is complete, remove the conical flask from it and allow it to cool down at room temperature only in Laminar Chamber. Before using laminar air flow Chamber, it should be cleaned with 70% ethanol solution. Then pour the solution to 4 test tubes and 2 Petri plates in Laminar Chamber only and allow it to cool for approximately 1 hour. Properly dip the Loop to *E. Coli* liquid culture, and then spread the *E. Coli* containing Loop to the surface of solidified Agar plates and test tubes. Place the Conical flask solution to
Incubator at 37°C and 120 RPM. Leave it for at least 12 hours to 16 hours so that fresh *E.Coli* can grow in it. After 16 hours, the Petri plates and test tubes have shown whitish growth on its surface which indicates that the *E.Coli* has growth has taken place. Table 3 illustrates the initial concentrations of *E.coli* spiked in bore well water, tap water and bore well water with targeted bacteria before treatment expressed in an average count of cfu/ml.

Table 4: E.Coli Concentration in bore well water, Tap water, and Pond Water sources before and after treatment

<table>
<thead>
<tr>
<th>Water source</th>
<th><em>E.Coli</em> concentrations (cfu/ml) before filtration</th>
<th><em>E.Coli</em> concentrations (cfu/ml) after filtration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 Filter Used</td>
<td>2 Filter Used</td>
</tr>
<tr>
<td>Bore well water</td>
<td>4x10^6</td>
<td>1x10^6</td>
</tr>
<tr>
<td>Tap water</td>
<td>7x10^6</td>
<td>1x10^6</td>
</tr>
<tr>
<td>Pond water</td>
<td>137x10^6</td>
<td>74x10^6</td>
</tr>
</tbody>
</table>

Developing countries have made de-centralized systems vital for the development of new technologies, especially in scattered communities depending on total nylon groundwater supplies. The study explored the use of nylon membrane which is cost-effective materials locally available for use in drinking groundwater disinfection.

**CONCLUSION**

The filtration system was used to improve the purity of the groundwater sample. Nylon membrane was tested for antibacterial activity and showed excellent antibacterial filtration performance against *E. coli*. Number of *E.Coli* was detected in the output water, based on the performance of the membrane as an antibacterial water filter system. The single membrane is not so good systems for the disinfection of drinking water because of low bacterial removal. A combined membrane system is the sole drinking-water purification system suggested. Complete anti-microbial solutions to rural communities can be an
offer by this technology. Future studies will be conducted on the coating of the nylon filter system with some cheap available antibacterial agent.

REFERENCES


[RJC-5041/2018]