Characterization of binding sites and Optimization of cell free bacteria condition for metal bio-sorbents

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ABSTRACT: Bacteria a Microscopic organisms are the most inexhaustible and flexible of microorganisms and constitute a huge division of the whole living earthly biomass, certain microorganisms were found to amass metallic components at a high limit Was Known as Bacterial Bio-sorption Due to their little size, capacity to become under controlled conditions, and their Accommodation to an extensive variety of ecological situations; Potent metal bio-sorbents among microorganisms, at low pH esteems, cell divider ligands are protonated and contend essentially with metals for official. With expanding pH, more ligands, such as amino and carboxyl groups, could be exposed, leading to attraction between these negative charges and the metals, and consequently increment bio-sorption onto the cell surface. Starting with Isolation and identification of heavy metal-resistant bacteria from rock Ore. Studying Factors Affecting Uranium Bio-sorption, Optimization of bacterial growth conditions and optimum for metal uptake by free and immobilized bacterial cells and Desorption ratio of uranium ions adsorbed by Coli. /alginate, All this evidence suggest that functions groups Represented in our study are responsible for metal uptake in our bacterial biomass beside change in peaks position which assigned for it's groups confirm bio-sorption of metal ions from waste due to ions charge interaction comparing with immobilized we found increase in no of binding sites indicate that immobilized bacterial have high efficiency for metal up take which also change in peaks position which assigned for its groups confirm bio-sorption of metal ions from waste due to ions charge interaction, Where the high bio-sorption yield obtained by
bacteria, the Uranium & heavy metal bioremediation process expects microorganisms to be joined to a strong surface.

Key words:

Binding sites, Optimization, cell free bacteria condition for metal bio-sorbents.

Introduction:

1.3. Bacterial biomass Biosorbents

Mann 1990 Reported that Bacteria a Microscopic organisms are the most inexhaustible and flexible of microorganisms and constitute a huge division of the whole living earthly biomass, whose mass is estimated as ~1018 g. In the mid 1980s, certain microorganisms were found to amass metallic components at a high limit Was Known as Bacterial Bio-sorption (Vijayaraghavan and Yun 2008a, b). Due to their little size, capacity to become under controlled conditions, and their Accommodation to an extensive variety of ecological situations; furthermore, inexpensive nutrient sources are readily available for microbes. Potent metal biosorbents among microorganisms incorporate genera Bacillus, Pseudomonas, Streptomyces, Micrococcus, and Escherichia coli. Biosorbents got from bacterial biomass have since turned out to be common. The Limits of bio-sorption not just relies upon the sort of metal particles, yet in addition on the bacterial variety, because of varieties in cellular Components. Metal particles in solution are adsorbed on bacterial surfaces through interactions with chemical functional gatherings, for example, carboxylate, amine, amide, imidazole, phosphate, thioether, hydroxyl, and other utilitarian gatherings found in cell divider biopolymers. The fast energy saw with bacterial biomass speaks to a favorable viewpoint for the plan of wastewater treatment frameworks. Bio-sorption incorporates a blend of a few components, for example, Charge fascination, composition, particle trade, covalent authoritative, van der Waal’s forces, adsorption, and micro-precipitation. Very short contact times are for the most part adequate to accomplish a metal-bacterial biomass enduring state. This is because biomass is used in the form of either fine Pellets or wet viable cells, where mass transfer resistances are usually negligible.

1.4. Bacterial Cell Constituent
The diameter of typical bacterial cells range from 0.5 to 1.0 mm; however, some are more extensive than 50 mm., despite the fact that there is an incredible assortment of shapes because of contrasts in hereditary qualities and environment. Microorganisms have basic morphology; the most widely recognized microscopic organisms are available in three essential shapes: circular or ovoid (coccus), bar (bacillus, with a round and hollow shape), and winding (spirillum. The little size of microscopic organisms guarantees quick metabolic procedures. A bacterial cell (e.g., E. coli) contains a cell divider, cell film, and the cytoplasmic network which comprises of a few constituents that are not layer enclosed (inclusion bodies, ribosomes, and the nucleoid with it's genetic material). Bacteria are classified as either Gram-positive or Gram-negative. Are Classified by the Gram staining Microbes into two principle group cell wall qualities (Beveridge 2001). This grouping partitions (Beveridge 1989; Sleytr and Beveridge 1999). Both cell divider sorts contain a peptidoglycan layer that is rich in carboxylate gatherings and totally encompasses the phone (Beveridge 1989; Langley and Beveridge 1999). Cell divider a general negative charge, because of the nearness of phosphodiester bonds between teichoic acid monomers. The exceptionally charged nature of lipopolysaccharides presents a general negative charge on the Gram-negative cell divider. The anionic useful Gram- gatherings show in the peptidoglycan, teichoic acids, and teichuronic acids of positive microscopic organisms, and the peptidoglycan, phospholipids, and lipopolysaccharides of Gram-negative microbes are the segments fundamentally in charge of the anionic character and metal-restricting capacity of the cell divider (Moat et al. 2002; The peptidoglycan layer in the Gram-positive cell divider is ca. 25 nm thick, while the Gram-negative peptidoglycan layer is much thinner (ca. 7.5 nm). The walls of Gram-positive bacteria consist of three primary components: cytoplasm mixed with peptidoglycan, to which teichoic acids are covalently bound. The envelope of Gram-negative bacteria is more complex than that of Gram-positive bacteria. It consists of two membrane bilayers (the outer and plasma membrane) that are chemically and functionally distinct from one another and sandwich a thin peptidoglycan layer between them. Teichoic acids enable the Gram-positive. Prescott et al. 2002). Extracellular polysaccharides (EPSs) are also Provide ability of binding metals; however, their availability rely on the bacterial species and growth conditions, what's more, they can without much of a stretch be expelled by basic
mechanical disturbance or concoction washing (Yee and Fein 2001). The cell dividers of bacteria contain a large number of surface functional groups, in which carboxyl is generally the most acidic group in the bacteria. At low pH estimates, cell divider ligands are protonated and contend essentially with metals for official. With expanding pH, more ligands, such as amino and carboxyl groups, could be exposed, leading to attraction between these negative charges and the metals, and consequently increment bio-sorption onto the cell surface. A few microscopic organisms have extraordinary structures, for example, flagella and the S-layer. Sumin Park and Minhee Lee 2017 Explained that The inner and outer void spaces of the Ca alginate spherioul were filled During the bio-sorption process with heavy metals such as Cu, Fe, and S, suggesting that heavy metal removal by Ca alginate beads occurs by not only ion exchange but also by Framing edifices and precipitation.

1.5. The S-layer action in bio-sorption

The S-layer action is a surface and Para crystalline envelope present in several gatherings of microbes and archaea. This layer is formed from protein or glycoprotein monomers that can self-collect in two-dimensional structures (Sleytr et al. 2003). S-layers are related with lipopolysaccharides Urrutia 1997; Madigan et al. 2000). Porosity is between 30 and 70% and the diameter of the pore between 2 and 8 nm. This characteristic can be used for metal binding. An imperative normal for this protein is its ability to reassemble once disconnected from the cell (Pollmann et al. 2006). Due to this effect, it can be used for bioremediation. S-layer proteins may execute a catching part of metallic particles in both living and dead cells, being a potential option for bioremediation of substantial metals in the field. Some bacterial cells can produce a capsule outside the bacterial cell wall. They are much hydrated and approximately organized polymers Cases are made out of starches and proteins of polysaccharides and a couple of comprise of proteins or polymers of amino acids called polypeptides ,Container course of action is imperative to metal official (Madigan et al. 2000; Channel et al. 2002). EPSs have a high sub-atomic weight with a wealth of adversely charged practical gatherings (ligands), e.g., carboxyl, hydroxyl, and uronic acids (Sobeck and Higgins 2002; Yan et al. 2008). Where the arrangement of bacterial EPS is perplexing, contingent upon the strain and its way of life conditions. EPS amalgamation is likewise announced for a few pseudomonads,
Zoogloea ramigera, Rhizobium sp., Klebsiella sp., and Bacillus sp. Ordinary constituents of EPS are for the most part polysaccharides and proteins, frequently joined by nucleic acids, lipids, or humic substances (Flemming and Wingender 2001; van Hullebusch et al. 2003).

1.6. Uranium sequestering mechanism (Bio-sorption)

Research is in progress to establish bio-sorption as a financially reasonable strategy to trap and accumulate metals. Bio-sorption can serve as a tool for the recovery of precious metals (e.g., from processing solutions or seawater) and for the elimination of poisonous metals (particularly from industrial wastewaters) (Schiewer and Volesky 2000). Adsorption and micro-precipitation involve binding of electrically neutral metals without the arrival of a stoichiometric amount of previously bound ions. In precipitation reactions, the main impetus is interaction between the solute and the solvent, whereas in adsorption affinity amongst sorbent and sorbate is the driving force. On account of physicochemical, the mechanism by which interaction based on physical adsorption, ion exchange, and complexion between metal and functional groups of the cell surface, metal binding does not depend on cellular metabolism. Tunali et al. (2006) indicate that the bio-sorption of lead and copper by Bacillus sp. involve a particle-exchange mechanism. Since the main mechanism involved in bio-sorption is ion exchange, protons compete with metal captions for the binding sites and for this reason; pH is the operational condition, which influences the process most strongly.

2. Factors Affecting Uranium Bio-sorption

There are many factors affecting the bio-sorption process of Uranium -metal ions by the microorganisms. Some of these components were recognized from the examinations of the bio-sorption procedure, for example, particle fixation, biomass concentration, time, pH and temperature, while other factors belong to types of biomass such as living or dead, free or immobilized cells, and the biosorptive capacity of cell biomass. Kerkar, S., & Das, K. R. (2017) Stated that The contact time where The obstruction between restricting locales because of expanded Biomass dosages cannot be overruled, as this will result in low specific uptake Which played a vital part in the evacuation of Zn²⁺ as (Mishra and Tadepalli 2014) Initial solute
concentration appears to have an effect on bio-sorption, with a higher concentration resulting in a high solute uptake (Öztürk 2007; Bueno et al. 2008; Uzel and Ozdemir 2009). Also in our study U bio-sorption capacity increase with increasing contact time, due to as time passed the vacant sites of biosorbent available for adsorption were occupied by the pollutant. While the initial concentration is the significant independent parameter associated with Chrome study Mishra and Tadepalli 2014 in U study increase in initial conc. associated with increase in bio-sorption % Ion removal Toxic pollutant, therefore, initially the effluent concentration was very less and then started to increase up to 30 h and then decreased Gupta, A., Balomajumder, C. (2016). The important independent parameter for Pb²⁺ and U removal is the temperature where optimum temperature for growth of bacteria ranged from 30 - 40°C. However, pH value affects strongly in the removal of Zn²⁺, Pb₂⁺ and Cr³⁺ as (Mishra and Tadepalli 2014) stated in his study also in our study it has significant effect in U removal. Bio-sorption by living cells is temperature dependent, and hence change in this parameter will strongly affect while the bio-sorption processes by nonliving biomass is not significantly affected by the temperature. Gupta, A, Balomajumder, C. (2016).

**Immobilization (capsulation) of Bacteria**

In addition to the high bio-sorption yield obtained by bacteria, the heavy metal bioremediation process requires microorganisms to be attached to a solid surface. Surface fixation and cell entrapment are the two methods of immobilization. Distinctive lattices were tried for cell immobilization (Beolchini et al., 2003; Xiangliang et al., 2005). Bolster frameworks appropriate for biomass immobilization incorporate alginate, polyacrylamide, polyvinyl liquor, polysulfone, silica gel, cellulose, and glutaraldehyde (Wang 2002; Vijayaraghavan and Yun 2008a, b). The polymeric grid decides the mechanical quality and synthetic protection of the last bio-sorbent molecule to be used for progressive sorption–desorption cycles, so it is imperative to pick the right immobilization lattice. Akar et al. (2009) measured the bio-sorption of 100 mgL⁻¹ of nickel at pH 6.5 to be 33.83 and 7.50 mgg⁻¹ for silica gel and Proteus vulgaris, respectively, whereas the immobilized bio-sorbent had a bio-sorption capacity of 45.48 mgg⁻¹ under the same conditions. Maximum bio-
sorption obtained using immobilized biomass provides promise for immobilized cells in a column reactor for the remediation of heavy metals. At pH 5.0, the Cd\(^{2+}\) biosorption capacity of *E. coli* biomass-free PVA beads was 1.30 mg/g, which was significantly lower than the adsorption capacity of PVA-immobilized cells, displaying a capacity of 2.18 and 4.41 mg/g for biomass loading of 8.42 and 19.5 wt %, respectively (Kao et al. 2009). Although cell entrapment imparts mechanical strength and resistance to chemical and microbial degradation upon the bio-sorbent, the costs of immobilizing agent cannot be ignored. Free cells are not suitable for use in a column, due to their low density and size they tend to plug the bed, resulting in marked declines in pressure. For industrial applications of bio-sorption, it is important to utilize an appropriate immobilization technique to prepare commercial biosorbents which retain the ability of microbial biomass to adsorb metal(s) during the continuous treatment process. The immobilization of biomass in solid structures would create a biosorbent material with the right size, mechanical strength, rigidity, and porosity necessary for use in practical processes. The immobilized materials can be used in a manner similar to ion-exchange resins and activated carbon such as adsorption–desorption cycles (i.e., recovery of the adsorbed metal, reactivated and reuse of the biomass) (Veglio and Beolchini 1997). In different matrices, tested surface fixation was chosen as the immobilization methodology instead of cell entrapment. Cell immobilization has successfully been achieved mostly in calcium alginate beads, but this matrix also has a high affinity for heavy metals. Metal retention kinetics studies with calcium alginate confirmed that almost 100% of the metal assayed was retained by the beads (Vullo et al. 2003) and that it is pointless to try to improve heavy metal retention by bacterial cell entrapment in calcium alginate beads (Arica et al. 2001; Davis et al. 2003; Vullo et al. 2003; Arica et al. 2004). Although calcium alginate is useful for entrapping cells in its gel structure, its advantage resides mostly in the re-utilization of the entrapped cells. However, the high heavy metal affinity of alginate makes it unusable for the development of continuous industrial processes, as the recovery of the alginic acid would increase the final costs of effluent treatment.
Materials and methods

1- Sampling

U-resistant bacterial strain were separated from the rock ore using nutrient agar (NA) medium and were prepared using peptic Digest of animal tissue (5 g/L), beef extract(3 g/L), NaCl (5g/L) and agar 15g/L.

2- Isolation and identification of heavy metal-resistant bacteria from rock Ore.

The isolated metal-resistant bacteria were amended with different conc. Of U metal. Pour plate was performed in NA medium and was brooded at 37°C for 24 h.

3-Determination of heavy metal-resistant bacterial isolates by plate diffusion method

Heavy metal resistant bacteria were determined by plate diffusion method (Hassen, et al. 1998). U solutions were prepared in different concentrations, say 10, 20, 50, 100, 250, 500 and 1000 ppm. Each plate was spread with overnight societies of proper living beings. To each of the plate 100 µl of appropriate U metal salt solutions were added in each wells of 10 mm in diameter and 4 mm in depth. NA plates were incubated at 37°C for 24 h. After incubation, the zone of inhibition was measured. A zone size less than 1 mm scored as resistance strain.

4. Optimization of bacterial growth conditions

Studying factor affecting bacterial growth Like pH, Temperature, The cultures were incubated at 37 °C for 24,48 h and By Detecting O.D. The development was checked using a spectrophotometer (at 600 nm) 120 Min.

5- Optimization for heavy metal uptake

(Gourdon, et al. 1990), (Gong, et al. 2005) (Kiran, et al. 2005) Reported that Temperature, pH, biomass, heavy metal concentrations are factors which influences
the bio-sorption procedure. Especially, pH biomass concentration and U metal concentration on bio-sorption tests were explored by advancement process. The bacterial isolates were inoculated into a series of test tubes containing 5 ml of nutrient broth. The pH was varied from five to nine (5, 6, 7, 8 and 9) by adjusting the medium amended with U. The biomass concentration was varied from 1 to 5% (1, 2, 3, 4 and 5) in the medium containing 25 mg/L of U. The heavy metal concentration was varied from 20 mg/L to 100 mg/L (20, 40, 60, 80 and 100 mg/L).

6- Heavy metal adsorption by the immobilized bacterial cells

the immobilized bacterial cells were set up as dabs agreeing o the strategy of Leung, et al. (2000) and were maintained in the conical flask containing 50 ml of samples for incubation, after which the specimens were pulled back for substantial metal examination by utilizing Titration method.

7. Cultivation of E. coli.

Cultivation of E. coli. Was done in 250 mL cone shaped flasks with 100 mL culture medium on a rotary shaker at 200 rpm at a constant temperature of 37°C. The way of culture medium contained The pH of the medium was adjusted to (6).

8. Determination of Uranium:

The uranium content of the sample and prepared standard and treated solution were determined according to the method Described by (Davies &gray, 1964).

9. Application of the FTIR spectra of U loaded and unloaded free and immobilized cells.

10. Preparation of immobilized bacterial biomass beads/alginate–chitosan microcapsule was composed of E. coli sodium alginate, chitosan and calcium chloride. Therefore, under sterile conditions, the bacteria were mixed with sodium alginate solution, and then the mixed solution was dropped into calcium chloride solution for immobilization using a microcapsule preparation instrument. The E. coli Capsule-loaded calcium alginate gel beads were obtained after immobilization, and the loaded calcium alginate gel beads were mixed with chitosan solution to obtain E. coli Capsule/alginate–chitosan microcapsules. The microcapsule system had good mechanical strength, flexibility and biocompatibility between the E. coli Capsule and
the microcapsule. In addition, the internal three-dimensional network structure of the microcapsule provided a sufficient space for the E. coli Capsule growth and good encapsulating stability.

Results & Discussion

1. Characterization Uranium-resistant bacterial isolates
1.1. Screening of Uranium-resistant bacterial isolates growth in presence of Different Uranium conc. Isolated from Aboshid Uranyle rock sample. From 10 bacterial isolate 6 stable isolate where studied to elect the most potent bacterial isolate to be utilized as bacterial capsule in our study.

| Uranium conc. | Isolate no. | 100ppm | 200ppm | 300ppm | 600ppm | 1000ppm |
|---------------|-------------|--------|--------|--------|--------|---------|
|               | S6          | ++     | +      | +      | +      | +       |
|               | S4          | +      | +      | +      | +      | +       |
|               | S5          | +      | +      | +      | +      | +       |
|               | S7          | +      | +      | +      | -      | -       |
|               | S8          | +     | -      | -      | -      | -       |
Table -1.a Test for Screening of uranium resistant isolates

Table 1 represents that we was have 6-10 Isolates tested for incubation with different conc. Of Uranium and investigate strong of growth against U conc. We found that the most potent isolate S6, S5. Which it's Growth continue with stability up to 1000 ppm Uranium conc. This will continue with us

Table. 1. b. Growth of bacteria in different concentrations of Waste.

| WASTE CONC. | 10% | 20% | CONC. soln. |
|-------------|-----|-----|-------------|
| S6          | ++  | +   | +           |
| S5          | +   | +   | +           |
| S7          | -+  | -+  | -           |
| S8          | -+  | -+  | -           |
| S3          | +   | +   | +           |

Previous Table indicates that also the most potent isolates were S6, S5.

| U conc. | S6 | S4 | S3 |
|---------|----|----|----|
| 10      | 2  | 2  | 2  |
| 20      | 2  | 2  | 2  |
| 40      | 2  | 2  | 2  |
| 60      | 2  | 4  | 2  |
| 80      | 2  | 4  | 2  |
| 100     | 2  | 4  | 2  |
| 200     | 2  | 5  | 5  |
| 300     | 2  | 5  | 5  |
| 600     | 2  | 6  | 8  |
| 1000    | 2  | 6  | 8  |

Table .1.C. Growth of bacteria in different concentrations of Uranium by plate diffusion method

This Table confirm that what we found in Previous Tables a,b that inhibition zone indicate metal resistant bacterial isolates .
1.2. **Optimum condition for Growth bacterial isolates:**

Optimizing a growth medium is very important to study the performance of microbes in bio-sorption Processes.

| pH | O.D.(24hr) |
|----|------------|
| 4  | 0.313      |
| 5  | 0.409      |
| 6  | 0.469      |
| 7  | 0.354      |
| 8  | 0.213      |

**Table: 2-a Cellular Growth of Bacteria (Growth Curve of bacteria at different PH).**

Hypothesis, that 6.0 is the optimal pH for *E. coli* growth was accepted. The radii of the restraint zones around the test plates were little contrasted with those of the anti-toxin test. This could be due to the concentration of the varying pH substances being too low to show the full affect of the pH. The results were not considered anomalous, however, because they do show a direct correlation between the pH and inhibition of the *E. coli*. To further improve this experiment, higher concentrations of each of the varying pH solutions would be used so that the effect that the pH has on the *E. coli* is much clearer. In addition, more varying pH solutions would be tested. The difference between a pH of 2.4 and 7.0 is quite large, as is the difference between pH 7.0 and 11.6. The maximum tolerable pH of *E. coli* may not have been 7.0, but pH 11.6 was the next tested pH after 7.0, and 11.6 was too basic for the *E. coli* to survive. So, all that can be concluded about the maximum pH is that it is between 6.0 and 11.6. Further pH values would need to be tested in order to obtain a more accurate estimate of the maximum pH for *E. coli* growth. Similarly, the minimum pH for *E. coli* growth may be somewhere between
pH 0.0 and 2.4, however further pH values would need to be tested to find the actual minimum pH for *E. coli* growth. Furthermore, pH 2.4 may have been an anomalous result. However, because there were no other tested pH values between 2.4 and 7.0, it is not certain that the pH of 2.4 is the true minimum pH that *E. coli* can tolerate.

Table: 2-bCellular Growth of Bacteria (Growth Curve of bacteria at Different pH)

| Time   | PH. 4  | PH. 5  | PH. 6  | PH. 7  |
|--------|--------|--------|--------|--------|
| 0 time | 0.0411 | 0.0511 | 0.0551 | 0.0501 |
| 20 min | 0.029012 | 0.03412 | 0.03912 | 0.04012 |
| 40 min | 0.0342 | 0.0542 | 0.0642 | 0.0542 |
| 60 min | 0.0442 | 0.0642 | 0.0542 | 0.0642 |
| 80 min | 0.0605 | 0.0505 | 0.0705 | 0.0705 |
| 100 min | 0.0459 | 0.0459 | 0.0459 | 0.0459 |
Table: 2-c Cellular Growth of Bacteria (Growth Curve of bacteria at Different Temperatures)

| Time     | 10-20 | 30-50 | 50-60 |
|----------|-------|-------|-------|
| 0 min.   | 0.0411| 0.0511| 0.0551|
| 20 min   | 0.029012| 0.03412| 0.03912|
| 40 min   | 0.0442| 0.0642| 0.0542|
| 60 min   | 0.0542| 0.0562| 0.0532|
| 80 min   | 0.0705| 0.0755| 0.065|
| 100 min  | 0.0449| 0.0459| 0.0439|

Temperature, like pH, affects the activity of enzymes. The results show a direct correlation between the temperature and E. coli growth as shown in Table 4 and Figure 7. The hypothesis, that the optimal temperature for E. coli growth is 37 °C, was conformed. The most and largest colonies were present on the temperature test plate that was placed in the incubator set at 37 °C. As with the case of varying pH, when the temperature is changed to a temperature outside the tolerable range of an enzyme; the enzyme becomes denatured and cannot function. This is what was happening in the E. coli bacteria that were on test plates in the refrigerator and. The E. coli on the test plates in 45 °C and 50 °C were able to survive, but as the temperature increased from 37 °C
C to 50 °C, the size and number of colonies decreased. The hypothesis, that the optimal temperature for *E. coli* growth is 37 °C was accepted.

**Table: 2-d Cellular Growth of Bacteria (Growth Curve of bacteria at Different Initial Biomass)**

| Biomass conc. | O.D. at 500 | 24 hr. |
|---------------|-------------|-------|
| 0.1ml         | 0.02661     | 0.152 |
| 0.5ml         | 0.0549      | 0.688 |
| 1ml           | 0.07311     | 0.698 |

In our study bacteria capsule as we mentioned have standard dosage as 3% of the volume but we study the bacteria inoculum to get optimum turbidity, which reflect optimum growth to obtain high viability with available active sites.

**Table: 2-ECellular Growth of Bacteria (Growth Curve of bacteria at Different Initial Biomass)**

| Time     | Biomass conc. 0.1ml(O.D.600nm) | Biomass conc. 0.5ml(O.D.600nm) | Biomass conc. 1ml O.D.600nm |
|----------|---------------------------------|---------------------------------|-----------------------------|
| 0 time   | 0.02661                         | 0.0149                          | 0.07311                     |
| 20 min   | 0.029012                        | 0.03412                         | 0.03912                     |
| 40 min   | 0.0342                          | 0.0442                          | 0.0540                      |
| 60 min   | 0.0542                          | 0.0502                          | 0.0641                      |
| 80 min   | 0.0605                          | 0.0725                          | 0.0705                      |
| 100 min  | 0.0409                          | 0.0439                          | 0.0459                      |

Regarding to Table 2 the dosage of a bacterial biomass strongly influences degree of bio-sorption. An expansion in biomass fixation by and large builds the measure of solute biosorbed, because of the expanded surface zone of the cell wall, which thusly expands the quantity of restricting destinations while the amount of metal solute per unit weight of biomass diminishes with an expanding biosorbent dose, which might
be because of the perplexing connection of a few elements. An essential factor at high sorbent doses is that the accessible solute is deficient to totally cover the accessible replaceable destinations on the biosorbent, as a rule bringing about low solute take-up. The obstruction between restricting locales because of expanded biosorbent doses can't be overruled, as this will bring about low particular take-up.

**Optimum Condition for bio sorption Process**

Initial solute concentration appears to have an impact on bio-sorption, with a higher concentration resulting in a high solute uptake

| U CONC. | Qe (sorption %) | O.D. |
|---------|-----------------|------|
| 20 ppm  | 100%            | 0.05 |
| 50 ppm  | 100%            | 0.0490 |
| 100 ppm | 100%            | 0.0459 |
| 250 ppm | 100%            | 0.0438 |
| 500 ppm | 100%            | 0.0468 |
| 1000 ppm| 93 %            | 0.0389 |

Bueno et al. 2008; Uzel and Ozdemir 2009 reported that. This occurs because at lower initial solute concentrations, the ratio of the initial moles of solute to the available surface area is low; subsequently, the fractional sorption becomes independent of the initial concentration. However, at higher concentrations, the sites available for sorption become fewer compared with the moles of solute present and, hence, the removal of solute is strongly dependent upon initial solute concentration.

**2.1. Effect of pH bio sorption Process**

As a rule, expanding pH builds the negative charge on the cell surface until the point that all applicable practical gatherings are deprotonated, which favors electrochemical fascination and adsorption of inscriptions. Moreover, the expansion in metal take-up with an expansion in pH might be the aftereffect of more proficient rivalry of captions with H⁺ for restricting locales on microscopic organisms. Where it influences the arrangement science of metal particles and the surface practical gatherings of the bacterial cell divider it additionally influences the solvency of the metal particles in
the arrangement, where H\(^+\) particles supplant a portion of the positive particles from the biomass (Long et al. 2014). Detailed that the pH is an essential parameter, which. The bio-sorption limit of metal subtilises increments with increment in pH esteem, and this might be because of the more negative restricting destinations uncovered on Bacterial surface (Aksu and Gulen 2002). Announced that at low pH esteem are blocked and connected with hydrogen particles that prevent the entrance of metal cations because of ghastly powers to the surface useful gatherings. While (Aryal et al. 2010; Zia-gova et al. 2007) Stated that, biosorption proficiency of metal anions increments with diminish in pH esteem because of the expansion in emphatically charged Bacterial surface gatherings, while at higher pH, the shocking powers between metal anions and adversely charged biomass surface reduce the metal take-up limit (Aryal and Liakoupolou Kyriakides 2013b) Reported that bio-sorption of metal increased with increasing pH from 1.0 to 5.0 and decreased upon further increase to 7.0. The decrease in the sorption efficiency at pH higher than 5.0 may be due to the precipitation of \(\text{metal} \times \text{hydroxide}\), now in our study as previous mentioned above Table 3 that pH 5 and 6 are the most favorable for bio-sorption in that waste where higher pH availability of precipitation occur prevent accurate detection of uranium bio-sorption But in our case of waste of high content 100, 250, 500 and 1000 ppm First result waste of 100 ppm content after 24 hr incubation time U reside is Nil at that pH(6)

**Table: 3 a. Cellular Growth of bacterial isolates (O.D. at 500) in response to various biomass concentrations and 100 ppm. Temperature 37°C, incubation time: 24h.**

| Biomass conc. | 0 time  | 24 hr  | Qe %  |
|--------------|---------|--------|-------|
| .1ml         | 0.02661 | 0.598  | 100%  |
| 0.5ml        | 0.0149  | 0.152  | 100%  |
| 1ml          | 0.07311 | 0.698  | 100%  |

**2.2. Effect of biomass concentration on bio-sorption process.**

Bio-sorption of Uranium substantial metals wards on biomass fixation utilized as the sorption medium. An expansion in biomass fixation as a rule brings about increment
of bio-sorption productivity, likely because of the increment in the quantity of restricting destinations. It was watched that the sorption productivity expanded with increment in biomass fixation, however biomass focuses over 1.0 and 2.0 g/L had bring down effect in sorption proficiency of As (III) and As (V), individually (Aryal et al. 2010) This lower augment in rate expulsion above ideal biomass fixation might be credited to the obstruction between Capacity Gathering. A few investigations have brought up that take-up limit of Uranium& overwhelming metals diminishes with expanding the biomass focus because of solid impediments of ionic species portability in the bio-sorption medium, abandoning some coupling locales for metal particles unsaturated. (Aryal et al. 2012; Tangaromsuk et al. 2002). In Incharoensakdi et al study expressed that The particular take-up of Zn (II) decreased, when Microbial biomass fixation surpassed as 0.2 g/L (Incharoensakdi and Kitjaharn 2002). Ziagova et al. (2007) detailed the huge increment in the Cr (VI) take-up productivity, when the biomass convergence of Staphylococcus xylosus expanded metal anion did not change fundamentally over 1.0 g/L of Bacterial biomass from 1.0 to 8.0 g/L, though expulsion effectiveness if there should be an occurrence of our investigation we utilize immobilized mass by infuse 3 ml of bacterial biomass in 100 ml calcium alginate water.

Table: 3b. Sorption % of bacteria at Different Temperatures.

| Temperature Range | O.D. at 500 | 24 hr. | Qe %  |
|-------------------|-------------|--------|-------|
| 10-20             | 0.02461     | 0. 1520. | 70%   |
| 30-50             | 0.0149      | 0. 678 | 100%  |
| 50-60             | 0.07311     | 0.398  | 60%   |

We know that optimum temperature for growth of bacteria in general ranges from 30-50 so optical densities increase consequently binding sites increase so bio-sorption capacity of bacterial isolates increase. Where it in low temperature low O.D. so low binding sites so sorption capacity decrease where in high temperature moderate O.D. but also still binding sites active and capacity moderate high comparing with low temperature. So optimum temperature for bio-sorption process rages between 30-50.
Table: 3C. Cellular growth at 100-ppm U standard

| Time  | O.D. at 500nm |
|-------|---------------|
| 0 time| 0.0511        |
| 20 min| 0.03012       |
| 40 min| 0.0542        |
| 60 min| 0.0442        |
| 80 min| 0.0505        |
| 100 min| 0.0359       |

Table: 3d. Incubation time w U20ppm O.D.at 500nm

| Incubation time w U 20 ppm | O.D.at 500nm |
|---------------------------|--------------|
| 0 time                    | 0.0411       |
| 20 min                    | 0.0301       |
| 40min                     | 0.0602       |
| 60 min                    | 0.0542       |
| 80min                     | 0.0605       |
| 100min                    | 0.0459       |

The uptake of Uranium is examined at time interval and the result are shown table 3 as can be seen in table that during the first 50 min of the experiment the concentration of Uranium adsorbed by immobilized bacterial isolate increases with increase from 50 to 120 min no change in concentrations observed in time our experiment 24 hr. were selected to confirm adsorption equilibrium.

| Incubation time | U Conc. 250 | U con.500 | Uconc.1000 |
|-----------------|-------------|-----------|------------|
| 24 hr.          | 100%        | 100%      | 98%        |
| 72 hr.          | 100%        | 100%      | 100%       |

Table: 4 for Effect of contact time on bio-sorption efficiency.
2.3. Techniques Used in Metal Bio-sorption Studies

In study of Jian-hua et al. (2007) In investigation of Jian-hua et al. (2007) he effectively associated the amount of acidic gatherings introduce on Bacillus cereus biomass, decided through potentiometric titrations, with the metal take-up limit. The idea of the coupling locales and their inclusion amid bio-sorption can be roughly assessed utilizing FTIR. So we utilize FTIR systems to decide The idea of the coupling locales where investigation of Mishra and Doble (2008) indicated that carboxyl and amino groups were responsible for the binding of chromate, Carboxyl groups are negatively charged and abundantly available, actively participate in binding of metal cations. Kang et al. (2007) observed that amine groups protonated at pH 3 and attracted negatively charged chromate ions via electrostatic interaction. Potentiometric titrations can provide information on type and number of binding sites. Kang et al. (2007) titrated Pseudomonas aeruginosa and determined the pKa values of available binding sites Loukidou et al. (2004).

2.4. Analyzed the FTIR spectra of U loaded and unloaded

we use FT-IR spectra as to confirmed availability of binding sites as shown in Table : 5a Table : 5b  for Uranium we found Amino acid(O-H) Stretching protein v(N-H) stretching , Phosphate C-O Stretching band ,P-H stretching, Protein amide I band mainly(C=O) Stretching, Protein (CH$_2$) and (CH$_3$)bending of methyle Lipid (CH$_2$) bending of methyl, Carbohydrate (c-o) of polysaccharides, Nucleic acid (other phosphate containing compound ) , $p=0$ stretching of phosphodiester, acid chlorides C-Cl stretch in S6 E.coli, and comparing with dead isolate we found the same Beside acid chlorides at position 550 cm$^{-1}$ C-Cl stretch Cayllahua et al. (2009) study who used FTIR spectra to confirm the presence of amide, carboxyl, and phosphate groups in Rhodococcus sp. Biomass
Table: 5a data of FT-IR OF un loaded E. coli. (S6) unloaded and Bacterial Isolate.

| Wave number range | Typical band | Intensity of loaded band live bacteria | Main peak(cm⁻¹) |
|-------------------|--------------|----------------------------------------|-----------------|
| 3029-3639         | Amino acid(O-H) stretching protein v(N-H) stretching | 77.9            | 1-3439.42       |
| 2344-2365         | Phosphate C-O Stretching band ,P-H stretching        | 93              | 2:2355.62       |
| 1583-1709         | Protein amide I band mainly(C=O) Streching           | 90              | 3-1638.23       |
| 1425-1477         | Protein (CH₂) and (CH₃)bending of methyle Lipid (CH₃) bending of methyl | 95              | 4-1428.99       |
| 1072-1356         | Carbohydrate (c-o) of polysaccharides, Nucleic acid (other phosphate containing compound) >p=o stretching of phosphodiester | 100             | 5-1101.15       |
| 730-550           | acid chlorides C-Cl stretch                          | 97              | 6-556.363       |

Table: 5b data of FT-IR OF unloaded E. coli. (S6) loaded Bacterial Isolate.

| Main peak(cm⁻¹) | Intensity of loaded band | Typical band | Wave number range |
|-----------------|--------------------------|--------------|-------------------|
| 1-3455.42       | 50.4                     | Amino acid(O-H) stretching protein v(N-H) stretching | 3029-3639         |
| 2-2088.62       | 99                       | Phosphate C-O Stretching band ,P-H stretching        | 2344-2365         |
| 3-1641.23       | 78                       | Protein amide I band mainly(C=O) Streching           | 1583-1709         |
| 4-1428.99       | 96                       | Protein (CH₂) and (CH₃)bending of methyle Lipid (CH₃) bending of methyl | 1425-1477         |
| 5-1109.15       | 105                      | Carbohydrate (c-o) of polysaccharides, Nucleic acid (other phosphate containing compound) >p=o stretching of phosphodiester | 1072-1356         |
| 6-593.363       | 92                       | acid chlorides C-Cl stretch                          | 730-550           |
| 7-550.577       | 92                       | acid chlorides C-Cl stretch                          | 730-550           |
A few band changes enabled the creators to anticipate the conceivable contribution of amino, carbonyl, carboxyl, and phosphate bunches in the bio-sorption of Uranium.

Table: 5c data of FT-IR OF unloaded immobilized E. coli. (S6) Bacterial Isolate.

| Main peak(cm⁻¹) | Intensity of loaded band | Typical band | Wave number range |
|----------------|--------------------------|--------------|------------------|
| 1-3436.42      | 71.14                    | Amino acid(O-H) stretching protein v(N-H) stretching | 3029-3639        |
| 2-1725.23      | 97                       | Phosphate C-O Stretching band ,P-H stretching       | 2344-2365        |
| 3-1636.3       | 92                       | Protein amide I band mainly(C=O) Streching          | 1583-1709        |
| 4-1380.99      | 86                       | Protein (CH₂) and (CH₃)bending of methyle Lipid (CH₂) bending of methyl | 1425-1477        |
| 5-1354.15      | 80                       | Carbohydrate (c-o) of polysaccharides, Nucleic acid (other phosphate containing compound ) >p=ø stretching of phosphodiester | 1072-1356        |
| 6-1038.48      | 91.90                    | Carbohydrate (c-o) of polysaccharides, Nucleic acid (other phosphate containing compound ) >p=ø stretching of phosphodiester | 1072-1356        |
| 7-924.7.363    | 99                       | acid chlorides C-Cl stretch                         | 730-550          |
| 8-879.577      | 99                       | acid chlorides C-Cl stretch                         | 730-550          |
| 9-808.992      | 98                       | acid chlorides C-Cl stretch                         | 730-550          |
| 10-597         | 88                       | acid chlorides C-Cl stretch                         | 730-550          |

All this evidence suggest that functions groups Represented in table (5a, 5b, 5c, 5d) are responsible for metal uptake in our bacterial biomass beside change in peaks
position which assigned for it's groups confirm bio-sorption of metal ions from waste due to ions charge interaction comparing table 5a, 5b with 5c, 5d we found increase in no of binding sites indicate that immobilized bacterial have high efficiency for metal up take which also change in peaks position which assigned for it's groups confirm bio-sorption of metal ions from waste due to ions charge interaction.

Table: 5 d data of FT-IR OF loaded immobilized *E. coli*. (S6)

| Bacterial Isolates | Main peak(cm⁻¹) | Intensity of loaded band | Typical band | Wave number range |
|--------------------|-----------------|--------------------------|--------------|------------------|
|                    | 1-3909.42       | 96.64                    | Amino acid(O-H) stretching protein v(N-H) stretching | 3029-3639 |
|                    | 2-3859.23       | 96                       | Phosphate C-O Stretching band ,P-H stretching | 2344-2365 |
|                    | 3-1636.3        | 92                       | Protein amide I band mainly(C=O) Streching      | 1583-1709 |
|                    | 4-1729.99       | 93                       | Protien (CH₂) and (CH₃)bending of methyl Lipid (CH₃) bending of methyl | 1425-1477 |
|                    | 5-1630.15       | 85                       | Carbohydrate (c-o) of polysaccharides, Nucleic acid (other phosphate containing compound ) >p=O stretching of phosphodiester | 1072-1356 |
|                    | 6-1429.4        | 90.90                    | Carbohydrate (c-o) of polysaccharides, Nucleic acid (other phosphate containing compound ) >p=O stretching of phosphodiester | 1072-1356 |
|                    | 7-1382.7        | 80                       | acid chlorides stretch C-Cl                      | 730-550  |
|                    | 8-1175.577      | 100                      | acid chlorides stretch C-Cl                      | 730-550  |
|                    | 9-1037.5,992    | 90                       | acid chlorides stretch C-Cl                      | 730-550  |
|                    | 10-936.27       | 103                      | acid chlorides stretch C-Cl                      | 730-550  |
|                    | 11-880.34       | 104.166                  | acid chlorides stretch C-Cl                      | 730-550  |
|                    | 12-818.634      | 101                      | acid chlorides stretch C-Cl                      | 730-550  |
|                    | 13-664.357      | 95                       | acid chlorides stretch C-Cl                      | 730-550  |
|                    | 14-562.148      | 92                       | acid chlorides stretch C-Cl                      | 730-550  |
2.5. Energy dispersive X-ray (EDX):

![Figure 1 - (EDX) to confirm Bio-sorption of U by Capsulated Cell-Free Extract Loaded Ca-Alginate Beads.](image)

The alginate beads (Fig. 1), predominantly ellipsoidal spheres, with average diameter of 3–5 mm were used in the packed bed to remediate 10-1000 ppm U (VI) in a synthetic Uranium solution. The effectiveness of different dosages of beads was considered and the optimized ratio of 1:5 (v/v) of beads to water was used in all batch
studies of isotherm kinetics. Scanning electron microscopic of these beads, Synthetic Solution (Fig.1), Control(Fig.2), showed that these were hollow from inside (having smooth inner wall In SEM/EDS analysis of the Ca-alginate beads after the experiment, void spaces of the beads were found to be filled with precipitates of heavy metals, showing that Ca-alginate beads can be successfully used as a biosorbent for the removal of Uranium and which agreed with (Sumin Park and Minhee Lee 2017) in in Substantial metal evacuation like Cu ,Cd. The picture of the control demonstrates nonappearance of any metal besides to Uranium, which as of now not present. While in Fig.1,2, 3,4, Indicated Uranium as Fig. 3,4. In the spot zone affirmed Uranium bio-sorption.
Figure 2- (EDX) to confirm Bio-sorption of U by capsulated Cell-Free Extract Loaded Ca-Alginate Beads

(Fig. 3). This established the fact that *E. coli* interacted with the metal present in the medium.

**Fig. 3** Bioremediation of 600 ppm U (VI) from synthetic solution in bottle
Fig. 4 the IR Spectrum of Live E. coli before Bio-sorption Process.
Kazy et al. (2006) utilized X-beam diffraction (XRD) examination and affirmed the inclusion of cell carboxyl and phosphate bunches in the authoritative of lanthanum by Pseudomonas biomass. SEM micrographs have supported analysts in dissecting cell surface morphology prior and then afterward bio-sorption. Tunali et al. (2006) envisioned the surface of metal-stacked Bacillus sp. while in our investigation we utilized SEM helped us in affirming U bio-sorption.

**IR Examination of Cell wall of **E. coli**:

Cell wall of **E. coli** used in bio-sorption showing Function group in cell wall and comparing it with autoclaved forms exploring difference in intensity of function group of cell wall which involved in bio-sorption this difference in intensity Reflected in bio-sorption capacity as showed in **figure 4** function group and its intensity in case of free cell.

**2.7. Comparative study showing different forms of Bacterial Biomass.**

**Table: 6. Treatment for different form of sample and applications.**

| Bio-sorption%                  | Immobilized Bacteria | Autoclaved bacteria | Free bacterial cell |
|-------------------------------|----------------------|---------------------|---------------------|
| Standard Uranium 100ppm       | 100%                 | 80%                 | 65%                 |
| Waste water (low conc.)       | 100%                 | 100%                | 70%                 |
| Phosphoric acid(40PPM)        | 90%                  | 50%                 | 60%                 |

**4.3. Relation between Immobilization of Cells and efficiency of bio-sorption:**

Immobilized individual enzymes can be successfully used for single-step reactions. They are, however, not suitable for multi-enzyme reactions and for the reactions requiring cofactors. The whole cells or cellular organelles can be immobilized to serve as multi-enzyme systems. In addition, immobilized cells rather than enzymes
are sometimes preferred even for single reactions, due to cost factor in isolating enzymes. For the enzymes, which depend on the special arrangement of the membrane, cell immobilization is preferred. Immobilized cells have been traditionally used for the treatment of sewage. The techniques employed for immobilization of cells are almost the same as that used for immobilization of enzymes with appropriate modifications. Entrapment and surface attachment techniques are commonly used. Gels, and to some extent membranes, are employed.

**Immobilized live Cells:**
The viability of the cells can be preserved by mild immobilization. Such immobilized cells are particularly useful for fermentations. Sometimes mammalian cell cultures are made to function as immobilized viable cells.

**Immobilized dead Cells:**
In many instances, immobilized non-viable cells are preferred over the enzymes or even the viable cells. This is mainly because of the costly isolation and purification processes. The best example is the immobilization of cells containing glucose isomerase for the industrial production of high fructose syrup.

Data showed in table 6 reveal that bacterial isolate can treat different forms of wastes and show that immobilized bacterial isolate gain high protection enable it to treat waste with high acidity like Phosphoric acid while free cell of the same bacterial isolate can not treat this type of waste so can't complete bio-sorption process with the same efficiency so we found that immobilized bacterial isolates with bio-sorption efficiency reach 100% and 90 % respectively in case of standard Uranium, and Phosphoric acid ,where immobilization Provide high number of active sites as mentioned above in table 5 c , 5d entrapment imparts mechanical strength and resistance to chemical and microbial degradation upon the bio-sorbent, the costs of immobilizing agent cannot be ignored. Free cells are not suitable for use in a column; due to their low density and size they tend to plug the bed, resulting in marked declines in pressure. For industrial applications, as in our study we made alginate beads like (capsule) of bio-sorption, it is important to utilize an appropriate immobilization technique to prepare commercial bio-sorbents, which retain the ability of microbial biomass to adsorb metal(s) during the continuous treatment process. The
immobilization of biomass in solid structures would create a bio-sorbent material with the right size, mechanical strength, rigidity, and porosity necessary for use in practical processes. The immobilized materials can be used in a manner similar to ion-exchange resins and activated carbon such as adsorption–desorption cycles.

2.8. Effect of desorption study

Table: 7. Desorption ratio of uranium ions adsorbed by Coli. /alginate

| No of cycle | 1st | 2nd | 3rd | 4th | 5th |
|-------------|-----|-----|-----|-----|-----|
| 100ppm      | 100%| 100%| 100%| 100%| 100%|
| 250ppm waste| 100%| 100%| 100%| 100%| 100%|
| ppm 500     | 100%| 100%| 100%| 100%| 100%|
| 1000ppm     | 95% | 100%| 100%| 100%| 95% |

The proposed procedure could be practical in performing remediation on location particularly for the emanating of enterprises. The real preferred standpoint of operation could be typical natural conditions under which the dots work productively. Subsequently, the curiosity of this proposed Work was because of the work of strain \( E. \ coli \) sp. whose healing limit in all structures (whole cell, crude Eluents % recovery Time taken (hr.) Desorption ratio Beads’ condition (post-desorption)

| Eluents 1M HN\(_0\)\(_3\),Cacl\(_2\) | % Recovery | Time(hr.) | Desorption Ratio | Beads(post Desorption) |
|--------------------------------------|------------|-----------|-----------------|------------------------|
| 10                                   | 100%       | 24        | 100%            | Intact Bead            |
| 50                                   | 100%       | 24        | 100%            | Intact Bead            |
| 100                                  | 100%       | 24        | 100%            | Intact Bead            |
| 250                                  | 100%       | 24        | 100%            | Intact Bead            |
| 500                                  | 100%       | 24        | 100%            | Intact Bead            |
| 1000                                 | 98%        | 24        | 98              | Intact Bead            |

Table 8 Desorption study of used CFE-loaded Ca-alginate beads

Along these lines, it is important to assess the appropriate elutant volume, which can be performed utilizing explores different avenues regarding distinctive strong to
liquid proportions. The strong to-fluid proportion is characterized as the mass of solute-loaded bio-sorbent to the volume of elutant. Davis et al. (2000) watched that strong to-fluid proportion influenced copper elution effectiveness of Cacl$_2$ arrangements, while it was about free on account of 0.1 M HCl. The reason for desorption is to unbind a contaminant from a bio-sorbent, so both the recouped solute and bio-sorbent can be reused. After desorption, the bio-sorbent ought to be near its unique shape, both morphologically and practically. Additionally, amid the desorption procedure; expulsion of all bound sorbate from bio-sorbent ought to be guaranteed. On the off chance that this does not happen, a decreased take-up ought not out of the ordinary in the following cycle. Puranik and Paknikar (1999) recovered and reused a polysulfone-immobilized Citrobacter strain more than three cycles for the bio-sorption of lead, cadmium, and Zinc, utilizing 0.1 M HCl and 0.1 M EDTA as elutants, yet just with constrained achievement, and accentuated the requirement for additionally screening work. Beolchini et al. (2003) immobilized Sphaerotilus natans into a polysulfone framework for the bio-sorption of copper, and with the guide of 0.05 M Cacl$_2$ recovered and reused the dabs more than ten cycles with tasteful outcomes.

Table: 9. Decrease in % bioremediation due to desorption from beads after each sorption cycle

| No of cycle | 1$^{st}$ | 2nd | 3rd | 4th | 5th |
|-------------|--------|-----|-----|-----|-----|
| 100ppm      | 100%   | 100%| 100%| 100%| 100%|
| 250 ppm waste | 100%   | 100%| 100%|     |     |
| ppm 500     | 100%   | 100%| 100%| 100%| 100%|
| 1000 ppm    | 97%    | 95% | 95% | 95% | 95% |

Desorption think about: Out of the different eluents utilized, 0.5 M HNO$_3$ ended up being the most productive. The desorption apportion is given by: Desorption ratio= [Amount of U (VI) particles in desorption medium/Same adsorbed in beads] $\times$100 (Table 9). It was discovered that a cluster of dabs dose could perform bioremediation up to 5 cycles all together (15 days). What's more, ready to finish
more than additional 5 cycle Past 5 cycles, the dots not swelled up and not contorted. This may be because of the corrosive increment it's strength and unbending nature denaturation of alginate-protein connection by desorption treatment and the came about medicinal rate likewise diminished sans cell separate and epitomized in bioploymeric hydrogel dabs) has been high (93–99.7 %) and utilization of an extraordinary reused connect stream model to semi-cluster method of adsorption. Kuyucak and Volesky (1989) analyzed a few synthetic operators to desorb Co$^{2+}$ from cobalt-loaded Ascophyllum nodosum, and distinguished CaCl$_2$ within the sight of HCl as a reasonable elutant. The execution of an elutant additionally unequivocally relies upon the sort of system in charge of bio-sorption. For example, electrostatic fascination was observed to be the essential system in charge of bio-sorption of contrarily charged color anions to a decidedly charged cell surface (O’Mahony et al. 2002). In this way, it is coherent to influence the cell to surface negative utilizing antacid answers for repulse the adversely charged responsive colors (Won and Yun 2008). Elution is likewise affected by the volume of elutant, which ought to be as low as for all intents and purposes conceivable to acquire the most extreme solute focus in the littles conceivable volume (Volesky 2001). In the meantime, the volume of the arrangement ought to be adequate to give most extreme solvency to the desorbed solute. Likewise, one needs to understand that the desorbed sorbate remains in arrangement, another harmony is set up amongst that, and the one (staying) still settled on the biosorbent. This prompts the idea of a "desorption isotherm" where the harmony is firmly moved toward the sorbate disintegrated in the arrangement (Yang and Volesky 1996).

Conclusion:

We endeavored to expel U from U wastewater tested at Nuclear Materials Authority utilizing microscopic organisms showing a noteworthy capacity to amass U. Suitable cells of E. coli were suspended in 100 mL of answer-immobilized microbes (pH 6.0) of wastewater containing U for 1 h at 25°C. E. coli confined from Egypt U Metal expelled 90% and 78% U, separately (Table 15, 16), when arrangement pH was balanced at first to 6.0. Arrangement pH step by step diminished, with E. coli cells being all the more unfavorably influenced by pH. Be that as it may, strains quantitatively expelled U when the pH was kept up at 6.0. These species would thus be able to expel U from U refining wastewater with a high productivity.
were likewise made to expel U from U refining wastewater utilizing immobilized microorganisms having a high capacity to adsorb U. Uranium refining wastewater (100 mL, pH 6.0) supplemented with 2.1 mM of U were adsorbed on a segment (bed volume 2 mL) of immobilized bacterial cells. Immobilized bacterial cells disconnected from U mines in the Egypt can also remove U from the U refining wastewater with high efficiency U Sorption at Different condition incubation time 100 min.

DECLARATIONS

- ETHICAL APPROVAL AND CONSENT TO PARTICIPATE
  Manuscripts Doesn’t reporting studies involving human participants,

- CONSENT FOR PUBLICATION
  Not Applicable

- AVAILABILITY OF DATA AND MATERIAL
  “Please contact author for data requests.”

- COMPETING INTERESTS
  “The authors declare that they have no competing interests.”

- FUNDING
  The authors have No Funds.

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