



enterprises, hardware and space businesses. This innovation manages little structures and little estimated materials of measurements in the scope of couple of nanometers to fewer than 100 nanometers. NPs show exceptional and extensively changed substance, physical and organic properties contrasted with main part of the same synthetic organization, in view of their high surface-to-volume extent. These particles moreover have various applications in different fields, for instance, healing imaging, nanocomposites, channels, drug transport and hyperthermia of tumors. A key domain of examination in nanoscience discussed the combination of nanometer range particles of diverse shapes, monodispersity and sizes [1-33].

### **Morphology of Nanoparticles**

The morphological uniqueness of nanoparticles is: flatness and aspect ratio.

- 1- High aspect ratio NPs.
- 2- Small aspect ratio NPs.

### **Nanomaterials types**

NMs might be natural, inorganic and composite in nature. Natural nanomaterials incorporate man-sized proteins, nucleic acids and carbon nanotubes and so forth. On the other side inorganic nanomaterials incorporate metallic NPs, mixes and mass nano-organized materials.

These metallic nanomaterials are Ag, Au, Pd and Pt [1,2]. It likewise incorporates bimetallic nanoparticles like Ag-Pt, Ag-Au, Pd-Au, Ti [TiO<sub>2</sub>], oxides of Fe [Fe<sub>2</sub>O<sub>3</sub>-] magnetite, Fe<sub>3</sub>O<sub>4</sub>-magnetite [3], Zn [ZnO, Si [SiO<sub>2</sub>] and in addition sulfides of various metals [CdS [4], FeS [3], ZnS [5,6]. Different nanomaterials incorporate mass nano-organized metals, precious stones and powders of Te [7], Ti [8], Se, Al [9] and move metals like Ni, Co, Cu, Cr, Zr [10] and Pb. Then again, nano-composite materials include quantum Dots, carbon nanotubes [11], nano-shells, nano-bars, nano-wires, nano-gels and nano-emulsions and so on. Nanoparticles can be sorted by measurements for instance zero-dimensional iota groups, one-dimensional regulated multilayered, two-dimensional NPs, three-dimensional, ultra-fine grained over layers, [12].

### **NMs properties**

The nanoscale is unique in light of the fact that no strong can be readied littler than it. It is likewise remarkable in light of the fact that a large portion of the components of the physico-synthetic and natural world capacity on estimation scales going from 0.1 nm to 100 nm. By and large, the lessening in size of the particles builds their surface to volume proportion which in this way expands their reactivity.

At nano-scale, particles of various components show amazing physical, compound and natural properties. In contrast with

traditional coarse-grained partners, NPs show lower softening point, electrical resistivity, particular warmth, diffusivity, pliancy and mechanical quality alongside changes in their electromagnetic and synergist properties. Researchers are attempting to decipher these properties connected with NPs in different advancements.

### Synthesis of nanomaterials

The nanomaterials synthesis involves chemical, physical and biological methods. Biological procedures are still in the stage of development [13] FIG 1.

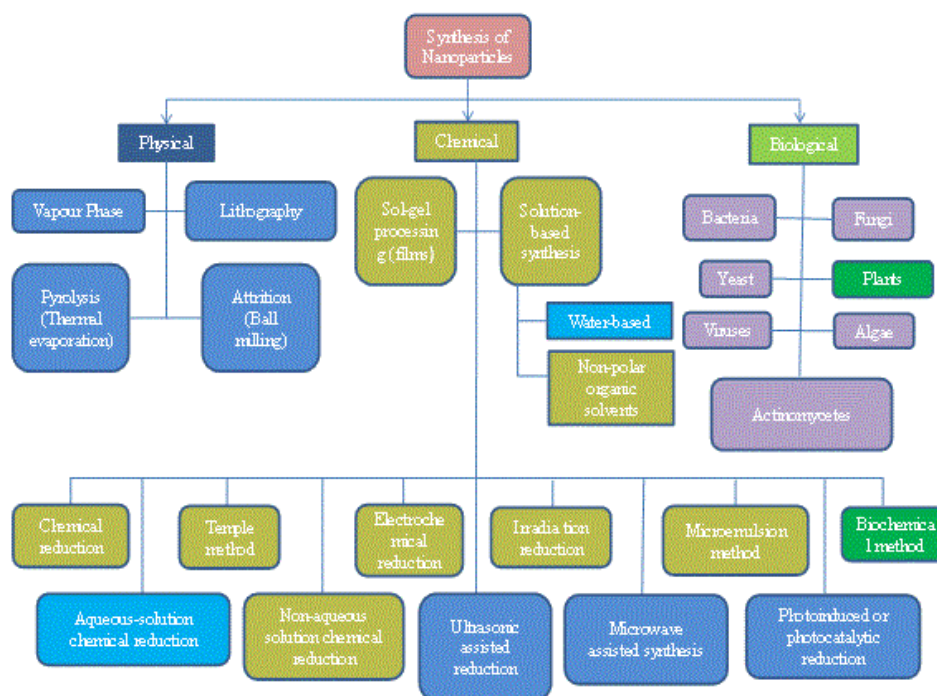


FIG.1. Different methods for the synthesis of NPs.

### Physical and chemical synthesis of nanomaterials

Physical methods for blending nanomaterials include crushing, physical vapor deposition (PVD), ball processing, lithography and pyrolysis. Among these, pounding and pyrolysis have been utilized most regularly. While, there should arise an occurrence of substance systems including electrolysis, sputtering, sol gel synthesis, CVD (chemical vapor deposition), sputtering and inactive gas buildup have been more mainstream. These types of techniques are advantageous however they are dangerous as well furthermore exorbitant.

### Biological synthesis of nanomaterials

Biological sources include plants and microbes like yeast, fungi, bacteria and actinomycetes. It has been found that biological sources act as reducing metal ions from the soil. This process is quite safe as comparison to other physical and chemical methods. It is a lesser amount of energy concentrated loom but the nanomaterials produced by these methods have found very reactive. This “enzymatic” proposal is extra workable by the reality that most of the microorganisms raise at climate situation of temperature and pH (TABLE 1).

TABLE 1. Advantaged and disadvantages of different techniques.

| Technique  | Advantages   | Disadvantages   | Chemicals/<br>Microorganims used  | References                      |
|------------|--|---|---|---------------------------------|
| Physical   |  | Firm join, baffled in data and effectiveness note being of NPs Deliver of the crystal expansion, synthesis of the particles   |   | Mandal et al. [9]               |
| Chemical   | High monodispersity (5–15%)                          | Costly feeble-minded in figure and sortie consequently claim of mortal chemicals Note env ironmental easy to deal with resolution of NP Manage of the crystal development, synthesis of the particles Low yield | Reducing agents, such as methoxypolyethylene glycol, sodium borohydride, potassiumbitartarate, hydrazine Stabilizing agents, such as polyvinyl pyrrolidone, sodium dodecylbenzyl sulfate. | Sastry et al. Mandal et al. [9] |
| Biological | Low monodispersity (~40–50%), environmental friendly |   | Yeast, bacteria, fungi, algae, plants, viruses and actinomycetes.   | Sastry et al.                   |

Synthesis of nanoparticles can occur either intracellularly or extracellularly.

Intracellular synthesis of nanoparticles requires extra steps, for example, ultrasound treatment or responses with appropriate cleansers to discharge the combined nanoparticles [14-16].

In the meantime, extracellular biosynthesis is inferior and it requires more straightforward downstream handling. As a result of this, numerous studies were focused on extracellular techniques for the blend of nanoparticles.

TABLE 2. Microorganisms involve in the synthesis of different NPs.

| Biological Entity             | Microorganisms   | Types of NPs synthesized                              | References  |
|-------------------------------|--|---|-------------|
| Bacteria 1 nm–200 nm diameter | <i>Bacillus licheniformis</i> ; <i>Bacillus subtilis</i> ; <i>Bacillus stearothermophilus</i> ; <i>Clostridium thermoaceticum</i> ; <i>Desulfobacteriaceae</i> ; <i>E. coli</i> ; <i>Klebsiella aerogenes</i> ; <i>Klebsiella pneumonia</i> ; <i>Lactobacillus increase</i> ; <i>Rhodopseudomonas capsulata</i> ; <i>Magnetospirillum magnetotacticum</i> ; <i>Desulfovibrio desulfuricans</i> ; <i>Pseudomonas aeruginosa</i> ; <i>Pseudomonas stutzeri</i> AG259; <i>Rhodopseudomonas palustris</i> ; <i>Thermoanaerobacter ethanolicus</i> (TOR-39) | Ag, Au, CdS, pd, Fe <sub>3</sub> O <sub>4</sub> , ZnS | [4,9,15,16] |

#### Why choose bacteria?

Nanoparticles are constantly being studied and developed. Three methods are used to synthesize the nanomaterials. Physical methods include lithography, pyrolysis, vapour pressure etc. but these methods are quite expensive. It is important that physical and chemical methods are low yield, energy intensive, difficult to scale up, often produce high levels of hazardous wastes and may require the use of costly precursors.

Besides, a chemical method includes Irradiation reduction, micro-emulsion method and electrochemical reduction but chemical methods are quite hazardous this synthesis may still lead to the existence of some lethal chemical species adsorbed on the surface that may have unpleasant effects in medical applications.

Hence, there is a need to develop inexpensive, clean, nonlethal and environmentally kind synthesis methods. Therefore, in last year's researchers have turned to biological systems for inspiration in nanoparticle synthesis. Microorganisms are recently found as possible environment friendly nanofactories, even though they have many biotechnological uses such as remediation of lethal metals.

It has confirmed that prokaryotes earnest powerfully as action of synthesizing nanoparticles. Suited to their over-sufficiency in atmosphere and their gift to accommodate to original broadcasting situation, bacteria are a good selection for research. Bacteria are in addition remaining maturation, economical to grow and easy to control. Heaping up diffusion such as oxygenation, incubation time and temperature can be easily manipulated. It was discovered that irregular the pH of the collecting operation by means of incubation mean in the play of nanoparticles of incompatible display and suit, predominant such transfer is standard, as unconditional morphologies of NPs are necessary for different applications such as catalysts, optics or anti-microbials.

### NPs synthesis by bacteria

Bacteria have the remarkable ability to reduce and synthesize heavy metals and nanomaterials. Some bacterial species have developed the ability to remedy to specific defense mechanisms to control stresses like toxicity of nanomaterials. Research has paying attention heavily on prokaryotes as a means to synthesize nanoparticles.

Due to their plenty in the surroundings and their ability to adjust to extreme conditions, bacteria are a good option to learn. They are also fast growing, cheap to develop and easy to control. Growth conditions such as temperature, oxygenation and incubation time can be easily controlled.

TABLE 3. Bacteria involve in synthesis of different nanoparticle.

| Bacteria   | Nanoparticle  | Size      | Morphology                  | Reference             |
|--|---|-----------|-----------------------------|-----------------------|
| <i>Aeromonas sp.</i> SH10  | Silver  | 6.4       |                             | Rai et al. [17]       |
| <i>Bacillus Cereus</i>   | Silver  | 20–40     | Spherical                   | Sunkar et al. [18]    |
| <i>Bacillus megatherium</i><br>D01   | Gold  | 1.9 ± 0.8 | Spherical                   | Wen et al. [19]       |
| <i>Bacillus subtilis</i>   | Silver  | 5–50      | Triangular and<br>spherical | Beveridge et al. [20] |
| <i>Clostridium thermoaceticum</i>  | Cadmium sulfide   |           | Amorphous                   |                       |
| <i>Desulfobacteraceae</i>  | Zinc sulfide  | 2–5       | Spherical                   |                       |
| <i>Desulfovibrio</i><br>( <i>desulfuricans, vulgaris, magneticus</i><br>strain RS-1) | Palladium, selenium,<br>gold, uranium,<br>chromium and<br>magnetite | Up to 30  | Crystalline                 |                       |

|   |   |  |  |                            |
|---|---|--|--|----------------------------|
| <i>Escherichia coli</i><br>(DH5 $\alpha$ , MC4100)  | Silver, gold                                      | Less than 10-50                            | Spherical, triangular,<br>hexagonal and<br>rod shape                                   | Mahanty et al. [21]        |
| <i>Geobacillus sp.</i>  | gold  | 5-50                                       | Quasi-hexagonal  |                            |
| <i>Klebsiella (aerogenes, pneumonia)</i>  | Cadmium sulfide,<br>silver                        | (average size of<br>52.2)                  | Spherical  | Shahverdi et al. [22]      |
| <i>Lactobacillus strains</i>  | Gold, silver, gold-<br>silver alloys,<br>titanium | 100-300                                    | Crystalline and<br>cluster   |                            |
| <i>Magnetospirillum<br/>Magnetotacticum</i>   | Magnetite   |  | Cluster  |                            |
| <i>Nocardiopsis sp.</i><br>MBRC-1   | Silver  | 45   | spherical  | Manivasagan et al.<br>[23] |
| <i>Pseudomonas</i><br>( <i>aeruginosa</i> ,<br><i>fluorescens</i> , <i>putida</i><br>NCIM 2650, <i>stutzeri</i><br>AG259) | Gold, silver,<br>lanthanum                        | 35–46 and<br>up to 200                     | Crystalline silver,<br>Hexagonal,<br>equilateral triangle,<br>and monoclinic<br>silver |                            |
| <i>Rhodobacter<br/>sphaeroides</i>  | ZnS   | Average diameter<br>of 8                   | Spherical  |                            |
| <i>Rhodopseudomonas</i><br>( <i>capsulate</i> , <i>palustris</i> )  | Gold, CdS   | 8.01 $\pm$ 0.25                            | Crystalline, FCC   | Bai et al.                 |
| <i>Shewanella algae</i><br>(strain BRY,<br><i>putrefaciens</i><br>(Gs-15) )   | Gold, magnetite                                   | Various<br>morphologies<br>altered with pH |  | Konishi et al. [15]        |
| <i>Thermoanaerobacter<br/>ethanolicus</i><br>TOR-39   | Cobalt, chromium,<br>magnetite and nickel         |  | octahedral   | Rai et al. [17]            |

### Synthesis Types

Synthesis is of two types; intracellular and extracellular.

### **Intracellular nanosynthesis**

A large number of bacterial species have been investigated for intracellular bio-nanosynthesis of metallic and non-metallic nanoparticles the details are as under;

#### **Intracellular synthesis of metallic nanomaterials**

Intracellular metal nanoparticles combine has been accounted for in different microorganisms as bio-nanofactories [25,26]. Generally, both bacterial synthesis indicated has been very important part in bio-geo cycles. This capacity has helped them in change of different mixes and basic metal in nature. Profoundly lethal centralization of metal particles is detoxified by microscopic organisms through lessening or oxidation, precipitation or complexation, transportation or efflux frameworks and so on. These properties have marked them as potential bioremediation specialists in soil and oceanic situations. In this point of view, microscopic organisms part in biogenesis of nanoparticles has additionally been explained last couple of decades. In this manner, the synthesis of intracellular metal nanoparticles including gold, silver, gold-silver alloy, platinum, palladium, uranium has been researched in a few bacterial strains.

**Gold nanoparticles:** *Bacillus subtilis* 168 encouraged gold ( $\text{Au}^{+3}$ ) particles to  $\text{Au}^0$  nanoparticles with octahedral morphology has been accounted for onto their cell dividers [26]. Another bacterium, *Geobacter ferrireducens* diminished Au particles in periplasmic space to deliver AuNPs. *Shewanella* algae likewise decreased  $\text{Au}^{+3}$  particles to natural AuNPs at 25°C in periplasmic space and on its cell surface. Under various pH conditions distinctive sizes of NPs were watched [15].

A cyanobacterium, *Plectonema boryanum* UTEX485, synthesized AuNPs at 25 to 200°C with the assistance of some external layer proteins, lipopolysaccharides and phospholipids. This blend has been connected with indicated detoxification systems in microscopic organisms [25]. Morphologically, round, triangular or hexagonal AuNPs has been bio-diminishment of wrote about cell surface in *E. coli* DH5 $\alpha$  with morphologies. Besides, this synthesis has been connected with plasma layer related NADPH-subordinate proteins and carotenoids in *Rhodobacter capsulatus* intervened biosynthesis of AuNPs.

**Silver nanoparticles:** Comprehensively, AgNPs synthesis has been connected with bacterial cell surfaces. If there should be an occurrence of *Pseudomonas stutzeri* AG259 crystalline of silver sulfide acanthite ( $\text{Ag}_2\text{S}$ ) were situated in periplasmic space with monoclinic, triangular, equilateral and hexagonal morphologies. It was connected with cell surfaces which were at first connected with biosorption and afterward diminishment  $\text{Ag}^+$  particles to frame AgNPs in *Lactobacillus* sp. A09 at 30°C [26]. For the arrangement of AgNPs, the nucleation destinations are thought to be given by silver-restricting proteins which give amino corrosive moieties. Silver precious stones with face-focused cubic morphology were accelerated by AG3 and AG4 (silver encouraging peptides). *Bacillus* sp. additionally integrated AgNPs in its periplasmic space. This aggregation or precipitation of Ag at cell divider at temperature 60°C has likewise been considered as a detoxification procedure encouraged through periplasmic proteins [5].



**Other metals:** Synthesis of bimetallic NPs has likewise been accounted for in couple of microscopic organisms, Biogenesis of silver, gold and gold-silver composites was examined by utilizing *B. subtilis* [20], *P. stutzeri*, *Lactobacillus sp.*, *Corynebacterium sp.* [5], sulfate-diminishing microscopic organisms [25], *P. boryanum* [26], *E. coli*, *R. capsulatus* and *Bacillus sp.* with circular, hexagonal and cubic morphologies. Intracellular bacterial synthesis of platinum (Pt) NPs was seen in *S. algae* in its periplasmic space amongst inward and external films. Palladium (Pd) NPs were intracellularly synthesised by *Desulfovibrio desulfuricans* NCIMB 8307 which diminished Pd nanocrystals on its cell surface with the assistance of formate (electron contributor). In a comparative study, *S. oneidensis* MR-1 was uncovered to frame Pd nanocrystals inside periplasmic space and in addition on the cell divider [27]. Uranium nanocrystals as urinate ( $UO_2$ ) were observed to be encouraged by *Desulfosporosinus sp.*, on its cell surface, which can be useful in minimizing the defilement of solvent radionuclides in soils and dregs by changing the dissolvable to insoluble structure.

**Magnetic nanomaterials:** Intracellular bacterial biosynthesis of attractive nanocrystals of magnetite ( $Fe_3O_4$ ) nanoparticles was observed to be shown by different microscopic organisms including *Aquaspirillum magnetotacticum* [3], *Magnetospirillum magnetotacticum*, *M. magnetotacticum* MS-1, *M. gryphiswaldense*, *Candidatus Magnetoglobus multicellularis*, *Magnetotactic bacterium* MV-1 [28], Sulfate decreasing microscopic organisms. Biomineralized magnetite NPs are generally synthesised by magnetotactic microorganisms which have particular structures called magnetosomes in their phones. These magnetotactic microscopic organisms live in marine and new water silt and deliver intracellular, layer bound magnetite, pyrrhotite, ferromagnetic iron sulfide, greigite ( $Fe_3S_4$ ) [28] and at some point non-attractive mineral (e.g. iron pyrite) as chains [5]. The attractive NPs are isolated from arrangement by utilizing high-angle attractive separators. The magnetosomes create both crystalline and non-crystalline attractive nanocrystals of prevalent morphologies including octahedral, exceedingly requested single or different chains, unpredictable faceted cub-octahedral shape, parallelepiped, octahedral or hexagonal crystal-like shapes, tied down and collected in phospholipids layers of microbes.

**Sulphide nanomaterials:** The semiconductor nanocrystals of cadmium sulfide (CdS) were intracellularly incorporated by different microorganisms including, *Klebsiella pneumonia* [29], *Clostridium thermoaceticum* and *E. coli* [4]. It was seen that, CdS nanocrystals were accelerated by the activity of cysteine desulphydrase which brought about desulphydration of cysteine either on cell surfaces or in medium. These bio-interceded semiconductor CdS nanoparticles displayed optical and photoactive properties and indicated round and circular shapes. In another study, round NPs of ZnS were naturally orchestrated by organisms having a place with *Desulfobacteriaceae*.

### **Non-metallic nanomaterials**

Bionanosynthesis of non-metals like selenium have likewise been examined. Different microscopic organisms including *Stenotrophomonas maltophilia* SELTE02, *Enterobacter cloacea* SLD1a-1, *Rhodospirillum rubrum*, *Desulfovibrio*

*desulfuricans*, *E. coli* [30], *P. stutzeri*, *Tetrathlobacter kashmirensis*, *P. aeruginosa* SNT1 have been found to store selenium NPs after bio-decreasing selenite to essential selenium in cell cytoplasm, periplasmic space and extracellularly with various shapes like granular, circular, fibrillar or in totals [31].

TABLE 4. Intracellular synthesis of nanoparticles by bacteria.

| Organism                             | Metal/Non-metal                | Size (nm) | Location of synthesis | Shapes         | References                |
|--------------------------------------|--------------------------------|-----------|-----------------------|----------------|---------------------------|
| <i>Idiomarina</i> spp. PR58-8        | Ag                             | 26        | Intracellular         |                |                           |
| <i>Pseudomonas</i> spp.              | Ag                             | 156-265   | Intracellular         |                | Thomas et al. [30]        |
| <i>Bacillus subtilis</i> 168         | Au                             | 5–25      | Intracellular         | Octahedral     | Beveridge and Murray [20] |
| <i>Shewanella algae</i>              | Au                             | 10-20     | Intracellular         |                | Lengke et al. [25]        |
| <i>Plectonema boryanum</i> UTEX485   | Au                             | 10        | Intracellular         | Cubic          | Lengke et al. [25]        |
| <i>Escherichia coli</i> DH5 $\alpha$ | Au                             |           | Intracellular         | Spherical      |                           |
| <i>P. stutzeri</i> AG259             | Ag, Ag <sub>2</sub> S          | 200       | Intracellular         |                |                           |
| <i>Corynebacterium</i> spp. SH09     | Ag                             | 10-15     | Intracellular         |                | Zhang et al. [5]          |
| <i>Bacillus</i> spp.                 | Ag                             | 5-15      | Intracellular         |                |                           |
| <i>Lactobacillus</i> spp.            | Au, Ag, Au–Ag                  | 20-50     | Intracellular         | Hexagonal      |                           |
| <i>P. aeruginosa</i> SNT1            | Se                             |           | Intracellular         | Spherical      | Yadav et al. [29]         |
| <i>Desulfovibrio desulfuricans</i>   | Pd                             | 50        | Intracellular         |                |                           |
| <i>S. oneidensis</i> MR–1            | Pd                             |           | Intracellular         |                | De Windt et al. [27]      |
| <i>Aquaspirillum Magnetotacticum</i> | Fe <sub>3</sub> O <sub>4</sub> | 40-50     | Intracellular         | Octahedral     | Mann et al.[3]            |
| <i>Magnetotactic bacterium</i> MV-1  | Fe <sub>3</sub> O <sub>4</sub> | 40×40×60  | Intracellular         | Parallelepiped | Bazylinski et al. [26]    |

|                           |           |        |               |                                  |  |
|---------------------------|-----------|--------|---------------|----------------------------------|--|
| <i>M. gryphiswaldense</i> | Magnetite | 35-120 | Intracellular | Cubo-<br>octahedral<br>hexagonal |  |
|---------------------------|-----------|--------|---------------|----------------------------------|--|

### Extracellular nanosynthesis

A number of bacteria have been studied for their potential of extracellular bionanosynthesis.

### Extracellular synthesis of metallic nanomaterials

**Gold nanoparticles:** Bioreduction of  $\text{Au}^{+3}$  into metallic AuNPs was seen by *Rhodopseudomonas capsulata* at room temperature. The particles were obtained with round and triangular morphology. Arrangement of various sizes and states of these NPs were discovered pH division of the response mixture.

Besides, nearness of diminishing and topping proteins of 14 kDa to 98 kDa size was confirmed through SDS-PAGE investigation. Iron ( $\text{Fe}^{+3}$ ) thinning archea and bacteria, for example, *Pyrobalaculum islandicum*, *S. algae*, *G. sulfurreducens*, *Thermotoga maritim* and *Pyrococcus enraged*, interceded precipitation of gold from ionic to metallic structure within the sight of hydrogen.  $\text{Au}^{+3}$  reductases close to the cell surfaces brought on this precipitation. *P. aeruginosa* (ATCC 90271, strain1 and strain 2) interceded synthesis of AuNPs was uncovered. Because of surface plasmon sound of greater NPs, medium shading changed from pink to blue.

*B. megatherium* D01 in dried structure synthesized spherical AuNPs at 26°C by reducing gold salts by utilizing dodecanethiol as topping specialist for balancing out NPs furthermore adjusting size, shape and monodispersity of NPs [20]. The morphologies of AuNPs manufactured by *P. aeruginosa*, *R. capsulata* and *B. megatherium* D01 were observed to be nanowires triangular and round fit as a fiddle extending in size from 1.9 nm to 400 nm.

**Silver nanoparticles:** Dried cells of *Aeromonas* sp. SH10 were utilized to deliver mono dispersed AgNPs with uniform size and security. A fast combination of AgNPs was seen by bioreduction of silver ( $\text{Ag}^{+}$ ) particles to metallic  $\text{Ag}^0$  by the activity of society supernatants of *E. coli*, *Enterobacter cloacae* and *K. pneumonia* [23]. The bioreduction was hindered when piperitone was added to the response blend which affirmed the nearness of nitro-reductase. Extracellular amalgamation of AgNPs was additionally seen by *B. licheniformis* [33].

Biogenesis of AgNPs was additionally exhibited in cellulose layers of *Acetobacter xylinum* when bacterial societies were presented to the arrangement containing  $\text{Ag}^{+}$  particles and tri-ethanol-amine ( $\text{Ag}^{+}$ -TAE). Extracellular blend of AgNPs with

round morphology was seen by microbes, for example, *E. cloacae*, *Aeromonas* sp. SH10, *E. coli*, *A. xylinum*, *B. licheniformis*, *Morganella* sp. and *K. pneumonia*.

**Other nanoparticles:** Nanoparticles of platinum were accounted for extracellularly by the activity of different microscopic organisms, for example, *P. boryanum* UTEX 485 and Cyanobacterium with different morphologies containing dendritic, circular and dab like chains [25]. *P. boryanum* UTEX 485 integrated PtNPs at 25 to 100°C. Round molded NPs of titanium as totals were combined extracellularly by *Lactobacillus* sp. society filtrate [8].

Acellular concentrates of *Micrococcus lactilyticus* accelerated uranium NPs by lessening dissolvable  $U^{6+}$  to insoluble  $U^{4+}$  [34]. *Alteromonas putrefaciens* created uranium NPs responded with hydrogen (electron giver) and  $U^{6+}$  (electron acceptor).

Uranium NPs were delivered by *G. metallireducens* GS-15 when under anaerobic conditions acetic acid derivation (electron benefactor) and  $U^{6+}$  (electron acceptor) did bioreduction of uranium particles. Because of nearness of MtrC (c-sort cytochrome) on outside layer of *S. oneidensis* MR<sup>-1</sup>, it did bioreduction of uranium particles with UO<sub>2</sub>-EPS (polymeric substance) extracellularly and in addition in periplasm.

**Magnetic nanomaterials:** Extracellular biosynthesis of magnetite NPs has been done by some non-magnetotactic microorganisms. *Geobacter metallireducens* GS-15, a non-magnetotactic bacterium disconnected from waterway base, created ultrafine magnetite NPs by lessening ferric oxide. In this procedure ferric particles (electron acceptors) responded with natural matter (electron benefactors).

Magnetite nanoparticles with octahedral morphology were incorporated extracellularly at 25°C by another bacterial strain named as TOR-39. Microscopic organisms were noted to go about as biocatalysts for precipitation of magnetite NPs. Ferric reduction microscopic organisms, TOR-39 and *Thermoanaerobacter ethanolicus* intervened electrochemical response which octahedral move metals (Ni, Cr, Co) substituted attractive nanocrystals were integrated.

The creation of semi round magnetite nanocrystals were exhibited by [8] by utilizing *Actinobacter* sp. (non-magnetotactic bacterium). Magnetite NPs with semi round and octahedral morphologies were incorporated extracellularly by TOR-39, *G. metallireducens* and *Actinobacter* sp. [8]. Cubic spinel-molded single-crystalline ferromagnetic Co<sub>3</sub>O<sub>4</sub> nanoparticles were synthesised by *Brevibacterium casei* [35] which is a metal-tolerant bacterium by utilizing watery cobalt acetic acid derivation as antecedent.

**Sulfide nanomaterials:** *Klebsiella aerogenes* was set up to create circular formed nanocrystals of CdS bound on cell divider after reduction of Cd<sup>2+</sup> in society medium [31].

A photosynthetic bacterium *Rhodospseudomonas palustris* as showed by Bai et al. [36], created extracellular nanocrystals of CdS with round morphology. The creation of CdS nanocrystals was thought to be interceded by activity of cysteine desulphydrase. If there should be an occurrence of *Gluconoacetobacter xylinus*, after precipitation CdS nanoparticle were observed to be stored on bacterial cellulose nanofibers [6]. *R. palustris*, *K. aerogenes* [29], *R. spheroids'* and *G. xylinus* [6] integrated semiconductor NPs with circular shape.

Rhodobactersphaeroides [5], in immobilized structure, created monodispersed, circular semiconductor zinc sulfide (ZnS) and lead sulfide (PbS) nanoparticles extracellularly.

### Non-metallic nanomaterials

Selenium has semiconducting and photo-optical properties with applications in microelectronic circuits and scanners like gadgets. Uniform stable nanospheres of selenium were extracellularly synthesized by couple of microbes including *B. selenitireducens*, *Selenihalanaerobac shriftii* and *Sulfurospirillum barnesii* by changing essential selenium ( $Se^0$ ) to monoclinic crystalline structure with one of a kind and complex nanosized collection of selenium particles by the procedure of bioreduction [37]. Reduction of telluride to natural tellurium by *S. barnesii* and *B. selenitireducens* brought about the synthesized semiconductor tellurium NPs. If there should arise an occurrence of *S. barnesii*, little sporadic crystalline nanospheres were made [7].

TABLE 5. **Extracellular synthesis of nanoparticles.**

| Organisms                          | NPs    | Synthesis location | Method    | References             |
|------------------------------------|--------|--------------------|-----------|------------------------|
| <i>Thermomonospora sp.</i>         | Au     | Extracellular      | Reduction | Kasthuri et al. [36]   |
| <i>Escherichia coli</i>            | Pd, Pt | Extracellular      | Reduction | Park et al. [37]       |
| <i>Rhodospseudomonas capsulata</i> | Au     | Extracellular      | Reduction |                        |
| <i>Pseudomonas aeruginosa</i>      | Au     | Extracellular      | Reduction | Narayanan et al. [38]  |
| <i>Delftia acidovorans</i>         | Au     | Extracellular      | Reduction |                        |
| <i>Shewanella sp.</i>              | AsS    | Extracellular      | Reduction | Raveendran et al. [39] |
| <i>Desulfovibrio desulfuricans</i> | Pd     | Extracellular      | Reduction | Cai et al. [11]        |

|                                   |                      |               |                              |                       |
|-----------------------------------|----------------------|---------------|------------------------------|-----------------------|
| <i>Bacillus sphaericus</i> JG-A12 | U, Cu, Pb,<br>Al, Cd | Extracellular | Reduction and<br>Biosorption | Das et al. [30]       |
| <i>Klebsiella pneumonia</i>       | Ag                   | Extracellular | Reduction                    | Kalimuthu et al. [14] |
| <i>Escherichia coli</i>           | Ag                   | Extracellular | Reduction                    | Kalimuthu et al. [14] |
| <i>Enterobacter cloacae</i>       | Ag                   | Extracellular | Reduction                    | Kalimuthu et al. [14] |
| <i>Lactobacillus sp.</i>          | Ag                   | Extracellular | Reduction and<br>Biosorption | Shahverdi et al. [22] |
| <i>Enterococcus faecium</i>       | Ag                   | Extracellular | Reduction and<br>Biosorption | Shahverdi et al. [22] |
| <i>Lactococcus garvieae</i>       | Ag                   | Extracellular | Reduction and<br>Biosorption | Shahverdi et al. [22] |

## Materials and Methods

### Silver NPs synthesis by *Bacillus sp.*

The sample was collected from the most contaminated area common toilet in university campus. The isolated organism was maintained for nutrient agar medium. The organism was cultivated in 1 litre of medium containing 1.0 gm of beef extract, 2.0 gm of yeast extract, 5 gm of sodium chloride, 5 gm of peptone, and 15 gm of agar. Than incubated at 35°C. The isolates were morphologically and microbiologically characterized as *Bacillus sp.*

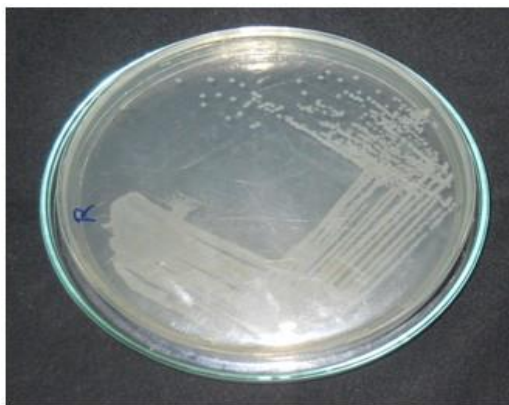


FIG. 2. *Bacillus sp.* in Nutrient agar containing plate.

For the silver nanoparticles synthesis, the optimized biomass and without optimized *Bacillus* sp. strain was used. One loopful culture of *Bacillus* sp. was inoculated in nutrient broth. Then the broth was placed in the orbital shaker for 24 h for the growth of culture. Then 1 mM of Silver Nitrate crystals were weighed and added with 24 h incubated 100 ml of culture and kept it in a shaker at 200 rpm per minute.

The UV-visible spectrum of the solution was recorded in Perkin – Elmer spectrophotometer. The particles wavelength ranges from 300 nm to 700 nm. The shape and particle sizes were determined by Scanning electron microscope (SEM) that was performed by focusing on nanoparticles.

### Visual identification

When bacterial biomass mixed with silver nitrate solution (pH 7.5) and incubated for 24 h at 37°C, it shows no change in color. The appearance of milk fish white color is apparent sign of the formation of silver nanoparticle in the mixture after 12 h incubation (FIG.3 (a)). The production of nanoparticle is whitish brown in color (FIG.3 (b)). Due to excitation of surface plasmon vibrations of silver nanoparticle brown color appears.



FIG.3. Biosynthesis of silver nanoparticle using *Bacillus* sp. (a) with addition of cadmium sulphate 12 h incubation (b) 24 h incubation.

### UV-Vis spectrophotometer

The characterization of synthesized silver nanoparticle was primarily done by UV spectrophotometer. At different time intervals, the UV-visible spectra recorded showed increased absorbance with increasing time of incubation. Figure describes the absorbance spectra of reaction mixture containing 1 mM silver nitrate aqueous solution and the culture supernatant of *Bacillus* sp. after incubation. The intensity of the color from the cells harvest at the stationary phase was stable and also

maximum. The band related to the surface plasmon resonance is at 410 nm to 430 nm. The strongest peak is at 420 nm. The accurate mechanism for the NPs synthesis has not been visibly established but an enzyme NADH dependent nitrate reductase is concerned in the process.

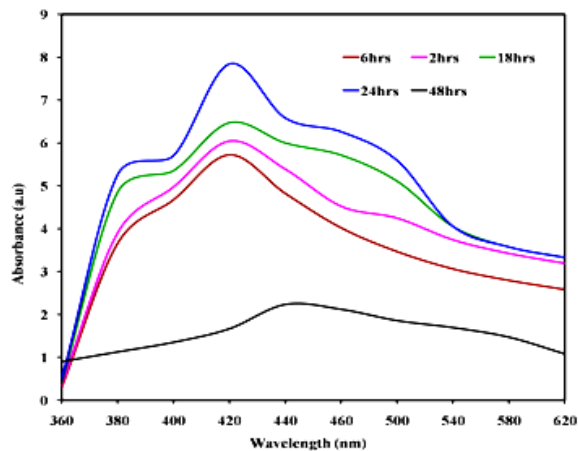


FIG. 4. UV-visible spectrum of silver NPs biosynthesis by using *Bacillus* sp.

#### SEM with EDX

The shape of the silver NPs was analyzed by using the Scanning Electron Microscope. SEM images of the optimized bacterial culture moderate silver nano powder having spherical, pseudo-spherical and some in determinate shapes with traces of agglomeration because of the biological molecules combine with the NPs in the bacteria.

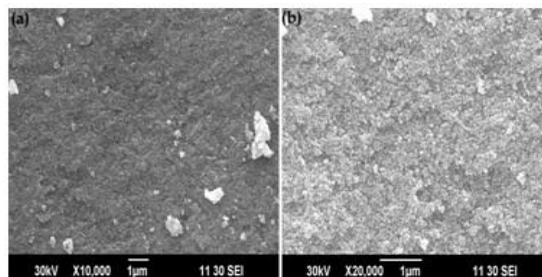


FIG.5. SEM images of the silver NPs created by the reaction of 1 mM  $\text{AgNO}_3$  and *Bacillus* sp. broth.

In the EDX spectrum of the bacterial moderate silver NPs, the strongest peak detected was from silver and weaker peaks detected from carbon and oxygen. This shows that the silver NPs biosynthesis is relatively pure in chemical composition. This result was closely related with the SEM results.



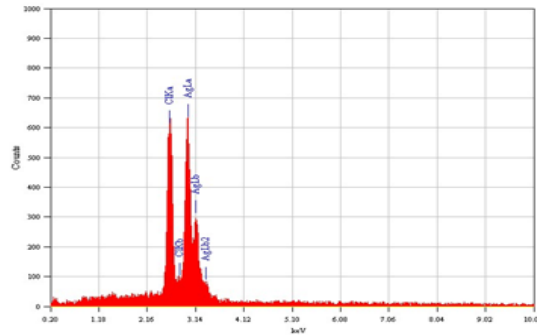


FIG. 6. SEM images of the silver NPs produced by the reaction of 1 mM AgNO<sub>3</sub> and *Bacillus* sp. broth.

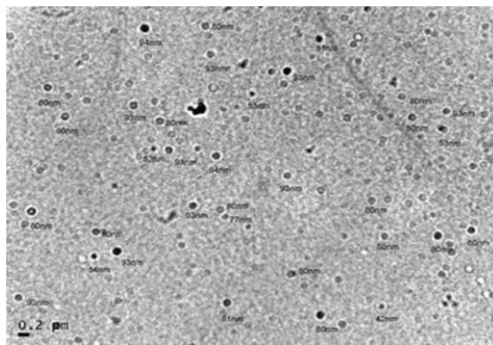


FIG. 7. TEM analysis of the silver nanoparticles synthesized by *Bacillus* sp.

## Mechanism

### Ability of bacteria

The capacity of bacteria to survive and develop in unpleasant circumstances may be because of particular instruments of resistance which incorporate efflux pumps, metal efflux frameworks, inactivation and complexation of metals, impermeability to metals and the absence of particular metal transport frameworks, adjustment of solvency and poisonous quality by changes in the redox condition of the metal particles, extracellular precipitation of metals and volatilization of lethal metals by enzymatic responses [42].

For instance, *Pseudomonas stutzeri* AG259 disengaged from silver mines has been appeared to create silver NPs [43]. There are a few cases of microorganisms-metal associations which are imperative in biotechnological applications, including the fields of biomineralization, bioremediation, bioleaching and microbial influenced corrosion (MIC) forms.

### Acquaintance of MIC

Acquaintanceship of MIC procedures in structure of microbially intervened restricted changes in the surface science of carbon steel, stainless steel, copper combinations, or different ones is increasing developing consideration [44]. Microscopic organisms likewise mediate in mineral precipitation responses specifically as impetuses of watery substance responses as

geochemically receptive solids [45] and demonstrated the capacity to oxidate minerals [46]. These procedures are economically utilized as a part of bacterial filtering operations, for example, the pretreatment of gold metals which contain arsenopyrite (FeAsS) [46].

Microbial metal decrease can be a system for *in situ* and *ex situ* remediation of metal pollutions and squanders. With a specific end goal to discover the significance of nanoparticle combination and metal lessening, bio-recovery of substantial metals and bioremediation of dangerous ones, analysts have researched the systems of nanoparticle synthesis and bio-reduction and centered their consideration on decreasing specialists in microbes (e.g., proteins and catalysts) and biochemical pathways prompting metal particle diminishment.

As a result of the basic part of these specialists, there were more examinations in comprehension the part and utilization of regular and hereditarily designed bacterial strains and different microorganisms in bioremediation of lethal metals and radionuclide-debased earthbound situations. Besides, these microorganisms were equipped for preparation and immobilization of metals [47] and at times, the microscopic organisms which could diminish metal particles demonstrated the capacity to accelerate metals at nanometer scale.

These studies would then prompt check the likelihood of hereditarily designed microorganisms to over express particular lessening atoms and to create microbial nanoparticle blend systems, which may conceivably control size, shape, solidness and yield of NPs. Really, hereditarily designed microorganisms have begun being created so as to build protein discharge and accordingly clarify the most plausible diminishing specialist.

For example, Kang and his co-workers [48], investigated interestingly the deliberate methodology toward the tunable combination of semiconductor CdS nanocrystals by hereditarily designed *E. coli*. To investigate the plausibility of utilizing *E. coli* as a biofactory for the controlled blend of CdS nanocrystals, a strain was blessed with the capacity to create phytochelatins (PCs) by communicating the PC synthetase of *S. pombe* (SpPCS). PCs serve as a coupling format site for the metal particles and balance out the nanocrystals center against proceeded with accumulation.

### **Use of recombinant strains**

Recombinant strains have been investigated for growing more productive life forms in the *in vivo* synthesis of NPs. For example, recombinant *E. coli* strains communicating Arabidopsisthaliana phytochelatin synthase (AtPCS) and *Pseudomonas putida* metal-lothionein (PpMT) were utilized for the blend of Cd, Se, Zn, Te, Cs, Sr, Fe, Co, Ni, Mn, Au, Ag Pr and Gd NPs. Changing the groupings of supplied metal particles brought about controlling the extent of the metal NPs. It was accounted

for that the designed *E. coli* framework can be material to the organic blend of metal NPs [39]. Mutant strains of a few microscopic organisms utilized for union of NPs could clarify the atoms required in the bioreduction procedure.

If there should arise an occurrence of *Acidithiobacillus thiooxidans*, gold (I)-thiosulfate was gone into the cell of a thiooxidans as a component of a metabolic procedure [26]. This gold complex was at first decomplexed to Au (I) and thiosulfate ( $S_2O_3^{2-}$ ) particles. Thiosulfate was utilized as vitality source and Au (I) was apparently lessened to basic gold inside the bacterial cells. Amid the late stationary development stage, the gold NPs which were at first accelerated inside the cells were discharged from the cells, bringing about the arrangement of gold particles at the phone surface. At long last, the gold particles in the mass arrangement were developed into micrometer-scale wire and octahedral gold [26].

As per Lengke and Southam [25], the precipitation of gold (I)-thiosulfate complex by sulfate-diminishing microorganisms was brought about by three conceivable components: (an) iron sulfide development, (b) confined decreasing conditions and (c) a metabolic procedure.

### **Reduction**

If there should be an occurrence of *D. desulfuricans* and *E. coli*, fractional restraint of periplasmic hydrogenases with Cu (II) demonstrated that these metal reductase compounds assume a part in Au (III) diminishment [49]. Periplasmic hydrogenases were conceivably in charge of Pd (II) lessening and repressed by Cu (II).

Moreover, Au (III) decrease was done within the sight of  $H_2$  (as the electron contributor) utilizing microorganisms, for example, *T. maritima*, *S. alga*, *D. vulgaris*, *G. ferrireducens*, *D. desulfuricans* and *E. coli*. Perhaps, hydrogenases assume an essential part in Au (III) lessening, however more examinations were expected to know accurate instruments of these reductions. Moreover, it was accounted for that hydrogenase is included in  $U^{+6}$  decreases by *Micrococcus lactyliticus*, notwithstanding  $Se^{+6}$  diminishment by *Clostridium pasteurianum* [34]. Hydrogenases from the sulfate-diminishing microorganisms have been appeared to be fit for lessening  $Tc^{+7}$  and  $Cr^{+6}$  [50].

In another study, it was accounted for that the hydrogenases separated from phototrophic microscopic organisms could lessen  $Ni^{+2}$  to  $Ni^0$  under a  $H_2$  environment [51]. Matsunaga and his co-workers have indicated that MagA quality and its protein (confined from *Magnetospirillum* sp. AMB-1) were required for biomagnetic nanoparticle arrangement [52].

Magnetotactic microscopic organisms (e.g. *M. magnetotacticum* and *M. gryphiswaldense*), contain magnetosome film (MM) proteins which assume an essential part in magnetite biomineralization. Consequently, analysts have concentrated on ID of these proteins and their qualities. Late sub-atomic studies including genome succession, mutagenesis, quality expression and

proteome investigations showed various qualities and proteins which assume basic parts for bacterial attractive particles biomineralization [53].

Moises cu and his co-workers have concentrated on the concoction arrangement and microstructural attributes of bacterial magnetosomes removed from the magnetotactic bacterial strain *M. gryphiswaldense* [54]. They reported the created octahedral magnetite particles with a normal distance across of  $46 \pm 6.8$  nm.

The particles displayed a high synthetic virtue (only  $\text{Fe}_3\text{O}_4$ ) and the dominant parts fall inside the single-attractive area range. In instances of *Rhodopseudomonas capsulata* and *Stenotrophomonas maltophilia*, the creators trusted that the particular NADPH-subordinate chemical present in the detached strains lessened  $\text{Au}^{+3}$  to  $\text{Au}^0$  through an electron transporting system, prompting the blend of monodispersed NPs. A two-stage procedure is expected to lessen gold particles.

Among the initial step, the  $\text{AuCl}_4^-$  particles are decreased to the  $\text{Au}^+$  species. At that point, the last item is lessened by NADHP to a metallic gold [55].

#### **Size of magnetic crystals**

To control the morphologies and sizes of NPs, there were a few examinations concentrated on utilizing proteins. Strangely, relationship of proteins with spheroidal totals of biogenic zinc sulfide nanocrystals reported that extracellular proteins began from microorganisms could restrain the biogenic NPs [56].

Controlled arrangement of magnetite precious stones with formally dressed size was accomplished within the sight of Mms6 (a little acidic protein confined from *Magnetospirillum magneticum* AMB-1) [57]. The normal size of magnetite precious stones combined within the sight of Mms6 was around  $20.2 \pm 4.0$  nm. Be that as it may, without Mms6, the orchestrated magnetite gems were around  $32.4 \pm 9.1$  nm.

Along these lines, the gems integrated with Mms6 were littler than precious stones created without Mms6 and were disseminated over a smaller reach than gems orchestrated without the protein. Mms6 additionally advanced development of uniform, isomorphic, superparamagnetic nanocrystals [58].

With a bioinspired technique, Prozorov and his co-workers have reported the utilization of the recombinant Mms6 protein for union of uniform, all around characterized  $\text{CoFe}_2\text{O}_4$  nanocrystals *in vitro* [59]. To layout progressive  $\text{CoFe}_2\text{O}_4$  nanostructures, a recombinant polyhistidine-labeled full-length Mms6 protein and an engineered C-terminal area of this protein were covalently appended to triblock copolymers (poloxamers).

In the event of *Klebsiella pneumoniae*, it was accounted for that no arrangement of silver NPs by the supernatant was watched when the strategy occurred unconscious. The visible light depletion can essentially bring about the synthesis of NPs. It appears that for this situation, the silver particles diminishment was mostly because of conjugation transports with the investment of the reductase.

In this way, it gives the idea that the cell-related nitroreductase chemicals might be included in the photoreduction of silver particles [16]. Also, systems of cadmium sulfide nanocrystals union by *E. coli* cells were clarified through the control tests (brooding of  $\text{CdCl}_2$  and  $\text{Na}_2\text{S}$  without bacterial cells) which showed that nanocrystals were not incorporated outside the cells and after that shipped into the cells [4].

These examinations have demonstrated that CdS nanocrystals could be blended after  $\text{Cd}^{2+}$  and  $\text{S}^{2-}$  particles transported into the cells. In the event of zinc sulfide (ZnS), the NPs could be framed intracellularly through the organic manufactured technique recommended by Bai and his co-workers [5].

They clarified that solvent sulfate diffused into immobilized dots and after that was conveyed to the inside layer of *R. sphaeroides* cell encouraged by sulfate permease.

## **Applications**

### **Multidisciplinary in nature**

Nanotechnology is a multidisciplinary field in nature regarding investigations and applications [60]. In most recent couple of decades, inquiries about in building, physical science, natural chemistry and microscopy have prompted magnificent additions of worry in portrayal minor particles and their promising ramifications in various territories of material science. Enhance the analytic and treatment Nanotechnology is a multidisciplinary field in nature regarding investigations and applications [60]. In most recent couple of decades, inquiries about in building, physical science, natural chemistry and microscopy have prompted magnificent additions of worry in portrayal minor particles and their promising ramifications in various territories of material science. Enhance the analytic and treatment

### **Nano-biomedicine**

Discrete NMs are typically connected in nano-biomedicine as fluorescent natural labels and middle people for medication and additionally drug deliverance. Although this they can likewise be utilized for discovery of pathogens, tissue designing [7], tumor destruction, contrast change in (MRI) and phagokinetic examinations.

TABLE 6. Applications of NPs.

| Nanoparticle                   | Applications  |
|--------------------------------|---|
| Ag                             | ⇒ home appliances as antimicrobial agents<br>⇒ clothing for odor resistance       |
| TiO <sub>2</sub>               | ⇒ paints and coating for antimicrobial properties<br>⇒ cosmetics as a UV absorber |
| Carbon nanotubes (CNT)         | ⇒ consumer electronics<br>⇒ sports equipment for light weight and durability      |
| Fe <sub>2</sub> O <sub>3</sub> | ⇒ contrast agent for targeted tumor imaging                                       |
| Fullerenes                     | ⇒ drug delivery   |
| Fe                             | ⇒ environmental remediation   |
| Au                             | ⇒ medical diagnostics   |

### Incomprehensible as antimicrobial

Amongst various metal NPs, AgNPs have been widely considered surgical gloves and covers, antibacterial injury dressings, bed lines and so forth [61]. They likewise have various applications in the fields of gadgets, catalysis and indicative therapeutics.

### Biosensors

Similarly, gold nanoparticles (AuNPs) are additionally used for various purposes e.g. as labels for biosensors, for cure of hyperthermia and being equipped for conveying vast estimated biomolecules, give non-dangerous intends to quality and medication liberation to the objective locales [62].

Platinum nanoparticles (PtNPs) are utilized for the treatment of various illnesses, for example, growth and oxidative anxiety issue; in addition, they are utilized as a part of gadgets for planning of a novel memory component.

### Environmental remediation

NMs offer the potential for the productive expulsion of toxins and natural contaminants from the earth. NMs capacity as adsorbents and catalysts in different shapes, for example, nanoparticles, tubes, wires, filaments and so forth and their composites with polymers are utilized for the discovery and evacuation of gasses, for example, (SO<sub>2</sub>, CO, NO<sub>x</sub> and so on.),

violated chemicals (nitrate, arsenic, iron, substantial metals, manganese and so forth.), natural poisons (aliphatic and fragrant hydrocarbons) and characteristic substances, for example, (parasites, infections, microorganisms and anti-infection agents). Due to high surface region and high reactivity NMs demonstrate a superior execution in natural remediation than other regular procedures.

In sort to live in environment, containing high amounts of metals, life forms should have adjusted by advancing instruments to handle with them. These systems may include changing the potion way of the lethal metal so that it no more causes danger, bringing about the development of NPs of the metal concerned.

### **Clinical operations**

These days in the clinical operations, *in vivo* sign of NPs in helpful capacities is a developing order. Although, there are still a few ambiguities with respect to their configuration and ramifications of NMs, for example, their pharmacokinetics, conveyance in the body, poisonous quality and security assessment earlier and consequent to their conjugation in therapeutic methods. Therefore, the biosimilarity of these NMs is still unclear and distinctive planning trials are in progress to maintain a considered distance from these injurious impacts. As yet, inferable from the nontoxic nature, AuNPs are broadly utilized as a part of healing and imaging operations.

Despite the fact that, it is impractical to make NMs that are non-hazardous to every single living cell, as there is no all-inclusive instrument for this reason, yet some essential examinations have been completed keeping in mind the end goal to make them more biocompatible and mitigate their cytotoxic impacts both *in vitro* and *in vivo* conditions.

### **Desired response**

With the utilization of least conceivable dosage is typically regarded as an alluring answer for biocompatibility issues. Besides, the covering of NMs is additionally essential as broken, split or harsh packaging could trigger the safe reaction, in this way diminishing their bioavailability to the sought target locales [63].

Although worthy metals, NMs have developed as promising instruments against a standout amongst the most difficult human sickness i.e. tumor, however their definitive destiny in the body is still unclear with a specific end goal to interpret their general helpful potential.

### **For cancer therapy**

### **Optimization and Methodical Protocols**

Optimization may topple reduce to the enhanced biosynthesis of NPs. Methodical protocols could be used for alloy of decidedly definite and well-characterized NPs right away on the quinine aspects, such as types of organisms, hereditary and genetical offering of organisms, incomparable publication for partition collecting and enzyme activity, optimal counteraction

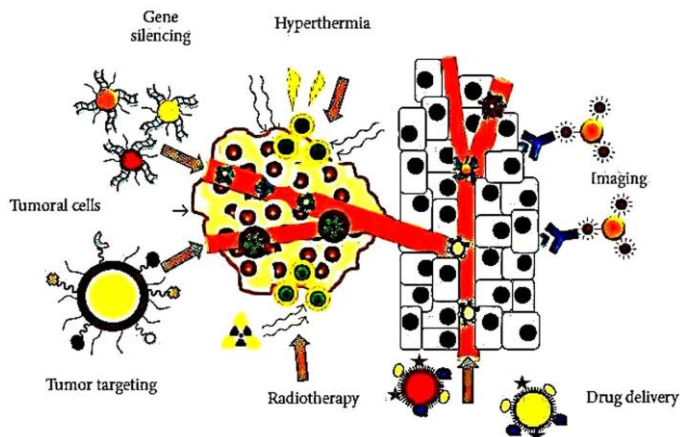


FIG.8. Applications of noble metal NPs for cancer therapy.

issuance and selection of the biocatalyst state have been considered. Morphology of the NPs depths is comfortable by regulating the prior reaction conditions. Industrial scale combine of metal NPs gulp biomass needs diverse processes, in addition transmission enchant, insusceptibility of the transmit into the biomass, harvesting the cells make up of NPs by additionally metal ions to the cells, breaking of cells by classiness, homogenization of the cells to isolate the produced NPs, stabilization of the NPs, product formulation and quality control.

TABLE 7. Nanoparticle applications and their toxicity.

| Nanoparticle     | Applications   | Toxicity  | References  |
|------------------|--|---|---|
| Titanium dioxide | Titanium nanoparticles have been applied in the pharmaceutical industry as drug delivery vehicles and in excipient formulations. | The composite material combines the high adsorption capability of apatite with the photocatalytic activity of titanium. Apatite coatings may thus become useful in the attenuation of the toxicological effects of inorganic metal oxide nanoparticles. | (Acosta-Torres et al. [62]; Driscoll et al., [63] Salata et al. [64] Cantado et al. 2008; [65] Guo et al. [66]; Sikong et al.[67] |
| Iron oxide       | Used in cellular therapy, such as cell labeling and  | No toxicity reported.   | (Acosta-Torres et al. [62] Arbab et   |



|        |   |   |   |
|--------|---|---|---|
|        | targeting and as a tool for cell biology research to separate and purify cell populations.<br>Also used in:<br>tissue repair;<br>drug delivery; magnetic resonance imaging;<br>hyperthermia;<br>magnetofection.   |   | al. [68] Sanpo et al. [19]                                      |
| Silver | Used for covering urinary catheters, surgical instruments and bone prostheses.<br>Additionally, silver has been used in water and air filtration to eliminate microorganisms.<br>AgNPs have been added to soft tissue Conditioners for prosthetic devices | Exposure of metal-containing nanoparticles to human lung epithelial cells generates ROS, which can lead to oxidative stress and cellular damage (AgNPs).<br>Silver nanowires resulted in the strongest cytotoxicity and immunological responses, whereas spherical silver particles had negligible effects on cells when tested in human cells. | (Limbach et al. [69] Niño-Martínez et al. [70] Ahn et al. [71]) |

### Conclusion

Nanoparticles are having a great deal of utilizations in different fields like antimicrobials, additives, paints, biosensors and makeup. The use of microbes is the great way to deal with the creation of Eco-accommodating and costs adequate nanoparticles. Advancement of nanoparticles union by enhanced generation of society development will giving a decent prospect for the most extreme manufacture and it is exceptionally helpful for the numerous nanoparticles based applications. The utilization of microscopic organisms offers a method for creating "manufactories" for generation of nanoparticles essentially and reasonable. It is likewise apparent that NPs have extraordinary prospective in a wide range of businesses. The

field of organic creation of nanoparticles is moderately new and underexplored; in any case it demonstrates awesome potential in the biotechnology part.

## Future Prospects

The research work clearly predicted the NPs synthesis abilities of bacteria; nonetheless, it could be enhanced as far as rate and quality by exploring proficient microorganisms which are safe and expressive against metals by delivering certain synergist proteins activated through various qualities.

Assessment and comprehension the fundamental systems in term of biomolecules required in changing and balancing out the metal particles into Nano-composites.

Choose the best bacteria, which have the best growth rate and characteristics to synthesize NPs with specific and smaller shapes.

Scaling up the laboratory methods to the industrial scale that may lead to the enhanced synthesis of NPs.

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