**Title: Plant-Microbe Interactions: A Brief Overview**

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**Abstract**

Interactions between plants and microbes significantly impact how plants behave, grow, and evolve. Numerous microbial species, such as bacteria, fungi, viruses, and archaea, which colonise the rhizosphere, phyllosphere, and endosphere of the plant, are involved in these complex associations. Depending on the traits and functions of the microorganisms and how they affect the plant, these interactions may be advantageous or detrimental. Positive relationships between plants and microorganisms are crucial for nutrient uptake, stress tolerance, and disease resistance. Plant-associated microorganisms can improve nutrient availability through several methods, including nitrogen fixation, phosphate solubilization, and iron mobilisation. They may also create phytohormones, which promote the growth and development of plants. Additionally, certain beneficial microorganisms function as biocontrol agents, inhibiting pathogenic growth and safeguarding plants from illnesses. Complex molecular signalling networks, such as the flow of chemical signals between plants and microorganisms, frequently facilitate these interactions. On the other hand, certain microorganisms can infect plants, resulting in serious output losses. Plants may become infected by pathogens through wounds, holes in the environment, or direct plant tissue penetration. They create chemicals and enzymes that interfere with the plant's defences and impair its immune system. Pathogens can also hinder the intake of nutrients and interfere with regular physiological functions, which compromises the health of the plant. For sustainable agriculture and the proper operation of ecosystems, it is essential to comprehend the subtleties of plant-microbe interactions. Utilising advantageous interactions can result in the creation of innovative techniques including biofertilizers, biocontrol agents, and bioremediation. These strategies have the potential to lessen the environmental effects of agriculture while increasing crop output and reducing chemical inputs. The study of plant-microbe interactions has been transformed by advancements in next-generation sequencing technology, omics techniques, and bioinformatics tools, which have made it possible to identify and characterise complex microbial populations linked to plants. Our comprehension of these intricate linkages will be improved by more research in this area, which will also open the door to creative agricultural techniques that might help solve problems with environmental sustainability and global food security.

**1. Introduction**

The coexistence of plants and microbes from time immemorial has shaped the ecosystem and environmental balance. The association between these two are either positive (confers protection in biotic and abiotic stress condition) or harmful by the act of pathogenesis (Newton et al., 2010; Adeleke et al., 2019). These associations are classified into rhizosphere, endosphere and phyllosphere microbiomes based on niche (interaction of microbes with their ambient environment). The study of these three microbiomes gives us insights into the characteristics and functions of the microbial communities. Whereas, in the perspective of plants, we have a generalized concept that the root’s only function is anchorage and uptake of nutrients and water from the soil. But the study of these interactions shows that roots are also involved in the secretion of several chemicals and exudates which invites microbes from different parts of soil to the rhizosphere and endosphere too.

Plant-microbe associations are influenced by several biotic and abiotic factors like host species, genotype, immunity, and climatic changes. In the recent scenario of “organics” and “natural” microbes can be utilized as a potential tool which acts as biocontrol organisms, biofertilizers, and bioremediation. All such applications are not only beneficial to humans but also help to restore the environmental balance, which has been destroyed by several anthropogenic activities and industrialization.

The recent surge in new omics technologies like genomics, transcriptomics, proteomics, metagenomics, and metabolomics paved us the way to discover more insights to determine the characteristics of the microbial community as well as its genomics and proteomics data, addressing the limitations of conventional techniques. It has also categorized microbial interactions based on their beneficiality at a very fast pace with great depths which was beyond imagination a few years back (Nadarajah et al., 2021).

This chapter discusses on the plant-microbe diversity based on niche, factors affecting these interactions (both biotic and abiotic), the mechanism behind the beneficial associations and pathogenesis, novel technologies used to analyse the microbial community and finally the applications and utilities of the plant-microbe interactions.

**2. Plant microbiome diversity**

Plant-microbe relationship is a very complex one and specific too. Plants secretions only signals to those microorganisms with whom the plants are suitable to make the bond and calls them to their immediate environment (Sharma et al., 2021). The soil microbes or the ones dwelling in the underground part colonizes the rhizosphere (the portion of soil regulated by roots). On the other hand, the microbes present aboveground colonizes the phyllosphere which includes microbes dwelling in seeds (spermosphere), flowers (anthosphere), fruits (carposphere), leaves (phylloplane), and stem (caulosphere) (Sivakumar et al., 2020; Hardoim et al., 2015; Nelson, 2018; Stanley and Fagan, 2002). However, there is existence of one more microbiome in addition to these two, endosphere which comprises the plant internal tissues (Sharma et al., 2021). Depending on niche, diversity among the plant-associated microbes can be seen (Trivedi et al., 2020). In general, these associations can takes place in any one of the following ways – pathogenic, parasitic or mutualistic (Newton et al., 2010).



***Fig 1.*** *Different plant-microbe interactions based on niche*

**2.1. Underground plant microbiome**

A substantial element of the underground plant microbiome is the rhizosphere. Several microorganisms form a community in the rhizosphere in conjunction with plant roots (Hinsinger et al., 2009). It is physiologically active as a result. According to general estimates, each gramme of root contains roughly 1011 cells from 30,000 different species (Berendsen et al. 2012; Pathma et al. 2019). Exudates from the roots, such as hormones, flavones and flavonoids, amino acids, phenolic substances, and organic acids, signal soil-dwelling microbes to begin the process of microbial colonisation. These secretions also affect how microbial genes are expressed (Compant et al., 2021; Egamberdieva et al., 2017; Patel et al., 2015; Nadarajah et al., 2021). Pseudomonads, Actinobacteria, Proteobacteria, Bacteroidetes, Copiotrophs, and Oligotrophs are a few of the prominent communities identified in the rhizosphere (Donn et al., 2015). Protozoa, archaea, oomycetes, fungi, algae, nematodes, viruses, and arthropods are among the other creatures found in the rhizosphere in addition to bacteria (Bonkowski et al., 2009). The beneficial ones interact with the host plants in symbiotic or synergistic ways, whereas those with parasitic and pathogenic effects have adverse effects (Haldar and Sengupta, 2015). When a community has developed in tandem with the root, the root exudate concentrates on the development of biofilms that are encased in extracellular polymeric materials (Mendes et al., 2013).

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| Plant | Dominant communities in rhizosphere | References |
| Maize | *Azospirillum*, *Ideonella*, *Bradyrhizoium* | Roesch et al., 2007 |
| Oats | Actinobacteria, Nitrospira, Firmicutes, Proteobacteria | DeAngelis et al., 2009 |
| Potato | Proteobacteria, Firmicutes, Acidobacteria, Actinobacteria, Bacteroidetes | Weinert et al., 2011 |

***Table 1.*** *Dominant microbial communities in rhizosphere with respect to the host plant*

Even the area found inside the roots, which makes up the endosphere microbiome, can be home to microbes (Hinsinger et al., 2009). An assortment of internally present bacterial and fungal endophytes colonise the plant roots. According to Sharma et al. (2022), the endophytes penetrate plant roots passively by infiltrating the cracks at the root emerging area, root tips, and lateral roots, as well as by an active process. Proteobacteria, Actinobacteria, Planctomycetes, Verrucomicrobia, and Acidobacteria make up the majority of the endophytic population. Among them, Proteobacteria forms dominant communities (Romero et al., 2014). The endosphere varies from the rhizosphere in that the host is involved in the identification and selection of microbial communities that maintain the plant's homeostasis (Compant et al., 2010). This maintains the concept of coevolution between plant and microbial symbionts. Numerous colonising endosphere bacteria have been revealed to exhibit chemotaxis pathways (Compant et al. 2010, Santoyo et al. 2016).

**2.2** **Aboveground plant microbiome**

The aboveground plant microbiome is held in the phyllosphere, representing the aerial plant parts (Parasuraman et al., 2019). According to estimates, the phyllosphere has a surface area of around 109 km2 and serves as a home for a variety of good and bad bacteria, with a density of 107 cells/cm2 of microbes on the foliage's surface (Lindow and Brandl, 2003; Farre-Armengol et al., 2016). Temperature, precipitation, light, pH variations, O2 concentration, availability of organic compounds, and other variables all have a large impact on the phyllosphere region (Sharma et al., 2022). It is densely populated with tiny creatures such as algae groups, bacterias like Bacteriodetes and Proteobacteria, actinomycetes, Ascomycota and Basidiomycota (fungi), viruses, and others (Sharma et al., 2022). Phyllosphere microorganisms are linked to plant development concerning biological processes like phytohormone production, biological nitrogen fixation, and defence against pathogen attacks (Cappelletti et al., 2016).

On the other hand, the aboveground endophytes which are dominant in the tissues play a variety of roles in biomass increment, stress tolerance as well as resistance, decreased intake of water, and many more by the development of symbiosis among the two (Panaccione et al., 2014; Rodriguez et al., 2009). Bacterial endophytes can be spread by several natural agents like wind, water, atmosphere, pollen, seed and insects (Frank et al., 2017). These endophytes receive their nutrition from several plant tissues like the xylem in fruits and flowers (Frank et al., 2017; Kandel et al., 2017). The location of endophytes throughout plant tissue will be heavily influenced by the food supply available within the organ to sustain their growth and development. Studies show that there exist discernible genus distinctions between endophytic and phyllosphere communities (Vishwakarma et al., 2020).

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| --- | --- | --- | --- |
| Plant | Dominant microbial community in phyllosphere | Dominant microbial community in endosphere | Referemces |
| Tomato | *Acinetobacter* | *Pseudomonas, Xanthomonas, Rhizobium, Methylobacterium, Sphingomonas* | Campisano 2014;Dong et al., 2019 |
| Grapes | *Bacillus, Frigoribacterium, Curtobacterium, Citrobacter Enterobacter, Acinetobacter, Erwinia, , Pantoea* | *Bacillus, Ralstonia, Propionibacterium Burkholderia, Dyella Pseudomonas, Staphylococcus* |

***Table 2.*** *Difference in dominant microbial communities in phyllosphere and endosphere*

**3. Factors influencing plant-microbe interaction**

**3.1. Host species**

According to Sharma et al. (2021), plant species actively influence the microbiome's microbial composition. The organisation of the root-associated microbiome is more variable in plants that are phylogenetically distantly connected, exhibiting difference in phenotypic traits like root architecture, chemical secretions. The host species also influences the spatial and temporal distribution of a microbial community. Dependency on plant species was demonstrated in an experiment in which a grapevine and four species of weeds (*Stellaria media, Lepidium draba, Veronica arvensis, Lamium amplexicaule*) grown side by side in the same vineyard had substantially distinct microbial community compositions (Samad et al., 2017).

**3.2. Host genotype**

Another important factor influencing interactions is the host plant's genotypic makeup, which determines phenotypic characteristics such as root hair density, nature of root exudates and rhizodeposits, root hair length, and leaf morphological features such as stomata, veins, and many others (Lindow and Brandl, 2003; Bulgarelli et al., 2012; Lundberg et al., 2012). Specific microbial species may be selectively drawn to or kept out by particular genotypes. According to Fitzpatrick et al. (2018), in flowering plants, rhizospheric microbial populations varied less as root hair length increased compared to root hair density, which increased greater variance among endophytic bacteria.

**3.3. Plant-derived metabolites**

Plant exudates nourish rhizosphere bacteria and control the variety of microorganisms (Olanrewaju et al., 2019). The effectiveness of colonisation depends on the presence of root exudates in the rhizosphere, such as amino acids, organic acids, carbohydrates, and phenolic compounds (Sundaram et al., 2015). Terpenoids, tannins, alkaloid compounds, and flavonoids are only a few examples of the secondary metabolites that the phyllosphere produces above ground and which the microorganisms use as a carbon source. Additionally, it generates volatile substances like methanol, which provides energy to the methylotrophic epiphytic bacteria and yeast as well as mediates cell wall metabolism in microorganisms (Crozier et al., 2006; Galbally and Kirstine, 2002; Vorholt, 2012).

**3.4. Host immunity**

The plant's health and immunology, as indicated by its two-layered defence system, can influence microbiome composition (Sharma et al., 2021). The triggering of immunological responses and the activation of plant defence mechanisms, such as the generation of antimicrobial substances, can prevent the establishment and expansion of potentially harmful microorganisms. On the other side, certain advantageous bacteria have developed strategies to get around or decrease plant defence, enabling them to form positive relationships with plants.

**3.5. Host developmental stages**

Interactions among the microbial communities and plants are associated with the host's age and stage of development (Sharma et al., 2021). Although the chemistry of root exudates is determined by plant genotype, plant age affects their concentration, amount, and quality. Rhizodeposit secretion decline as the plant ages (Bulgarelli et al., 2013). Furthermore, the non-pathogenic endophytes are unable to mitigate the plant defence systems as it becomes more efficient as the plant ages. As a result, juvenile plants have more endophytes than older ones.

**3.6. Microbe-microbe interaction**

Apart from interactions among plants and microbes, microbe-microbe interactions are also an important factor which influences the plant microbiome (Sharma et al., 2021). The dominance of a particular species in a niche depends on the competetion that occurs among themselves for food, space, production of effectors and secondary metabolites, etc (Pathma et al., 2020). Competition, parasitism, or mutually beneficial behaviour among the bacteria are possible interactions. Keystone species are developing and revealing insight into how they coexist with other microbial species and may have regulatory effects on their environment and other microbiome members (e.g., Rhizobiales and Burkholderiales in the rhizosphere) (van der Heijden and Hartmann, 2016).

**3.7. Anthropogenic factor**

Crop domestication is an example of an anthropogenic activity that seeks to enhance the qualitative and quantitative factors of output relative to their wild counterparts through continuous selection for a desired characteristic. However, the genetic variety and environmental adaptation of the domesticated crops were lost in this process. The need for artificial fertilisers and plant protection chemicals has grown as a result, which has contaminated the environment and prevented many interaction which are beneficial to the plant (Pathma et al., 2020). In contrast to the native fields having wild plant species, domesticated agriculture fields have distinct bacterial populations. Continuous nitrogen fertiliser application has been cited as having inhibited the evolution of mutualistic rhizobium bacterial strains (Weese et al., 2015). So, we can say that anthropogenic activities tend to act against the natural forces and impacts the co-evolution of plants and microbes (Pathma et al., 2020).

**3.8. Edaphic factors**

The native microbial community structure is determined by the nature of the soil and physical properties, including texture, structure, colour, water-holding capability, pH levels, availability of nutrients,  organic matter content, etc., as well as by geographical locations like plains or hills (Islam et al., 2020; Pathma et al., 2020). As an illustration, dark brown soil promotes the diversity of fungal communities whereas black soil supports the diversity of rhizobacteria (Xu et al., 2009; Wang et al., 2009). In contrast to fertiliser amendments, soil particle size initially dictates the corresponding microbial taxa specificity (Sessitsch et al., 2001). According to pH, acidic soils have a low variety of microbiomes, while neutral soils maintain a great diversity (Fierer and Jackson, 2006). According to studies by Davidson and Robson (1986) and Zhang et al. (2009), plants are seen to release more flavones and flavonols in nitrogen-deficient soils, which in turn triggers the rhizobia-legume symbiotic associations in the rhizosphere. The variety and biomass of microbial communities are favourably influenced, strengthened, and stabilised by conservation agriculture, which advocates minimal tillage and uses organic manuring (Wang et al., 2017).

**3.9. Other environmental factors**

According to Gupta et al. (2002), environmental changes cause variations in plant physiology, which in turn produce unique microbiomes. Additionally, a phenotypic characteristic is a result of both the plant's genotype and its environment. As a result, a plant's behaviour deviates from its usual state in a number of ways, attracting various strains of microorganisms, suitable in that environment. Limited soil moisture caused plants to produce more amino acids, which in turn impacted the rhizosphere's microbiology and decreased the growth of mycorrhizal mycelium in plant roots (Katznelson et al., 1955). While sufficient precipitation boosts the activity of soil microorganisms, increasing the carbon content of microbial biomass. Microorganisms with modified respiration may grow more quickly at a higher temperature (Sharma et al., 2022). Because of this, microbial diversity is more favourable in the tropics than in temperate regions, especially in the phyllosphere (Vorholt, 2012).

According to Bardgett (2008) and Compant et al. (2010), the altered root exudate frequently affects the interactions between plants and microbes. Radiation is a significant consideration in this scenario. In comparison to rhizospheric communities, the phyllosphere colonies are more susceptible to increasing radiation (UV) (Sharma et al., 2022). Abiotic variables affect the diversity of the plant microbiome in this manner.

**4. Mechanism of plant-microbe interactions**

Initially classifying all microbial infections as destructive incursions, the immune system of plants later distinguishes between pathogenic and helpful microorganisms (Pel and Pieterse 2013). Pattern recognition receptors (PRRs) found in plant plasma membranes during microbial infection detect and differentiate between MAMPs (microbe-associated molecular patterns) and PAMPs (pathogen-associated molecular patterns). Accordingly, the PRR triggers either MTI (MAMP-triggered immunity) or, PTI (PAMP-triggered immunity), which prevents infection in cases of phytopathogens but does not prevent beneficial infections. These defence responses are a result of continuous dialogues between plants and microbes, facilitated by several phytohormones. This signalling by phytohormones is crucial for the defensive reactions induced by both good and bad microorganisms. Pathogens that overcome the plant's initial line of defence produce effector molecules that interfere with the plant's defence system. Plants have developed a more noticeable defence mechanism known as ETI (effector-triggered immunity) against those effectors. Plant resistance proteins, also known as R proteins, recognise and act against the effector molecules generated by pathogens during this second stage of defence. Prior to the pathogen entering the plant, PTI offers protection, whereas R proteins step in to help once the infection has started (Pathma et al., 2020). This plant-microbe interaction mechanism, which Jones and Dangl (2006) suggested, is commonly referred to as the "zigzag" model.

**5. Modern tools to analyse microbial community**

With the development of molecular tools, it is now possible to research microbial communities connected to plant microbiomes with novel information (Pathma et al., 2020). Only less than 1% of microorganisms could be studied using culture-dependent approaches (Staley and Konopka, 1985; Pathma et al., 2020). The recent development of a variety of culture-independent methodologies yields valuable information that is now being used to evaluate the hidden microbial diversity and functional characteristics of the same (Gupta et al., 2021). Table 3 lists some molecular biology-based approaches that are culture-independent, in contrast to the conventional methods. These techniques can be used extensively used in varied microbial diversity investigations without affecting ecosystem structure.

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| SL.No. | Techniques | Reference |
| 1 | Enhancer traps (using fluorescent protein) | Rediers et al., 2005 |
| 2 | Ultradeep sequencing | Velicer et al., 2006 |
| 3 | 454 pyrosequencing | Leveau, 2007; Erkel et al., 2006; Handelsman, 2004 |
| 4 | Analysis of eDNA |
| 5 | Metagenomics |
| 6 | 16s ribosomal RNA based microarray  | Ehrenreich, 2006; Sanguin et al., 2006 |
| 7 | Analysis of transcriptome | Mark et al., 2005; Yuan et al., 2008 |
| 8 | Isotope probes usage | Haichar et al., 2008 |
| 9 | Flow cytometry for in-situ antifungal gene expression | De Werra et al., 2008 |
| 10 | Real-time PCR (RT-PCR) | Wu et al., 2009 |
| 11 | Chromatography techniques |
| 12 | Fourier transform infrared spectroscopy (FTIR) |
| 13 | Nuclear magnetic resonance (NMR) |
| 14 | Signature tagged mutagenesis (STM) | Walder et al., 2017 |
| 15 | Differential fluorescence induction (DFI) |
| 16 | Single-molecule real-time (SMRT) sequencing |
| 17 | Fatty acid methyl esters (FAME)  | Gupta et al., 2018; Bodor et al., 2020 |
| 18 | Next generation sequencing (NGS) |

***Table 3.*** *A list of techniques used as a tool to study microbial communities*

The above-mentioned techniques have enormous possibilities for studying the microbiomes associated with a plant, but they have some limitations, such as a lack of population dynamics data, functional data, difficulties with data analysis due to the high frequency of interspecific and intraspecific differences, and a lack of assembled microbial species genomes (Wolfe, 2018).

**6. Impact and role in plant health and agroecosystem**

Plant microbiomes influence entire plant health and ecological fitness by supporting plant development and avoiding abiotic and biotic stress. Plant microorganisms' effects may be generally classified into numerous roles.

**6.1. Recycling of nutrients**

Microorganisms assume a pivotal role in the continuous recycling of numerous essential nutrients, including C, N, P, K, Zn, Ca, Mn and Si. Among these, nitrogen-fixing bacteria play a vital role by converting atmospheric nitrogen into a bioavailable form for plants, thereby facilitating the nitrogen cycle within ecosystems. In aquatic environments, the symbiotic interaction between bacteria and algae significantly influences the carbon and nutrient cycles. Bacteria, acting as decomposers, contribute to the breakdown of organic matter, while algae, through photosynthesis, not only harness energy from sunlight but also release oxygen into the ecosystem. Consequently, the identification and preservation of optimal densities and community compositions of these crucial microorganisms become of paramount importance for the efficient functioning of each respective nutrient cycle. Thus, meticulous management and maintenance of the microorganism population in these cycles are imperative for sustaining ecological balance and the overall health of ecosystems (Jacoby et al., 2017; Rashid et al., 2019).

**6.2. Production of PGR**

Plant growth-promoting chemicals such as auxin (IAA), gibberellins, cytokinins, and ACC deaminase are produced by the plant-related microbiome (Penrose and Glick, 2002; Glick, 1995). Auxins are known to be created by bacteria such as *Pseudomonas* and *Bacillus* via metabolic pathways, whereas cytokinins are produced by microorganisms such as *Agrobacterium* and some fungi. Cytokinins encourage cell division and have an impact on shoot development, leaf expansion, and nutrient absorption. Certain microbes, including bacteria like *Pseudomonas* and fungi such as *Penicillium*, can generate ethylene, a plant hormone involved in fruit ripening and senescence. Some microbes create elicitors, which cause plants to produce increased quantities of plant growth regulators and help in stress tolerance.

**6.3. Role in plant protection**

Microorganisms play an important role in plant protection by combating pests and diseases. Antimicrobial substances produced by beneficial bacteria include antibiotics, HCN, siderophores, and enzymes such as pectinase, chitinase, lipase, DNAase and others (hydrolytic in nature), which defend the host plant from invading phytopathogens and herbivores (Pathma et al., 2020). *Bacillus thuringiensis*, for example, produces toxins that are fatal to insects, making them efficient natural pesticides. Plant pathogens can be suppressed by fungi, such as *Trichoderma species*, colonising their environments, competing for resources, and creating antifungal chemicals. A varied microbial population can provide numerous ways of defence while also improving overall plant resilience.

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| Microbe | Action | Reference |
| *Bacillus* and *Pseudomonas*  | Inhibits the action of *Agrobacterium tumefaciens, Fusarium spp.* | Pathma et al., 2019; Ho et al., 2017 |
| *P. fluorescens* WCS417r | Induces systemic resistance in Carnation against *F. oxysporum* | Van Peer et al., 1991 |
| *Bacillus pumilus* INR-7 | Protects against cotton aphids | Stout et al., 2002 |
| PGPR strains | Protects against blue-green aphids, termites and green peach aphids | Kempster et al., 2002; Sindhu et al., 2011; Boughton et al., 2006 |

***Table 5.*** *Examples of some microbes used in plant protection*

**6.4. Role as biofertilizers**

The symbiotic feature of plant-microbe interaction has proven to be an effective biofertilizer in boosting plant development. In symbiosis, bacteria use inert nitrogen from the surrounding environment to transform it into a usable form (ammonium and nitrate) for the plants while obtaining carbon from the appropriate host plant. Azorhizobium, Allorhizobium, Sinorhizobium, Bradyrhizobium, Mesorhizobium, and Rhizobium are the most effective bacterial strain genera (Singh et al., 2019).

**6.5. Bioremediation/ Rhizoremediation**

Bioremediation is an economic process that uses bacteria, algae, fungi, or plants to remove heavy metal ions from polluted environments by adsorption, biosorption, segregation of heavy metals into intracellular molecules, vacuolar compartmentalization, metal binding, extracellular mobilisation, or metal immobilisation (Rashid et al., 2019; Ray et al., 2020). Rhizoremediation, on the other hand, is a process of pollutant breakdown by a plant's rhizomicrobial population in a polluted location (Hao et al., 2014). Some examples are cited in table 4.

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| --- | --- | --- |
| Bacterial strain | Action | References |
| Enterobacter asburiae | Ameliorate cadmium toxicity | Kavita et al., 2008 |
| Pseudomonas putida strain PCL1444 | Reduce the concentration of naphthalene around the roots | Kuiper et al., 2001 |

***Table 4.*** *Example of Rhizoremediation*

**6.6. Biocontrol agents**

The biocontrol agent effectively inhibits the growth and virulence potential of pathogenic organisms through various mechanisms including niche exclusion, nutrient competition, production of cell wall-degrading enzymes such as chitinase, synthesis of toxic secondary metabolites, and induction of induced systemic resistance in the host plant. Niche exclusion refers to the ability of the biocontrol agent to occupy and utilize specific ecological niches, thereby depriving pathogenic organisms of essential resources required for their growth and survival. By competing for nutrients, the biocontrol agent limits the availability of vital substances required by pathogens, thus impeding their ability to proliferate and cause damage. Additionally, the production of cell wall-degrading enzymes, such as chitinase, by the biocontrol agent enables it to degrade the structural integrity of pathogenic organisms' cell walls, leading to their lysis and subsequent inhibition of growth. Furthermore, the biocontrol agent synthesizes toxic secondary metabolites, which can directly or indirectly interfere with the metabolic processes of pathogens, further impeding their growth and virulence. Moreover, the biocontrol agent exhibits a remarkable ability to induce systemic resistance in the host plant. This induction triggers the plant's defense mechanisms, such as the production of antimicrobial compounds, reinforcement of physical barriers, and activation of signaling pathways, thus enhancing the plant's resistance against pathogenic attacks (Singh et al., 2019). Biocontrol agents can be used in integrated pest management (IPM) strategies, which combine multiple approaches to minimize chemical pesticide use and reduce environmental impact.

**7. Future opportunities and conclusion**

Plant-microbe interactions have a big impact on ecosystem processes such as biochemical cycling, the formation of microbial communities, and plant development. Understanding the fundamentals of the interactions between plants and microbes might help improve plant health, disease management, and risk management by revealing important biological phenomena. Even today, we still don't fully understand how these interactions work, and more research is required to pinpoint the sensors and signalling pathways involved, comprehend the molecular underpinnings of how various types of stresses and responses interact, and identify the crucial elements of such interactions. Additionally, using complete organisms, such as bacteria and fungi, to enhance plants has become a way of the past; modern techniques use molecular & sub-molecular microbial units (Bourras et al., 2015). In order to understand the disease development processes & suppression, growth and development of plants in association with microbes, immune response, nutrient cycling and absorption, and other processes, microbial ecology and molecular plant pathology must be combined with next-generation sequencing technology, multiple 'omics' tools, databases, and metabolic modelling, artificial intelligence and machine learning (Iman et al., 2016; Nadarajah et al., 2021). Given the increased attention being paid to GMOs, it is important to remember that genetically modified crops can have a significant negative influence on the native microbiomes of the plant, causing unexpected changes in the variety of related bacteria. Therefore, while developing higher-performing hybrids or GM crops, it is equally necessary to take into account the plant-associated microbiomes (Pathma et al., 2020). In conclusion, despite all odds and difficulties, we should pursue a comprehensive strategy to utilise the microbial communities related to plants in an eco-friendly and sustainable way.

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