





International Journal of Recent Advances in Multidisciplinary Research Vol. 02, Issue 08, pp.0677-0684, August, 2015

Review Article

ANTIBIOFOULING BIOMATERIALS

*Asifa Qureshi, Smita Pal, Saheli Ghosh, Atya Kapley and Hemant J Purohit

Environmental Genomics Division, CSIR-National Environmental Engineering Research Institute (NEERI), Nagpur Nehru Marg, Nagpur-440020, India

ARTICLE INFO

Article History: Received 12th May 2015 Received in revised form 30th June, 2015 Accepted 21st July, 2015 Published online 31st August, 2015

Keywords:

Antifouling, Biofilms, Biomaterials

INTRODUCTION

Biofilms and biofouling originated on earth nearly 3.25 billion vears ago in oligotrophic systems (Wimpenny et al., 2000) where roughness of the surface provides more suitable environment for their growth. Microbes form biofilms in response to various factors, such as recognition to specific or non-specific attachment sites on a surface, nutritional factors, detergents and sub-inhibitory concentration of antibiotics (Hoffman et al., 2005). Microbial biofilms leading to biofouling consists of organisms and their by-products. It occurs on any surface by colonization and accumulation of micro and macrofoulers on immersed structures. Adhesive property, biofilm formation as well as quorum sensing associated features like exopolysaccharide secretion, virulence factor all these together lead to biofouling. Majority of studies have been carried on pathogen treatment or in medical fields. However, literature about biofilm and quorum talking of microorganism for environmental isolates are still in infancy. Biofouling study appears as unsaturated as well as interesting ground to be exploited by environmental scientists. For example quorum sensing in fungi was first reported in 2001 and studies were generally encircled till now with pathogenic strains of Candida sp (Burke et al., 2007).

*Corresponding author: Asifa Qureshi,

Environmental Genomics Division, CSIR-National Environmental Engineering Research Institute (NEERI), Nagpur Nehru Marg, Nagpur-440020, India.

ABSTRACT

Antifouling refers to the process of control of fouling which occurs on liquid-solid surfaces. The term 'fouling' indicates an undesirable natural succession process during biofilm formation, in which a submerged surface or membranes becomes encrusted with material from the surrounding environment. It mainly involves microorganisms and their by-products developed on the surface by conditioning, attachment, biofilm formation followed by colonization. The accumulation of micro and macrofoulers on immersed structures results in economic as well as environmental losses. It is one of the major vulnerable problems currently disturbing many ecological niches as well as in shipping and other industrial aquatic processes. The existence of natural antifouling agents or biomaterials provides sustainable eco-friendly control and hence remains a challenge for future researchers. The use of biological tools for control of fouling is gaining importance day by day.

Biofilm formation is key step towards biofouling process. The biofilm concept was coined by Bill Costerton in 1978 and now it is widely embraced by microbiologists, engineers and computer scientists (Jenkinson and Lappin-Scott 2001). It is a microbial development concept that was first proposed by O'Toole and his colleagues (O'Toole et al., 2000). According to Martin Dworkin, 'development' refers to a series of stable and metastable changes occurring in a cell, in response to certain environmental stimuli that become a part of the normal life cycle of cells. It helps the cells to adapt and survive in their dynamic environment (Dworkin and Kaiser 1985). Initially microbiologists ignored the socio-biology concept but the studies on co-operative behaviour in Myxobacteria and quorum sensing/biofilm formation in Pseudomonas aeruginosa sparked biofilm research (Kalia and Purohit 2011). They are found nearly on every surface and interfaces exposed to oil, water or air (Halan et al., 2012). They can colonize on both biotic and abiotic surfaces such as industrial and hospital settings (Stoodley et al., 2004, Lear and Lewis 2012) and on human host (Jefferson 2004).

Why microbes form biofilm?

According to the Darwin's theory of evolution, the only true driving force behind the course of action of any organism is reproductive fitness that is any action that increases proliferation. Since it remains an inherent action it almost seems contradictory that a biofilm mode of growth would impart reproductive fitness over planktonic mode. Outside the laboratory, bacteria would rarely find themselves in an environment as rich as the culture media and hence the biofilm offers a more protective mode of bacterial growth in nature. (Jefferson *et al.*, 2004). The biofilm matrix provides its members resistance to many environmental stresses such as fluctuations in pH, temperature, osmolarity, UV damage (Elasri and Miller 1999), desiccation (Chang *et al.*, 2007), predation (Matz 2005) and specific secondary metabolites such as antibiotics (Stewart and Costerton 2001), an advantage not available to their planktonic counterparts.

Microbial Biofilm Development in Environment

Biofilm formation in the environmental biofilms begins with a transition of bacteria from the planktonic (free swimming) to its genetically distinct attached state (Singh *et al.*, 2006, Parsek and Tolker-Nielsen 2008, Stoodley *et al.*, 2004). The physiological and genetical transition occurs across the life cycle of the biofilm as shown in Fig. 1.

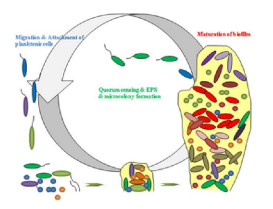


Fig.1 Biofilm developmental stages 1. Quorum sensing, EPS and microcolony formation 2. Maturation of biofilms 3. Migration and reattachment of planktonic cells

- Quorum sensing, EPS and microcolony formation
- Maturation of biofilms
- Migration and reattachment of planktonic cells

Quorum sensing, EPS and microcolony formation

During biofilm formation, cooperative communication known as quorum sensing — occurs to coordinate bacteria during different stages of development. By releasing some intercellular signalling molecules bacteria can sense each other presence and they continue to secrete these molecules until a sufficient population is reached thereby forming a 'quorum'. In this way bacteria can sense their quorum and start forming micro colony (Fig 1). This is called quorum sensing in bacteria. What is the relation between quorum sensing and biofilm formation? There is an acute relation between quorum sensing and biofilm formation because biofilms generally consists of clusters of cells. One can predict that the aggregates might be a product of quorum sensing. The prediction has truly provided evidence that quorum sensing is important for biofilm formation and dispersal. Three most common pathways used by bacteria are Acyl Homoserine Lactone (AHL) system in Gram positives, peptide based signalling in Gram negatives (Waters and Bassler 2002, Fuqua et al., 2001) and AI-2 signalling in both Gram positives and Gram negatives. In acyl homoserine lactone signalling system, a single enzyme synthesize the signal i.e acyl homoserine lactone from cellular metabolites for communication (Parsek et al., 1999). It can diffuse across the cell membrane, interacts with a cytoplasmic DNA binding receptor protein which belongs to Lux-R family of genes which then activates the expression of quorum sensing genes. These AHLs are derived from S-adenosylmethionine which consists of a hydrophilic homoserine lactone head and hydrophilic acyl side chain that varies from species to species. The side chain consists of 4 to 18 carbon atoms, the variation occurs in length from the 3rd carbon. These alterations are the main source of specificity in QS signals and facilitates cell to cell communication in bacteria.

The chemical structures of Quorum sensing peptides in Gram positive bacteria also varies in the number and type of amino acid residues (Dusane et al., 2010) and controls diverse physiological processes. Their biosynthesis processes are more complex than Gram negative bacteria on account of their post translational modifications in the peptides and their inability to diffuse through the membranes. Largest quorum sensing peptides in Gram positive bacteria are antibiotics which are having antimicrobial activity such as nicin produced by Lactobactococcus lactis (Lubelski.et.al.2008). Some other autoinducing peptides such as Type-1 autoinducing peptide produced by Staphylococcus aureus, Enterobacter faecalis, Listeria monocytogenes plays important role in quorum sensing process (Miller & Bassler 2001, Water and Bassler 2005). Some other non AHLs such as indole, small RNAs and secondary messengers are also involved in quorum sensing. Indole produces quorum in Escherichia coli. (Wang et al.,2001). Quorum sensing process also depends on the nutritional conditions (Shrout et al., 2006).

Many groups have demonstrated the link between quorum sensing and biofilm formation. In some species link was found while in some no relation was found between the two. Species in which quorum sensing mediates biofilm formation are through different stages initiated by attachment. Attachment or adherence of a bacteria to a surface or substratum is the initial step in biofilm formation (O'Toole.2000). 'Attachment' in deeper sense means bacteria forms bonds with the surface by their adherence factors. Bacteria employ many adherence factors for this purpose such as pili, fimbriae, carbohydrate binding proteins etc. Intestinal pathogen Helicobacter pylori has luxX genes homologues involved in attachment (Kirisits and Parsek 2005). Firstly when a bacteria adheres, it forms reversible attachment that is it can also come out of the surface if loosely attached then irreversible attachment occurs that is strongly bound to the surface and cannot come out. Then attached bacteria secrete extracellular polymers (EPS) and begins to multiply by microcolony formation and thus biofilm matures. By further growth and secretion of EPS, 3D structure in between the cell cluster channels are also formed to deliver water, nutrients and waste removal. After irreversible attachment bacteria begin to secrete EPS and through quorum sensing bacteria mutiply themselves until a specific cell density is reached and forms microcolony.

Further EPS secretion occurs and the biofilm matures by forming different shapes such as 3D mushroom like or flat microbial mats (O'Toole 2000). EPS or extracellular polymeric substances are the building blocks of the biofilm community. EPS serves variety of functions to support the biofilm mode of life by enabling cells to come in close proximty, facilitating cell to cell communication, store various extracellular enzymes which sequesters and digest colloidal, dissolved and solid substances leading to enhanced metabolism and growth. It also acts as a recycle centre which keeps and allows components of lysed cells to be used by live cells, which may also results in horizontal gene transfer by transformation, conjugation as well as transduction i.e involving viruses too. Although EPS represents external digestive system but complete digestion requires a large variety of enzymes on account of heterogeneity. It is also called 'dark matter of biofilms' because of heterogeneity of polymers and difficulty in analysing them (Flemming et al., 2002, Chelsea and Brennan 2010, Carsten Matz et al., 2011). This heterogeneity also varies from biofilms to biofilms depending upon the type of microbes, mechanical shear, temperature and availability of nutrients, Bacterial extracellular structures also stabilizes the matrix (Zogag et al., 2001). The forces that stabilizes EPS are not covalent bonds but weak interactions such as hydrogen bonds, Vanderwaals interactions, electrostatic and ionic interactions, entanglement of long molecules. They behave like elastic bodies until a threshold pressure is reached. Beyond that threshold pressure it liquefies to highly viscous liquids (Korstgens et al., 2001). During viscoelastic phase same partners react causing breakage and formation of bonds. After that equilibrium shifts in the breakage of bonds and EPS liquefies. This is the point which exceeded when we get slipped while walking in the streams containing biofilm coated rocks. The porous architecture of EPS allows convection of flow of fluid through the depth of the biofilm. Within EPS, substances flows by diffusion. So at the bottom, organisms get excess nutrients while those at top of the matrix competes among themselves to get nutrients from the bulk water phase.

Oxygen gradient also get formed in biofilms by a actively respiring heterotrophic organisms which consumes oxygen before it diffuses through the matrix thereby creating an anaerobic environment below the aerobic organisms allowing the growth of anaerobic organisms (Flemming 2010). Other gradients such as pH, redox and ionic gradients are also formed (Stewart 2001). Cells in a biofilm remain surrounded by EPS or extracellular polymeric substances and those cells forms capsules, are associated more closely to the surfaces than others (Flemming 2010). Multispecies biofilm community structure depends upon production and quantity of EPS (Sutherland 2001), concentration, cohesion, charge, sorption capacity, specificity, nature of components of EPS. Pores and channels determine mode of life in biofilms. CLSM examination revealed that EPS matrix provides physical structure which segregates different organisms in the biofilm community. These segregated regions contain different biochemical environments that are enzymatically modified in response to dynamic environment (Sutherland 2001).

Biofilm architecture depends upon nature and amount of EPS produced. EPS of *Escherichia coli* and colanic acid of *Bacillus subtilis* are essential for the formation of 3D structure.

Alginate is required for biofilm development but it is not essential in Pseudomonas aeruginosa (Flemming 2007). Acetyl groups also affect biofilm structures by modifying alginate with acetyl groups which increases cohesive and adhesive properties of EPS (Flemming 2007). Since the matrix is negatively charged and if it encounters multivalent cations it alters the structure. For example Ca2+ forms a bridge of polyanionic alginate molecules resulting in a thick and compact structure with increasing mechanical stability (Kortgens et al., 2001). EPS also possess optical properties. EPS have slightly different refractive index than water hence light tend to enter. Actually it enters the biofilm rather that reflect on at the surface. This is called 'forward scatters'. EPS function as a light conductor. EPS gel allows 'recapture'of scattered photons from an underlying surface and increases the absorption potential of the underlying cells by cellular chromophores (Decho 2010). EPS are formed by nucleic acids, polysaccharides, lipids, water proteins, ketal-linked pyruvates, detritus etc. Each of the components have indispensable roles which cannot be ignored (Decho 2010).

Maturation of biofilms

Maturation of the biofilm involves growing of microcolonies and production of extracellular polymeric substances to form a spatial structure which stabilizes the biofilm community (Kim et al., 2008, Marcato et al., 2012)). Building of this spatial structure is determined by three distinct layers of organisms. The inner layer is inhabited by early or pioneer colonizers which attaches to the surface which are generally facultative anaerobes. They serve as a foundation of the biofilm structure by remaining attached to the surface. In a developing biofilm strong oxygen gradients are formed by actively respiring aerobes at the upper stratas whose oxygen consumption rate is faster than oxygen diffusion rate. An anaerobic environment gets created at inner layer due to little or no oxygen diffusion (Stewart & Franklin 2008). The basis of genetic inheritance for the biofilm community remains in the inner layer. The middle layer consists of organisms which are arranged in close proximity to each other, which allows them to exchange nutrients and genetic information (e.g., for antimicrobial resistance). The process of cell-to-cell communication and genetic interaction between the cells occurs in middle layer of the biofilms, which allows the members to coaggregate with each other (Otami et al., 2007; Otzen et al., 2007). Reports suggests that in domestic showerheads (Vornhagen et al., 2013), biofilms of drinking water distribution systems (Simoes et al, 2008) and dental plaques (Kolenbrander et al1989), the middle bacteria acts as an adaptor or bridge which connects the inner and outer most layers of bacteria in a biofilm. The outer layer comprises of actively respiring bacteria which behave in a manner similar to individual planktonic (non-biofilmassociated) bacteria (Fig 1). Planktonic bacteria migrates and reattaches to new surface for proliferation.

Migration and reattachment of planktonic cells

It is very significant stage in the biofilm life cycle which allows the cells to inhabit new surfaces (Flemming, 2011). It is influenced by nutrient starvation and secretion of EPS hydrolyzing enzymes like hexosaminidase which breaks off EPS to release planktonic cells in the fluid-surface interface (Kaplan *et al*, 2003).

The dispersal is also affected by shear forces in its surroundings. Specifically, shear forces present in the microenvironment are high enough to cause detachment of a portion of the biofilm and formation of projections called streamers that migrates to new locations and environments. Consequently, more support are required for reattachment of planktonic cells to carry out their surface associated stages of life cycle (Leck 2005; Russel *et al.*, 2009). The association of bacterial layers in biofilms leads to biofouling at later stages.

Problems regarding biofouling

As learnt from literature survey, biofouling creates problems on any liquid-solid surfaces, for example on ship hulls. Roughness created by biofouling by bacteria results in high frictional resistance which leads to increasing weight and subsequent speed reduction of liquids with high power consumption. Relatively light biofouling that is made up diatom slimes results in increased power backing of 10-16%, whereas heavy calcareous fouling at full cruising speed results up to 86%. In the case of fuel consumption the loss rises can be up to 40% (Schultz 2007) involvement of higher efficiency machinery to overcome this problem leads to voyage overall costs as much as 77% higher. Biofouling debris clean up entails huge man power, machineries and high chances of time loss and wastage of resources. Toxic waste products and "alien species" get introduced to native ecosystems. One report showed introduction of new 16 species of barnacles at the port in Osaka Bay, Japan due to Biofouling (Oumi et al., 2007, Chelsea and Brennan 2012).

In many industries like water treatment, food processing, paper and milk industries biofouling have been found to be very serious problem regarding maintenance of different types of membranes (Anand *et al.*, 2014). It have been still a major challenge in terms of quality of water, plant performance and operating cost in different industries (Fig 2). Four major types of fouling occurs in membranes/filters viz;



Fig. 2. Biofouling (a) management in ship yard (b) membrane sheets in industry (Photocourtesy Yebra etal 2004, Chiellini et al 2012)

- inorganic salt precipitation (contributed by sparingly soluble salts),
- organic (mostly natural organic matter or effluent organic matter),
- colloidal (caused by accumulation of a colloidal cake layer on the membrane surface), and
- microbiological (usually governed by bacterial biofilms and subsequent microfouling formation).

If fouling could not be controlled, it could results in permeate flux decline of the membrane because of the accumulation of retained biofilms on it, which leads to increased differential pressure and feed pressure, increased salt passage, increased energy consumption. Other vigorous problem include membrane biodegradation caused by acidic by-products.

Antifouling strategies and necessity of biotools for antifouling

To control biofouling the proposed methods that concerned about physical, mechanical or chemical means were being a matter of question day by day. Physical or mechanical cleaning of ship hulls or submerged structure basically exposes the substrata for next event of biofouling in successive days. The chemical means of control actually employed different types of biocide or implication of antibiofouling paints (Yebra et al., 2004). Most antifouling paints composed of organotin (tributyltin) or heavy metals (copper plus organic booster biocides, zinc) that, even in very low concentration, served as broad spectrum toxins to target as well as non-target organisms. Use of toxic tributyltin (TBT) coatings has been increasingly banned at global scale (Magin et al., 2010). It has been shown that membrane biofouling chemical clean up stress-up the residual biota and triggers for readily biofilm formation for next session. The fouling organisms generally showed robust nature, that even if 99.9% cells were removed then even the chances exist that films could be easily formed from remaining biostratum (Nguyen et al., 2012). Even frequent chemical cleaning of membrane actually shortens the life time of membrane and it also includes extra maintenance cost as well as extra man power (Chiellinia et al., 2012). So to have alternative, safe, eco-friendly control use of biological tools are gaining importance day by day in biofouling treatment.

Antifouling biotools

Bioinspired biomaterial

Learning from nature's own defense and transferring the knowledge into application to combat biofouling biomaterials were gaining importance day by day. It has been seen that most of study regarding bioinspired biomaterials were used in the aspect of marine biofouling or ship hull management field. For example, many reports suggested sharkskin mimicking, resulted reduction in drag force and Reynolds number and deterred biofouling. To make nature's perfect replica there was urged to incorporate biosciences into physical models. Instead of only mimicking the surface topology and texture, many studies were carried on to add bioinspired biomaterials in the coating so that both the physical topography and biochemical phenomenon could be exploited (Salta et al., 2010). For example, besides having groovy scale topography (Baum et al., 2002) whale was also reported to resist micro-organisms as it contains micropores and nano-ridges surrounded by enzymatic gel coating that disintegrates proteins and carbohydrates. Sessile marine organisms do not possess mechanical or dynamic facilities to combat fouling, but they were notably free from macrofouling. Studies revealed that they produce some antimicrobial and antibiofilm exudates and these secondary metabolites keep them free from any fouling condition.

One of the very good examples has been sea weeds. Their unusual chemical structured secondary metabolites were very exclusive and do not share common features with terrestrial tissues. 40% less biofouling was observed when sugar kelp (Saccharina latissima) and Guiry's wrack (Fucus guiryi) were used as bioinspiration and matrix was replicated by polymorphic reproduction with doping of bromofuranone (Chapman *et al.*, 2014).

Quorum Sensing in biofilms are regulated by releasing and detecting small signaling molecules known as Auto-Inducers (AIs). Three types of AIs have been reported including oligopeptides, N-Acyl Homoserine Lactones (AHL), and autoinducer-2 (AI-2). Cellular communications in Grampositive and Gram-negative bacteria are achieved by oligopeptides and AHL, respectively. In the case of interspecies communication for both Gram-positive and negative bacteria AI-2 molecules are implied.

Table 1. List of some natural re	esources producing	antifouling	biomolecules

Source	Bio molecules	Anti biofouling property	
Macroalga (Delisea pulchra)	Furanone/ 2(5H)-Furanone, (5Z)-4-bromo -5-(bromomethylene)-3- butyl-2(5 H)-furanone.	Mimic AHLs and disrupt signaling, disrupt motility and biofilm formation	
Green macroalgae Ulva rigida Honaunau Bay coral reef bacterial community, specially marine cyanobacterium	Brominated furanone Honaucins A to C	Inhibit quorum sensing V. harvery biofilm formation and E. coli AHL inhibition	
Leptolyngbya sp. Seed exudates (Medicao sativa)	L-canavanine(L-α-Amino-γ- (guanidinooxy)-n-butyric acid)	Inhibit the expression of QS-regulated phenotype exopolysaccharide production.	
Streptomyces soil isolate	Tricyclic polypeptide siamycin	blocked QS regulated feature i.e gelatinase production	
Sweet basil Ocimum basilicum	Rosmarinic acid(R-O-(3,4- Dihydroxycinnamoyl)-3-(3,4- dihy- droxyphenyl) lactic acid)	Inhibit protease, elastase, hemolysin production, biofilm formation and virulence factor	
Vanilla beans extract (Vanilla planifolia)	Vanillin (4-Hydroxy-3-methoxybenzaldehyde)	Interfere with AHL receptors. Inhibit C4-HSL, C6-HSL, C8-HSL, 3-oxo-C8-HSL. Inhibit biofilm formation in Aer. hydrophila	
Dichotella gemmacea	Juncin	potent nontoxic antilarval settlement	
See grass Halodule pinifolia, Cymodocea serullata	carrageenan type	Antibiofilm	
Avicennia marina, Rhizophora mucronata	amino, carbonyl and phosphoryl functionalities, aliphatic (fatty acids), NH2	Antibiofilm	
Pseudomonas sp. strain PAI-A	AHL-acylase (PvdQ)	Degrade C10-HSL, 3-oxo-C10-HSL, C12- HSL, 3-oxo-C12-HSL, C14-HSL, C16-HSL	
Aspergillus niger IAM 2094 Bacillus megaterium	AHL-lactonase (Gluconolactonase) Oxidoreductase	Lactone ring hydrolysis Oxidizes; C12-HSL, 3-oxo-C12-HSL, C14- HSL, 3-oxo-C14-HSL, C16-HSL,	

More than 20,000 metabolites have been reported from marine weeds from 1970s and a considerable percentage showed antibacterial, antifungal, antiprotozoan activities. Though major part of their exact mode of action still remained a secret, they were well reported as antiadhesive, low pH content, toxic, anesthetics and sometime they act as biological signal breaking agents. Maximum secondary metabolites from sea weeds were of terpenoid group (the largest group of natural products) or they were brominated in their biochemical structure.

Quorum quenching molecules

Biofouling may be controlled by using Quorum Quenching (QQ) strategies now adays (Kalia et al., 2015, Feng et al., 2013). Inhibition of biofilm formation in liquid-solid interfaces through quorum quenching is promoted. It has been reported that 60% of bacterial species sampled from biofouled Reverse Osmosis membranes, collected from a water treatment plant produced Quorum Sensing (QS) molecules. Such microorganisms actively participate in biofilm formation on membranes, suggesting that biochemical control of biofilm formation by inhibiting Quorum Sensing signals could be an effective way to reduce membrane biofouling (Diggle et al., 2007, Feng et al., 2013).

Quorum Sensing inhibition can provide considerable and effective means to control biofilm growth without the application of growth-inhibiting agents (Lade *et al.*, 2014). Quorum Quenching (QQ) molecules which have been characterized and reported up to date, generally use three strategies to combat autoinducer systems. The molecules interfere with Quroum Sensing signal production and disturb the synthesis, disrupt accumulation or degrade the Quorum Sensing molecules.

Many reports showed that natural compounds such as vanillin, ajoene, furanones, flavonoids, curcumin, Iberin, patulin etc. (**Table 1**) and few enzymes notably group of acylase, lactonase, oxidoreductase showed potential quorum quenching activity (Lade *et al.*, 2014, LaSarre and Federle 2013) against biofouling bacteria without interfering with their growth. Extracts of sea grass, mangroves were also studied to reduce quorum sensing controlling phenomenon such as biofilm formation and reports stood at a considerable appreciation (Prabhakaran *et al* 2012). Furthermore, immobilization of quorum quenching bacteria as well as enzymes, by bead-entrapment has been implied to MBR as a new biofouling control technology (Suk OH *et al.*, 2012).

Conclusions

Biofilms develops into biofouling only when "threshold of interference" oversteps and microbiota becomes "nuisance". It is one of the major vulnerable problems currently disturbing many ecological niches as well as in shipping and other industrial aquatic processes like membrane technology and maintenance. System performance only gets hampered when bacterial count exceeds 10^4 cfu/cm². Ban on oraganotin compounds as antifouling coating agents on ship and ship hull raised the urge of introduction of safe biomaterials into the antifouling research platform. Gradually for controlling membrane biofouling in water systems, biotools has been explored for safe and sustainable management.

Authors' Contribution

AQ initiated and prepared the review text, SG and SP have contributed equally in compiling the technical literature of review, AK and HJP supported overall work.

Competing Interest

None of the authors have any financial and technical competing interest.

Acknowledgements

The authors wish to thank the Director of CSIR-National Environmental Engineering Research Institute (NEERI), Nagpur, CSIR-Network Project (ESC0306 Activity 3.4.2) Government of India for providing necessary funds and facilities. SG is thankful to CSIR, New Delhi and SP is thankful to DST-INSPIRE for granting Junior Research Fellowships.

REFERENCE S

- Anand S., Singh D., Avadhanula M. and Marka S. 2014. Development and control of bacterial biofilms on dairy processing membranes. Comprehensive Rev. *In Food Sci. and Food Safety*, 13: 18-33 doi: 10.1111/1541-4337.12048
- Burke, C., Thomas, T., Egan, S. and Kjelleberg, S. 2007. The use of functional genomics for the identification of a gene cluster encoding for the biosynthesis of an antifungal tambjamine in the marine bacterium Pseudoalteromonas tunicata. Environ Microbiol, 9(3), 814-818. doi: 10.1111/j.1462-2920.2006.01177.x
- Carsten Matz 2011. Competition communication, cooperation: Molecular crosstalk in multispecies biofilms highlights. Springler series on Biofilms 5, doi: 10.1007/978-3-642-19940-0 2
- Chang, W. S., van de Mortel, M., Nielsen, L., de Guzman, G. N., Li, X. and Halverson, L. J. 2007. Alginate production by Pseudomonas putida creates a hydrated microenvironment and contributes to biofilm architecture and stress tolerance under water-limiting conditions. J Bacteriol, 189(22), 8290-8299. doi:10.1128/JB.00727-07
- Chapman, J., Hellio, C., Sullivan, T., Brown, R., Russell, S. and Kitereington, E. 2014. Bioinspired Synthetic Macroalgae: Examples From Nature For Antifouling Applications Int. Biodeterior. Biodegr. 05/2013 doi: 10.1016/j.ibiod.2013.03.036

- Chelsea, K.M. and Brennan, A.B. 2012. *Bio-Inspired Antifouling Strategies* Ann Rev of Materials Research. 42(1), 211-229. *doi:* 10.1146/annurev-matsci-070511-155012
- Chiellinia, C., Iannellib, R., Modeoa, L., Bianchib, V. and Petronia, G, 2012. Biofouling of reverse osmosis membranes used in river water purification for drinking purposes: analysis of microbial populations Biofouling 28:9 969-84 *doi:* 10.1080/08927014.2012.724679
- Davey, M.E. and O'Toole, G.A. 2000. Microbial biofilms: from ecology to molecular genetics. Microbiol and Mol Biol Reviews, 64(4), 847-867. doi: 10.1128/MMBR.64.4.847-867.2000
- Decho, A.W., Norman, R.S. and Visscher, P.T. 2010. Quorum sensing in natural environments: emerging views from microbial mats. Trends Microbiol 18: 73-80 doi: 10.1016/j.tim.2009.12.008. Epub 2010
- Diggle, S.P., Griffin, A.S., Campbell, G.S. and West, S.A. 2007. Competition and conflict in quorum-sensing bacterial populations Nature 450: 411-414 PMID: 18004383
- Dusane, D.H. Zinjarde, S.S. Venugopalan, V.P. Mclean R.J.C. Weber M.M. and Rahman P. K.S.M. 2010. Quorum sensing: implications on Rhamnolipid biosurfactant production, Biotechnology and Genetic Engineering Reviews, Vol. 27, 159-184 doi:10.1186/1754-1611-2-13
- Dworkin, M. and Kaiser, D. 1985. Cell interactions in Myxobacterial growth and development. *Science*, 230(4721), 18-24. doi: 10.1126/science.3929384
- Elasri, M.O. and Miller, R.V. 1999. Study of the response of a biofilm bacterial community to UV radiation. Applied and Environmental Microbiology, 65(5), 2025-2031. PMID:10223995 ISBN 978-1-904455-96-7.
- Feng, L., Wu, Z. and Yu, X. 2013 Quorum sensing in water and wastewater treatment biofilms. J. Environ. Biol., 34: 437-444 ISSN 0254-8704 DODEN: JEBIDP
- Flemming, H. 2011. The perfect slime. *Colloids Surf B Biointerfaces*., 251-259.
- Flemming, H.C. and Wingender, J. 2010. The biofilm matrix Nature Reviews Microbiology 8, 623-633, doi:10.1038/nrmicro2415
- Flemming, H.C., Neu, T.R. and Wozniak, D.J. 2007. The EPS matrix: The "House of Biofilm cells" J Bacteriol. 2007 189 (22): 7945-7947 doi: 10.1128/JB.00858-07
- Flemming, H.S. and Leis, A. 2002. In Encyclopedia of Environmental Microbiology (ed. Bitton. G) 2958-2967 (Wiley, New York,) *doi:* 10.1002/9780470015902.a0000342.pub2
- Fuqua, C., Parsek, M.R. and Greenberg, E. P. 2001. Regulation of gene expression by cell-to-cell communication: acylhomoserine lactone quorum sensing. Annual review of genetics, 35(1), 439-468.
 driver 10, 1147 (Communication constrained for the sensitive constrained for the senset of the sensitive constrained for the sensitive constrained

doi: 10.1146/annurev.genet.35.102401.090913

- Halan, B., Buehler, K. and Schmid A. 2012. Biofilms as living catalysts in continuous chemical syntheses. Trends in biotechnology, 30(9), 453-465.
- Hall-Stoodley, L., Costerton, J. W. and Stoodley P. 2004. Bacterial biofilms: from the natural environment to infectious diseases. *Nature Reviews Microbiology*, 2(2), 95-108. doi:10.1038/nrmicro821
- Hoffman, L.R., D'Argenio, D.A., MacCoss, M.J., Zhang, Z., Jones, R.A. and Miller, S.I. 2005. Aminoglycoside antibiotics induce bacterial biofilm formation. Nature, 436(7054), 1171-1175. doi:10.1038/nature03912

- Jefferson, K.K. 2004. What drives bacteria to produce a biofilm?. FEMS Microbiol Letters, 236(2), 163-173. doi.org/10.1111/j.1574-6968.2004.tb09643.x
- Jenkinson, H. F. and Lappin-Scott, H. M. 2001. Biofilms adhere to stay. TRENDS in Microbiol., 9(1), 9-10. doi: org/10.1016/S0966-842X(00)01891-6
- Kalia, V.C. and Purohit, H.J. 2011. Quenching the quorum sensing system: potential antibacterial drug targets Crit Rev Microbiol. 37(2):121-40
- doi:10.3109/1040841X.2010.532479
- Kalia, V.C., Kumar, P., Pandian, S.T. and Sharma P. 2015. Biofouling control by quorum quenching. In Hb25_Springer Handbook of Marine Biotechnology (431-440). Springer Berlin Heidelberg.
- Kaplan, J. B., Ragunath, C., Ramasubbu, N. and Fine, D. H. 2003. Detachment of Actinobacillus actinomycetemcomitans biofilm cells by an endogenous beta-hexosaminidase activity. *J.Bacteriol*, 4693-4698.
- Kim, H. J., Boedicker, J. Q. and Ismagilov, J. W. 2008. Defined spatial structure stabilises a synthetic multispecies bacterial community. *PNAS*, 18188-18193.
- Kirisits, M.J. and Parsek, M.R. 2006. Does Pseudomonas aeruginosa use intercellular signalling to build biofilm communities? *Cellular microbiology*, 8(12), 1841-1849. doi:10.1111/j.1462-5822.2006.00817.x
- Kolenbrander, P. E., Anderson, R. N. and Holdemna, L. V. 1989. Coaggregation of *Fusobacterium nucleatum*, *Selenomonas flueggei*, *Selenomonas inflix*, *Selenomonas noxia and Selenomonas putigena* with strains from 11 genera of oral bacteria. *Infect Immun*, 3194-3203.
- Korstgens, V., Flemming, H.C., Wingender, J. and Borchard, W. 2001. Influence of Ca2+ ions on the mechanical properties of a model biofilm of mucoid Pseudomonas aeruginosa *Water. Sci. Technol.*, 43: 49-57, 02/2001; 43(6):49-57. pubmed/11381972
- Lade, L., Paul, D. and Kweon, J.H. 2014. Quorum Quenching Mediated Approaches for Control of Membrane Biofouling *Int. J. Biol. Sci.* 10(5): 550-565 doi: 10.7150/ijbs.9028
- LaSarre, B. and Federle, M.J. 2013. Exploiting quorum sensing to confuse bacterial pathogens Microbiol Mol Biol Rev. Mar; 77(1):73-111. doi:10.1128/MMBR.00046-12
- Lear, G. and Lewis, G.D. (editor) 2012. Microbial Biofilms: Current Res and Appli.Caister Academic Press ISBN 978-1-904455-96-7.
- Leck, C. and Bigg, E.K. 2005. Biogenic particles in the surface microlayer and overlaying atmosphere in the central Arctic Ocean during summer Tellus B 57:305-316 doi: 10.1111/j.1600-0889.2005.00148.x.
- Magin, C.M., Cooper, S.P. and Brennan, A.B. 2010 Non-toxic antifouling strategies Materials Today 13(4) 36-44 doi: 10.1016/S1369-7021(10) 70058-4
- Marcato, R.C.E., Pechaud, Y., Paul, E., Girbal-Neuhauser, E., Dossat-Létiss, V. 2012. Removal of microbial multi-species biofilms from the paper industry by enzymatic treatments Biofouling: *The Journal of Bioadhesion and Biofilm Research*, 28(2), 305-314 doi: 10.1080/08927014.2012.673122
- Miller MB and Bassler BL 2001. Quorum Sensing in Bacteria, Annual Review of Microbiology Vol. 55: 165-199 doi: 10.1146/annurev.micro.55.1.165
- Nguyen, T., Roddick, F.A. and Fan, L. 2012. Biofouling of water treatment membranes: A review of the underlying

causes, monitoring techniques and control measures. Membranes 2:4 804-840 doi:10.3390/membranes2040804

- O'Toole, G.A. *et al* 2000. Biofilm formation as microbial development. Ann. Rev. Microbiol. 54: 49-79 *doi*: 10.1146/annurev.micro.54.1.49.
- Otani, M., Oumi, T., Uwai, S., Hanyuda, T., Prabowo, R.E. 2007. Occurrence and diversity of barnacles on international ships visiting Osaka Bay, Japan and the risk of their introduction Biofouling 23: 277-86 doi:10.1080/08927010701315089
- Otzen, D. and Nielsen, P.H. 2007. We find them here, we find them there: functional bacterial amyloid Cell Mol. *Life Sci.*, 65, 910-927 doi: 10.1007/s00018-007-7404-4
- Prabhakaran, S., Rajaram, R., Balasubramanian, V. and Mathivanan, K. 2012. Antifouling potentials of extracts from seaweeds, seagrasses and mangroves against primary biofilm forming bacteria *Asian Pacific J of Tropical Biomedi*, S316-S322 doi:10.1016/S2221-1691(12)60181-6
- Russell, D.M., George, A. and O'Toole, 2009. The developmental model of microbial biofilms: ten years of a paradigm up for review for feature review Trends in Microbiol. Doi: 10/1016/j.tim 2008.11.001.Cell press
- Salta, M.S., Wharton, J.A., Stoodley, P., Dennington, S.P., Goodes, L.R., Werwinski, S., Mart, U., Wood, R.J.K. and Stokes, K.R. 2010. Designing biomimetic antifouling surfaces Phil. *Trans. R. Soc.*, A 368, 4729–4754 doi:10.1098/rsta.2010.0195
- Schultz, M.P. 2007. Effects of coating roughness and biofouling on ship resistance and powering. Biofouling 23(5-6):331-4 PMID 17852068
- Shrout, J. D., Chopp, D. L., Just, C. L., Hentzer, M., Givskov, M. and Parsek, M. R. 2006. The impact of quorum sensing and swarming motility on Pseudomonas aeruginosa nutritional condition Mol Microbiol 62, 1264–1277 doi:10.1111/j.1574-6968.2008.01089.x
- Simoes, L. Simoes, M. and Vieira, M.J. 2008. Intergenic coaggregation among drinking water bacteria: Evidence for a role of Acinetobacter calcoaceticus as a bridging bacterium. *Appl Environ Microbiol*, 1259-1263.
- Stewart, P. and Franklin, M. 2008. Physiological heterogeneity in biofilms. *Nat Rev Microbiol*, 199-210.
- Stewart, P.S., Costerton, J.W. 2001. Antibiotic resistance of bacteria in biofilms Lancet 358 (9276): 135-138 doi.org/10.1016/S0140-6736 (01)05321-1
- Suk, O.H., Yeon, K.M., Yang, C.S., Kim, S.R., Lee, C.H., Park, S.Y., Han, J.Y., and Lee, J.K. 2012. Control of Membrane Biofouling in MBR for Wastewater Treatment by Quorum Quenching Bacteria Encapsulated in Microporous Membrane doi: 10.1021/es204312u
- Sutherland, I.W 2001. The biofilm matrix-an immobilized but dynamic microbial environment Trends Microbiol 9:222-227 PMID:11336839
- Vornhagen, J., Stevens, M, and A.H., R. 2013. Coaggregation occurs amongst bacteria within and between domestic showerheads biofilms. *Biofouling*, 53-68.
- Wang, D., Ding, X. and Rather, P.N. 2001. Indole can act as an extracellular signal in Escherichia coli. J Bacteriol.; 183:4210–4216. doi: 10.1128/JB.183.14.4210-4216.2001
- Waters, C.M., Bassler, B.L. 2005. Quorum sensing: cell-to-cell communication in bacteria. Annu Rev Cell Dev Biol. 21:319-46. *doi*: 10.1146/annurev.cellbio.21.012704.131001.
- Yebra, D.M., Soren, K. and Kim, D.J. 2004. Antifouling technology-past, present and future steps towards efficient

International Journal of Recent Advances in Multidisciplinary Research

and environmentally friendly antifouling coatings, Progress in Organic Coatings 50: 75-104 *doi*:10.1016/j.porgcoat.2003.06.001.

Zogaj, X., Nimtz, M., Rohde, M., Bokranz, W. and Romling, U. 2001. The multicellular morphotypes of Salmonella

typhimurium and Escherichia coli produce cellulose as the second component of the extracellular matrix Mol. Microbiol. 39, 1452-1463. doi: 10.1046/j.1365-2958.2001.02337.x
