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Doctoral thesis

THE IMPACT OF SUBMARINE OUTFALLS OF TREATED WASTEWATER ON THE MICROBIAL COMMUNITY STRUCTURE AND THE RESISTOME OF THE COASTAL MARINE ENVIRONMENT

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Doktorski rad

UTJECAJ PODMORSKIH ISPUSTA PROČIŠĆENE OTPADNE VODE NA STRUKTURU MIKROBNE ZAJEDNICE I REZISTOM PRIOBALNOG MORA

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THE IMPACT OF SUBMARINE OUTFALLS OF TREATED WASTEWATER ON THE MICROBIAL COMMUNITY STRUCTURE AND THE RESISTOME OF THE COASTAL MARINE ENVIRONMENT

Marija Kvesić Ivanković

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Center of Excellence for Science and Technology-Integration of Mediterranean region (STIM), Faculty of Science, University of Split Abstract:

The impact of submarine outfalls on the coastal marine microbiome and resistome remains insufficiently analysed, emphasizing the main objective of this dissertation. Modern technologies, including fast-profiling probes and next-generation sequencing, were employed to fill knowledge gaps and provide new insights into submarine outfalls as epidemiologically significant sources as well as hotspots of environmental contamination with antibiotic-resistant bacteria (ARB) and resistance genes (ARGs). Analysis of the water column stability and its influence on the effluent plume movement confirmed the role of stratification in submerging effluents beneath the pycnocline. Additionally, probes' utility in describing plume shape and its real-time detection was emphasized and subsequently used as a sampling methodology in the chapter Additional results. Metagenomic analyses furthermore demonstrated the negative influence of submarine outfalls on the marine microbial community, leading to the proliferation of nonindigenous and pathogenic bacteria potentially carrying ARGs. Furthermore, identification of carbapenemase-producing Enterobacteriaceae (CPE) in the vicinity of the submarine outfall associated with hospital wastewater was notable, evidencing the initial detection of environmental CP-E. coli containing both blakpc-2 and blaOXA-48 carbapenemase genes. Additional findings highlighted differences in microbial community structure between outfall locations and areas unaffected by wastewater intrusion. Moreover, the integration of shotgun analysis across WWTPs and corresponding submarine outfalls provided insights into complex bacterial community and ARG variations, underlining AR's persistence and release into the marine environment. Emphasizing the need for holistic strategies, this work aligns with the One Health concept, recognizing the problem of antibiotic resistance in the environment.

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Sažetak:

U sklopu ove disertacije analiziran je utjecaj podmorskih ispusta na mikrobiom i resistom priobalnog morskog okoliša. Moderna tehnologija, uključujući multiparametarske sonde i sekvenciranje sljedeće generacije, korištena je kako bi se popunile postojeće rupe u znanju i pružile nove spoznaje o podmorskim ispustima kao epidemiološki važnim izvorima ali i žarištima onečišćenja okoliša rezistentnim bakterijama i genima antibiotske rezistencije. Analizirana je stabilnost vodenog stupca i njegov utjecaj na širenje efluenta, potvrđujući ulogu stratifikacije u održavanju efluenta ispod piknokline. Nadalje, naglašena je i korisnost korištenja sondi za opis oblaka efluenta i njegovu detekciju u stvarnom vremenu, a koje su zatim i korištene kao metodologija uzorkovanja morske vode za analize prikazane u poglavlju Dodatni rezultati. Paralelno, primjenom metagenomske analize utvrđen je negativan utjecaj rada podmorskih ispusta na sastav mikrobne zajednice, a koji je doveo do širenja alohtonih i patogenih bakterija. Nadalje, u blizini podmorskog ispusta povezanog s otpadnim vodama bolnica grada Splita, identificirane su karbapenemaza-producirajuće Enterobacteriaceae, a po prvi put se identificirala i okolišna KPC- i OXA-48-producirajuća E. coli. Dodatni rezultati istaknuli su razlike u strukturi mikrobne zajednice između lokacija u neposrednoj blizini podmorskih ispusta i područja koja nisu pod utjecajem otpadne vode. Nadalje, integracija shotgun analize provedene unutar uređaja za pročišćavanje otpadnih voda i u blizini pripadajućih podmorskih ispusta pružila je uvid u raznolikost bakterijske zajednice i gena antibiotske rezistencije, pri tome naglašavajući postojanost i ispuštanje gena antibiotske rezistencije u morski okoliš. Naglašavajući potrebu za holističkim strategijama, ovaj rad se uklapa i u 'One Health' koncept (Jedno zdravlje), prepoznajući i naglašavajući problem antibiotske rezistencije u okolišu.

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ABBREVIATIONS:

AR	Antibiotic resistance
ARB	Antibiotic resistant bacteria
ARG	Antibiotic resistance genes
CDOM	Colored dissolved organic matter
CPE	Carbapenemase-producing Enterobacteriaceae
CRE	Carbapenem-resistant Enterobacteriaceae
DO	Dissolved oxygen
ESBL	Extended spectrum β-lactamases
FIB	Faecal indicator bacteria
HGT	Horizontal gene transfer
HWWs	Hospital wastewaters
KPC	Klebsiella pneumoniae carbapenemase
MBLs	Metallo-β-lactamases
MDR	Multidrug-resistant
MGE	Mobile genetic elements
NGS	Next-generation sequencing
OTUs	Operational taxonomic units
OXA	Oxacillin-hydrolyzing β-lactamases
PBPs	Penicillin-binding proteins
VREfm	Vancomycin-resistant Enterococcus faecium
WHO	World Health Organization
WWTPs	Wastewater treatment plants

CONTENTS

1. INTRODUCTION
1.1.Submarine outfalls – transport and disposal of treated wastewater in coastal areas
1.2.Microbiome and resistome of the aquatic environment-role of WWTPs and submarine outfalls4
1.2.1. Microbial community – important aspect of aquatic health
1.2.2. Antibiotic resistance determinants in coastal area
1.2.3. Environmental metagenomics
1.3. Carbapenem resistance remains in hospitals?9
2. AIMS AND SCOPES OF THE THESIS
3. SCIENTIFIC PAPERS
3.1.Spatial and temporal vertical distribution of chlorophyll in relation to submarine wastewater effluent discharges
3.2. Microbiome and resistome of coastal waters under the influence of submarine outfalls35
3.3. Submarine outfalls of treated wastewater effluents are sources of extensively- and multidrug-resistant KPC- and OXA-48-producing Enterobacteriaceae in coastal marine environment
4. ADDITIONAL RESULTS AND DISCUSSION
4.1. Comparison of microbial community structure in coastal waters not affected by wastewater 63
4.1.1. Methodology
4.1.2. Result and discussion 64
4.2.Microbiome and resistome of WWTPs and submarine outfalls
4.2.1. Methodology
4.2.2. Results and discussion
5. CONCLUSION AND FUTURE REMARKS
6. ADDITIONAL REFERENCES
7. SUPPLEMENTAL MATERIAL 100
7.1.Supplemental material to Paper II
7.2.Supplemental material to Paper III
8. CURRICULUM VITAE
9. LIST OF PUBLICATIONS

1.INTRODUCTION

1.1.Submarine outfalls – transport and disposal of treated wastewater in coastal areas

Inherent complexities and dynamic nature render coastal areas among the most vulnerable ecosystems. Their importance goes beyond biodiversity and plays a crucial role in supporting the socio-economic aspects of human life and activities [1]. Unfortunately, these vital regions face escalating anthropogenic pressures, particularly due to tourism development and the growing coastal population, which now represents approximately one third of the world's population [2]. As a result, the quality of the coastal ecosystem is compromised, exerting substantial pressure on coastal water bodies. Moreover, as the expected human population will grow to 10 billion by 2050 [3], these environmental pressures are expected to increase. Therefore, effective protection of coastal ecosystems is now more important than ever.

The primary sources of pollution in coastal regions originate from untreated or inadequately treated wastewater, as well as runoff from agricultural areas and rivers. These sources contribute to various forms of pollution that can directly or indirectly impact human health. Various physico-chemical (suspended matter, nutrients, heavy metals etc.) and biological stressors (pathogens, antibiotic resistance (AR) determinants etc.), being introduced into the ecosystem by sewage water, contribute to water quality degradation. Disturbances in the intake and processing of organic matter and nutrients, both from natural sources and anthropogenic influence, disturb the fragile balance of this ecosystem. When the intake of nutrients, particularly nitrogen and phosphorus compounds, exceeds the ecosystem's assimilation capacity, it leads to eutrophication and degradation of water quality [4,5]. However, among all mentioned pollution sources, wastewater treatment plants (WWTPs) are known routes through which most of the mentioned contaminants are processed and disseminated further into the environment. WWTP receives and processes large volumes of wastewater enriched in physico-chemical and biological pollutants originating from municipal, industrial and hospital sewage. Thus, coastal environments are under the combined pressure of a complex matrix of contaminants being introduced by WWTP effluents. Discharging wastewater effluent, furthermore, promotes an intrusion of non-native bacterial populations, including pathogenic and antibiotic resistant bacteria (ARB), into marine environments [6,7].

Particularly for coastal areas, transporting and disposing of treated wastewater effluent into the marine environment is maintained by wastewater submarine outfalls, the discharge components of WWTPs. Outfalls consist of a pipeline, with varying length and diameter, attached to the seabed which mostly terminates with a multiport diffuser section to obtain a rapid mix and dilution effect (Figure 1). Effluent hydrodynamics can be defined as a mixing process occurring in two different regions. During the initial dilution, known as near-field, buoyancy flux and outfall geometry influence the effluent trajectory and degree of mixing. As distance from the source increases, the waste plume's characteristics become progressively less governed by the source and more by ambient environmental factors. These include passive diffusion and advection driven by tides, waves, pressure, and density gradients (known as far-field). Vertical mixing depends on the water column stratification, precisely vertical density gradient and the pycnocline formation. During the formation of stable vertical stratification, the effluent plume is trapped below the pycnocline layer, while upwelling to the surface is characterized during the disruption of vertical stratification (e.g. significant wind disturbances) [8,9].

Geometrical shapes of sewage plume, although rarely observed in detail, are indicators of plume dispersion and its impact on the ecosystem [10]. Dilution of the effluent plume can be estimated from in-situ measurements of its major components (i.e. initial dilution, rise height and vertical dispersion) performed in the near-field and far-field mixing zones. As the dilution effect often obscures the impact of effluent discharge into the sea, the difference between recipient and wastewater is minor and difficult to discern [11,12], even in cases involving substantial flows of inadequately treated wastewater [13]. Notably, the most pronounced effect of effluent discharge is visible in the near-field mixing zone, which weakens as it moves away toward the far-field mixing zone. While submarine outfalls are generally considered environmentally acceptable, they still pose a significant source of pollutants in the marine environment. Thus, precise measurements of variations of different parameters, which reflect the sewage impact, are crucial.

Water column density is one of the most crucial parameters for understanding the physical processes of wastewater effluent plume behaviour. It depends on temperature, salinity and pressure which all together describe vertical water column stability. In addition to the physical properties of the water column, chlorophyll *a*, turbidity, organic matter and dissolved oxygen are also important parameters to obtain information about the water quality status. Since primary producers are most sensitive to an increase in nutrient concentration,

phytoplankton biomass is considered a good indicator of the trophic state of coastal water [14], and due to the challenges in the direct measuring of phytoplankton, chlorophyll *a* is mainly used as a proxy for phytoplankton biomass [15–18]. Changes in chlorophyll *a* response will be followed by dissolved oxygen variation, as it primarily depends on respiration rate, but also the bacterial decomposition of elevated organic matter introduced by effluent.

The most traditional technique to determine the plume effect is based on bottle sampling and laboratory analysis. Although such analysis provides key information about the plume effect, it lacks high spatial and temporal resolution and does not permit real-time plume detection. Nowadays, different fast-profiling probes equipped with various physico-chemical sensors are used to obtain high-quality real-time data to map and describe effluent plume shapes. Consequently, continuous monitoring of effluent and physicochemical parameters within the water body is crucial in the intrusion area to obtain long-term ecosystem health.

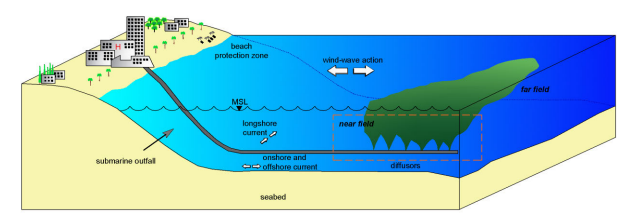


Figure 1. Schematic representation of submarine outfalls design (Credits: Morena Galešić Divić and Vladimir Divić)

1.2.Microbiome and resistome of the aquatic environment – role of WWTPs and submarine outfalls

1.2.1. Microbial community – important aspect of aquatic health

Aquatic ecosystems as a natural habitat have a high diversity of bacterial species, but at the same time are receptors of microorganisms originating from land (animal and human waste, runoffs etc.). Due to their pivotal role in water quality purification and nutrient circulation, microbial communities play a crucial role in these environments. Therefore, understanding their composition, abundance, and driving forces holds paramount importance for the effective management of coastal areas.

Along with all physico-chemical agents that WWTPs receive, treat and discharge further into the environment, diverse microbial communities from different sources make up the vast majority of sewage composition, as bacteria are the most abundant and diverse group of organisms living in wastewater. While WWTPs are designed to remove a considerable portion of pollutants, residues still expose environmental microorganisms to so-called "second-hand" pollution [19]. As a result, the release of insufficiently treated or untreated wastewater leads to significant changes in the microbial communities of lakes [20], rivers [21,22], and seas [23], and can impact the long-term biodiversity of the coastal marine microbiome too [24,25].

The link between microbial communities and different physico-chemical and biological stressors in WWTPs and the environment is well known, and all of them affect microbial communities in terms of structure [26] and function [27]. Generally, marine microorganisms are highly sensitive with prompt responses to most of the environmental variations. Thus, microbiome composition depends on nutrient concentration changes, promoting the shift in the composition which could lead to reduced biodiversity [28]. Additionally, other physicochemical parameters such as pH [29], temperature [30], salinity [31], and organic matter [32] are also closely linked to the composition and abundance of the microbial community. Hence, microorganisms are considered in situ biosensors, detecting environmental changes [33], and therefore, retrieving quantitative data and understanding the mechanisms that influence microbial community is essential to predict the impact of the microbial response to the changes and to determine their consequences on local and global scale [34]. However, among allochthonous microbial community discharged into the coastal area by submarine outfalls, the presence of different human pathogens (e.g., *Enterococcus faecalis, Escherichia*

coli, *Klebsiella pneumoniae*, *Pseudomonas aeruginosa* etc.) is of additional concern. If mentioned pathogens are resistant to different antimicrobials by harbouring AR genes (ARG), such a concern increases even more.

1.2.2. Antibiotic resistance determinants in coastal area

Whether produced naturally (from different microorganisms) or synthetically all antibiotics have the same role: to inhibit bacterial growth or reproduction (bacteriostatic) or to kill the bacteria (bactericidal) [35]. The historical lineage of antibiotics extends into the ancient past, with evidence of their presence dating as far back as 2 billion to 30 million years ago [36]. Anyhow, the human discovery of antibiotics dates back to Fleming's discovery of penicillin in 1929 [37], and there is no doubt, that antibiotics are one of the most important discoveries of modern medicine. These 'miracle drugs,' once referred to as such, transformed life-threatening bacterial infections into easily treatable conditions. [38]. However, a miracle has its own price, an AR, a phenomenon present all around the world. What has happened that switched 'miracle drugs' into the 'problem of a modern century'?

Antibiotics have shown high potential in the treatment of different diseases in humans, plants and animals, but their excessive and inappropriate use [39] became the main driving force of the AR spread, which has led to this phenomenon being widely recognized by international entities as a growing public health. It stands as one of the foremost challenges in modern medicine [40], giving rise to increased healthcare costs and elevated patient mortality. While earlier projections estimated approximately 10 million deaths attributable to AR by 2050 [41], recent analyses have shown that only in 2019 almost 5 million deaths were caused by multidrug-resistant (MDR) bacteria [42], which implies that the development of novel antimicrobials will not meet the future needs of human medicine [43].

While the AR phenomenon was initially regarded as a concern mainly within hospital settings, the widespread and excessive use of antibiotics has hastened the emergence and dispersion of acquired AR. Notably, WWTPs are recognized as primary sources of AR determinants into the environment [44–47], comprising ARB, ARGs, and mobile genetic elements (MGE) that facilitate successful horizontal gene transfer (HGT) even among unrelated bacterial species [48,49]. HGT can occur between two bacterial cells (donor-acceptor) through three different ways: 1) transformation – the most common pathway in which bacteria uptake DNA from their environment, 2) transduction – genetic elements are transferred

by bacteriophages, and 3) conjugation – pathway in which bacteria accepts new genes by physical contact between two cells (formation of conjugation bridge). ARB and ARGs experience heightened levels within patients and hospital environments. These elements are subsequently propagated into the broader community as a component of the biome or transported into WWTPs through hospital effluents. Additionally, sub-therapeutic antibiotic concentrations, along with their residues, are introduced into wastewater, intensifying the issue of AR by promoting selective pressures during prolonged microorganism-pollutant interactions (Figure 2). Consequently, hospital wastewaters deserves special attention as it directly introduces human opportunistic pathogens into the natural ecosystem [44,50,51], providing a platform for genetic material exchange with indigenous bacterial populations. However, from everything mentioned above, it is clear that WWTPs promote the persistence and proliferation of better adapted microorganisms [52], which makes the WWTPs considered reservoirs and hot spot sources of dissemination of AR determinants further into the environment. Although there are opposite findings about the removal of ARGs during different treatment processes in WWTPs, as some of the studies showed a reduction in abundancy [53,54], while others found elevated levels in treated effluent [55,56], most of the scientific literature agrees in one: most of the WWTPs are not capable to remove ARGs from the system. Given that AR poses a public health challenge on a global scale, in order to manage the risk of AR spread it becomes imperative to underscore the significance of researching this phenomenon in the environment too, as highlighted by the World Health Organization (WHO) under the One Health concept.

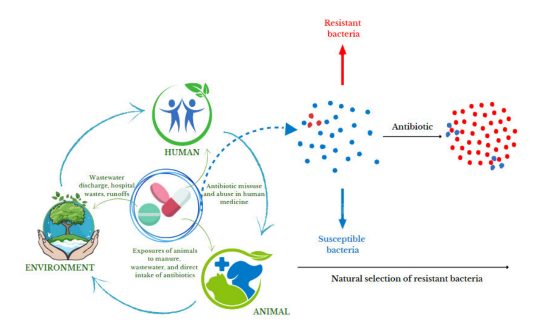


Figure 2. Distribution of ARGs through human, animal and environmental habitats providing an excellent opportunity for promoting AR

1.2.3. Environmental metagenomics

The determination of limits for microbiological parameters in bathing water is governed by the EU Directive on bathing water [57]. Similarly, in the Republic of Croatia, the Regulation on the quality of bathing water [58] places emphasis on monitoring and raising public awareness to ensure both environmental quality and human health. While this screening method is considered as 'the golden' standard, it relies on conventional approaches that assess seawater quality by quantifying indicator bacteria (*Escherichia coli* and intestinal enterococci). However, this method offers only a limited comprehension of environmental health. These traditional methods are time-consuming and yield information for less than 1% of cultivable bacteria [59]. To address these limitations, recent years have witnessed the growing adoption of advanced molecular biology and computer analysis techniques in environmental microbiology [60,61].

Slow growth with specific requirements, microbial competition and environmental stressors affect the bacterial culturing processes, remaining 80% of human gut bacteria and 99% of environmental bacteria being uncultured [62,63]. However, Next-generation sequencing (NGS), a powerful tool based on DNA sequencing directly from environmental samples [61], has found diverse applications, while enhancing our current understanding and generating novel insights across various environmental microbiology domains. Specifically, 16s rRNA amplicon sequencing can elucidate bacterial and archaeal community structures,

while shotgun sequencing is able to assess information about viral communities, ARGs and MGE too [64,65]. Metagenomics ensures non-targeted approaches to describe complex microbial communities and associations between such microorganisms in the environment. It enables the study of both, culturable and non-culturable bacteria, overcoming the limitations of laboratory isolation and cultivation. In this way, new insights about the current understanding of microbial structure, richness and distribution are and will be more provided [66]. Moreover, such a non-targeted approach has the potential for a wider application in public health surveillance as an early warning system [67].

How did it all begin? After NGS became commercially available in 2005, it evolved into a standard technique driven by the quest to compare different domains (especially the environment associated with the human gut) [68,69]. Consequently, various statistical analyses have been developed and used [70–72], and additionally, the comprehensive reference genome catalogue of the human microbiome has been compiled [73]. However, as the cost of NGS methods decreased, their application was no longer limited to the human gut, but was also used in other habitats (aquatic environment, WWTPs, animal microbiome, etc.). This led to several studies that provided insights on species or operational taxonomic units (OTUs), a species-like concept, conceived to explain the clusters obtained by NGS. These methods provided valuable new insights into species diversity around the world and how these communities change in response to environmental changes, as mentioned in earlier chapters.

1.3. Carbapenem resistance remains in hospitals?

Over the past decade, the alarming and widespread dissemination of gram-negative bacteria resistant to beta-lactam (β-lactam) antibiotics, notably carbapenems, has emerged as an imposing public health concern, transcending national boundaries and triggering a global alarm. Carbapenems, regarded as the last line of defence against infections caused by MDR bacteria [74], now face a dire threat from carbapenem-resistant bacteria, particularly those from the *Enterobacteriaceae* family (CRE). This has established itself as one of the most critical threats to public health [75,76]. Of particular distress is the limited progress in the development of antimicrobial drugs, especially for gram-negative pathogens [43].

Enterobacteria, commensals of the digestive systems of humans and animals and integral members of the normal intestinal flora, exhibit a dual nature, as particular species can also manifest as opportunistic human pathogens causing an array of diseases. These pathogenic species make up 70% of all infections caused by Gram-negative bacteria, and it includes infection of different human body systems (urinary, respiratory, gastrointestinal, etc.) with various organisms responsible for those diseases (*E. coli*, *Acinetobacter baumannii*, *Pseudomonas aeruginosa* and different *Klebsiella*, *Proteus* and *Enterobacter* species) [77]. The majority of these Gram-negative bacteria are resistant to multiple classes of antibiotics and are increasingly becoming resistant to almost all available antibiotics worldwide [78].

 β -lactams are frequently used antibiotics worldwide, and several antibiotics, including penicillins, cephalosporins, monobactams and carbapenems, belong to this class. They all share the common beta-lactam ring and act mostly by binding and inactivating the penicillin-binding proteins (PBPs), responsible for bacterial cell wall formation. Among all β -lactams, carbapenems (ertapenem, imipenem, meropenem, and doripenem) are the most potent members against Gram-negative and Gram-positive bacteria. Their structure ensures stability and effectiveness against most β -lactamases, enzymes which inactivate β -lactams, including AmpC beta-lactamases (AmpC), and the extended spectrum β -lactamases (ESBLs) [79]. Due to their wide and safe use, it is clear why the problem of carbapenem resistance is a globally disseminated problem. Therefore, in 2017, the WHO classified CRE in the highest-critical category among priority groups for multi-resistant pathogens requiring urgent drug discovery [40].

Resistance to carbapenems may be caused by different mechanisms (augmented drug efflux, a mutation in or loss of outer membrane porins, and production of carbapenemases), however, it primarily arises from the production of bacterial carbapenemases, enzymes that hydrolyse carbapenems, rendering them ineffective [80,81]. As mentioned earlier, the problem of AR is ancient and naturally founded in some species, and thus, carbapenem resistance is found intrinsically in some species like Stenotrophomonas maltophilia, Aeromonas spp., Chryseobacterium spp., and Bacillus spp. They possess intrinsic chromosomal β-lactamases, and therefore the use of carbapenems in a treatment for infections caused by those bacteria is not an option. Intrinsic carbapenem resistance is less common among clinically significant gram-negative bacterial species, yet it is often a result of mutational events or horizontal gene transfer. These acquired carbapenemases are classified into three molecularly distinct classes: A, B, and D (Figure 3). The respective genes are capable of dissemination among bacterial populations through mobile genetic elements such as plasmids, transposons, and integrons [82,83]. In class A, a plasmid-mediated K. pneumoniae carbapenemase (KPC) is the most important, as it is the most prevalent and widely distributed carbapenemase [84]. Metallo-βlactamases (MBLs) are part of Class B carbapenemases, not inhibited by avibactam, normally used for carbapenemase-producing Enterobacteriaceae (CPE) infections. Finally, in class D are counted plasmid-mediated oxacillin-hydrolyzing β-lactamases (OXA) i.e. OXA-48, OXA-23 and other derivatives, of which some are frequently founded in European countries [85]. The first report about CPE was back in the early 1990s [86], and from that, CPE has spread all across the world [87].

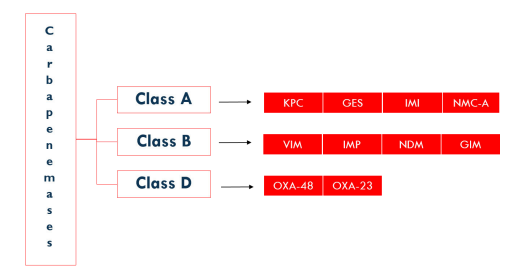


Figure 3. Molecular classification of acquired carbapenemases

While CRE presents a therapy challenge for patients, CPE pose a more significant issue for infection prevention and treatment. This is due to carbapenemase production in CPE, often facilitated by plasmids, enabling bacteria to transfer carbapenem resistance genes to other clinically pertinent species. Consequently, this hampers treatment options for various organisms, such as other *Enterobacteriaceae*, *Pseudomonas*, or *Acinetobacter*.

Information on ARGs in coastal areas is still scarce and mostly without mechanisms to explain certain significant relationships between environmental factors and ARGs. Moreover, it is difficult to make predictions about the impact of AR on environmental and health risks due to a lack of data [47]. However, it is clear that AR determinants occur in different environments, especially those under anthropogenic pressure, and WWTP effluents are considered to be important pathways for their spread. Besides WWTPs generally, hospital wastewaters (HWWs) play an important role in the spread of AR determinants as they are enriched with various ARBs and ARGs. However, only a few countries have recommended additional treatments for HWWs, although the enrichment of ARB are found through WWTPs [88–90]. Several studies have shown the spread of various CRE from WWTPs to the environment: CR *K. pneumonia*, [44,91,92], *E. coli* [91,92], *Citrobacter freundii* [93], etc. However, most of them have been found in freshwater rather than marine ecosystems. So far, carbapenemases have been identified in various pathogens worldwide, both in hospitals and outside hospitals [94].

In Croatia, *Enterobacteriaceae* producing KPC carbapenemase were first detected in hospitals in the northern part of the country [95], followed by observations in the southern region [96]. Notably, within Croatian hospitals situated in Zagreb and Split, the carbapenemases VIM and KPC exhibit the highest prevalence. Specifically, VIM-1 predominates in clinical *Enterobacteriaceae*, while KPC-2 is identified in *K. pneumoniae* [97,98]. In the natural environment of Croatia, the only case of KPC-producing *K. pneumoniae* was discovered in the Krapina River, downstream from the wastewater outlet of the Zabok General Hospital, indicating the hospital environment as its source [99].

A comprehensive review of the scientific literature shows that the focus is rather on CPE in hospitals, WWTPs, and freshwater ecosystems, while their occurrence in the marine environment has been insufficiently studied. Certainly, the effects of submarine discharges, especially in the near-field zone, have been insufficiently studied in the context of metagenomic analysis and the global problem of AR. Moreover, no research has been

conducted in Croatia on this critical issue. Considering the fact that AR is an urgent public health challenge on a global scale, it is essential to emphasise the importance of environmental research as highlighted by WHO under the One Health concept. Furthermore, the escalating trend of KPC-producing *Enterobacteriaceae* in hospitals in the Republic of Croatia points to their possible presence in the environment. This dynamic adds further motivation for this dissertation, which seeks to address the aforementioned global predicament and its growing impact within Croatia.

2.AIM AND SCOPES OF THE THESIS

Coastal areas under intense anthropogenic influence have emerged as significant reservoirs of bacterial resistance to antibiotics, where microbial community structure serves as an indicator of adverse influences. Framed within the One Health concept and considering the continuous increase of global AR, this research aims to provide new insights into the influence of submarine discharges on the spread of ARGs and resistant opportunistic pathogens in the coastal marine environment. Simultaneous analysis of the water column under the influence of the submarine outfall, by using fast-profiling probes and various molecular methods will provide solid insights into the extent to which WWTPs contribute to the spread of AR in the coastal marine environment.

Aims of this study:

- 1. Determination of the stability and quality of the water column by vertical profiling with multiparameter probes. Description of the effluent shape and behaviour by analysing physico-chemical parameters (temperature, salinity, density, buoyancy frequency (Brunt Väisälä frequency), chlorophyll *a*, turbidity, colored dissolved organic matter and dissolved oxygen)
- 2. Assessment of the biodiversity of the coastal marine bacterial community under the pressure of wastewater discharge through submarine outfalls using modern Next Generation Sequencing (NGS) methods
- 3. Analysis of the repertoire of ARGs available to the bacterial community using bioinformatics tools and shotgun sequencing
- 4. Analysis of the CRE genome with particular emphasis on resistance genes, virulence factors and types of resistant plasmids using the Whole Genome Sequencing (WGS) and appropriate bioinformatics tools

Main hypothesis from which this study proceeds:

H1: Vertical profiling of physico-chemical parameters using in-situ profiling probes will provide information on the effluent upwelling to the surface of the water column and provide better insight into the behaviour of the effluent plume

H2: Unstratified conditions allow effluent to upwell to the surface, leading to microbiological contamination of the surface layer of the water column

H3: Both the surface and bottom layers of the water column are under the influence of wastewater, as indicated by the composition of the microbial community, which is dominated by taxonomic groups characteristic of the human digestive tract and whose commensals also dominate the composition of the microbiome in WWTPs around the world

H4: There is a difference in the composition of the microbiome at the locations of the two different outfalls, suggesting that the composition of the effluent shapes the recipient microbial community

H5: The dissemination of carbapenem-resistant and carbapenemase-producing *Enterobacteriaceae* is facilitated by hospital wastewater, as indicated by the elevated prevalence of antibiotic-resistant opportunistic pathogens detected at the outfall which WWTP treats hospital effluents

H6: The most abundant antibiotic classes and ARGs in WWTPs reflect the most used antibiotics in Croatian community and hospitals

H7: WWTPs have limited removal efficiency, with the purification properties of seawater emerging as the predominant factor in the dilution of the effluent plume.

Specific scientific questions:

- What bacterial populations are prevalent at the bottom and surface of the water column at submarine outfall locations?
- Are clinically significant human pathogens such as *E. coli*, *K. pneumoniae*, *C. freundii*, and *A. baumannii* present at selected localities? If found, do they have lower susceptibility to antimicrobial drugs?
- Are there similarities between the carbapenem-resistant strains isolated in this study and clinical isolates at KBC Split or other hospitals in Croatia?
- Which classes of antibiotics and ARGs are the most abundant in both WWTPs and marine ecosystems? How much the concentration of pollutants (*E. coli*, intestinal enterococci, ARGs etc.) decreases during the treatments?

Scientific contribution:

The analysis of the environment, particularly those influenced by wastewater exposure, is extremely important for biodiversity preservation, environmental quality, and human well-being. Current public health challenges include monitoring of AR pathogens and the ubiquitous presence of ARGs, which exhibit robust survival within WWTPs. However, critical knowledge gap persists concerning the impact of submarine outfalls on the water column quality, with the specific focus on metagenomic analyses and the global distribution of ARGs.

This PhD thesis endeavours to make a scientific contribution, expanding existing knowledge and opening new perspectives on submarine outfalls as important epidemiological sources as well as potential hotspots of environmental pollution by ARB and ARGs. Furthermore, analyses such as these fit seamlessly into the One Health concept initiated by WHO revealing the intricate interdependencies between human, animal, and environmental health. Understanding the impact of submarine discharges on water column stability, microbial diversity and the spread of ARGs as well as AR human pathogens is important worldwide to assess the consequences for human health and to effectively control the problem of further resistance development. The results of this research can serve as a basis for a comprehensive risk assessment and the formulation of holistic strategies. These strategies in turn serve a dual purpose: reducing pollution and improving the protection of human health, both locally and globally.

3. SCIENTIFIC PAPERS

3.1. Spatial and temporal vertical distribution of chlorophyll in relation to submarine wastewater effluent discharges

Reproduced from:

Kvesić, M., Vojković, M., Kekez, T., Maravić, A., Andričević, R., 2021. Spatial and Temporal Vertical Distribution of Chlorophyll in Relation to Submarine Wastewater Effluent Discharges. Water 2021, 13, 2016. https://doi.org/10.3390/w13152016

The distribution of chlorophyll a in coastal waters is influenced by various factors, including hydrodynamics, biotic and abiotic processes. Spatial and temporal variations in chlorophyll a profiles provide valuable insights into changes occurring in the marine environment. Therefore, this study aims to characterise the vertical distribution of chlorophyll a and describe its variability as a function of factors such as stratification conditions and the presence of wastewater effluent. By obtaining high resolution data, these modern fast-profiling probes have been able to successfully describe the stability of the water column and the different vertical chlorophyll a profiles.

In addition to the successful use of fast-profiling probes in describing the characteristics of the effluent plume and the consequences left behind after intrusion (in terms of changes in physico-chemical parameters), these techniques can also be used to obtain real-time data and identify possible locations where wastewater has been discharged and could cause microbiological pollution and pose a health risk to humans. Such an application is used as a method to locate the effluent plume and perform sampling at that location. The results are discussed in the chapter 'Additional results'.





Article

Spatial and Temporal Vertical Distribution of Chlorophyll in Relation to Submarine Wastewater Effluent Discharges

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Abstract: The vertical distribution of chlorophyll in coastal waters is influenced by a combination of the hydrodynamic environment and different biotic and abiotic processes. The spatial and temporal occurrences of chlorophyll profiles provide a good representation of the changes in the marine environment. The majority of studies in the Adriatic Sea have so far been conducted in areas unaffected by anthropogenic pressure. Our study site is located near two marine outfalls, which are part of the public sewage system. This study aims to characterize the chlorophyll vertical distribution and describe its variability based on the stratification conditions and the presence of a wastewater effluent plume. Based on these conditions, we identified three characteristic scenarios/types of chlorophyll profiles. The first one occurs when the vertical mixing of the water column creates the upwelling of chlorophyll and nutrients to the upper part of the water column. The second and third scenarios occur during stratified conditions and differ by the extent of the effluent plume intrusion. Using modern fluorescence techniques, we identified and described three different vertical chlorophyll profiles, characterizing them according to their physical and biological parameters and processes. For cases with a visible effluent intrusion, we confirmed the importance of the pycnocline formation in keeping the effluent below and maintaining the higher water quality status at the top of the water column.

Keywords: chlorophyll vertical profiles; stratification; wastewater effluent plume intrusion



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1. Introduction

Phytoplankton has a major influence on marine ecosystems because of its role in oxygen production and overall significance in supporting and maintaining marine life. Due to the challenges in the direct measuring of phytoplankton, chlorophyll *a* is mainly used as a proxy for phytoplankton biomass [1–5]. Traditional techniques for chlorophyll analysis are based on bottle sampling, which is limited in terms of obtaining a high spatial and temporal resolution. This was corrected by modern in situ fluorescence techniques and remote sensing observations, which allow for measurements of chlorophyll in a high resolution. The research on chlorophyll temporal and spatial variability in the Mediterranean Sea is mostly restricted to the surface, as most of the collected data come from satellite observations and ex-situ laboratory measurements [5]. The oligotrophic status of the Mediterranean basin [6], with a decreasing trend from west to east [7,8], and seasonal and interannual variability [4,9,10] were observed based on surface chlorophyll analysis.

The vertical distribution of chlorophyll is under the influence of many biotic and abiotic processes, whose interplay affects the depth of the upper mixed layer and controls

Water 2021, 13, 2016 2 of 17

the vertical chlorophyll distribution [4,11]. So far, information about vertical chlorophyll profiles has been derived from different kinds of measurements [12–23].

A similar chlorophyll seasonality was noticed in the Adriatic Sea by remote sensing observations [24–26] and laboratory measurements [27,28]. The Adriatic Sea is a semi-enclosed basin of the Mediterranean Sea, where the biogeochemical parameters are influenced mainly by the circulation of the water masses. The chlorophyll trends in the Adriatic Sea have been investigated by remote sensing [29–31] and laboratory-based measurements [27,32] to find deep chlorophyll maxima [28,33] and to investigate the anthropogenic impact on the phytoplankton community structure and abundancy [34–36].

Coastal waters are highly sensitive ecosystems and may become even more vulnerable when exposed to the negative impact of human activities. Wastewater effluents contain high concentrations of pollutants, which can cause a large amount of deterioration to marine environments [37]. Bacteria are the most abundant and diverse group of organisms living in wastewater [38,39], and bathing water status is mainly determined using fecal indicator bacteria (FIB) as indicators of water quality and related human health issues [40,41]. Besides fecal pollution, eutrophication is another important aspect of coastal water quality. Since primary producers are the first components of an ecosystem to react to an increase in nutrient concentration, phytoplankton biomass is considered a good indicator of the trophic state of coastal water [42]. Changes in phytoplankton biomass abundance occur following nutrient enrichment, which can happen naturally (through riverine influence or the upwelling of nutrients from the bottom) or under anthropogenic influence (urban and industrial wastewater discharges and surface runoff). Anthropogenically introduced nutrients are the major problem of coastal ecosystems [43,44], and their mixing with seawater contributes to the eutrophication of coastal zone waters [45], thus increasing the biomass of macroalgae and seston algae [46].

This paper aims to characterize chlorophyll vertical profiles and describe their spatial and temporal variability based on the stratification status of a water column and the presence of a marine wastewater effluent plume. Due to the nature of fluorescence measurements, which change with the taxonomic species or environmental conditions [47] and can be affected by dissolved materials [48] or by different light irradiation statuses [49–51], the present study is focused on the relative distribution of vertical chlorophyll fluorescence in a water column by comparing different segments of the water column for each considered profile. To our knowledge, this is the first classification of chlorophyll vertical profiles in the presence of a wastewater effluent plume. The spatiotemporal variability of the different vertical profiles concerning stratification was also investigated.

2. Materials and Methods

2.1. Study Site

The study area is the eastern central Adriatic Sea, near the city of Split in Croatia (Figure 1). Two main marine wastewater outfalls are located within the study area, about 5 km away from each other, as a part of the Split–Solin public sewage system.

The submarine outfall at Location 1 discharges treated wastewater from the Katalinića brig wastewater treatment plant (WWTP) via a \varnothing 800 mm pipeline, ending with two diffuser sections 1.3 km away from the shore at a 43 m depth. The Katalinića brig WWTP is designed for the mechanical treatment of both municipal and rainfall-runoff wastewater, with a capacity of 122.000 population equivalents (PE) and an average flow rate equal to 35.000 m³/day approximately.

Water 2021, 13, 2016 3 of 17

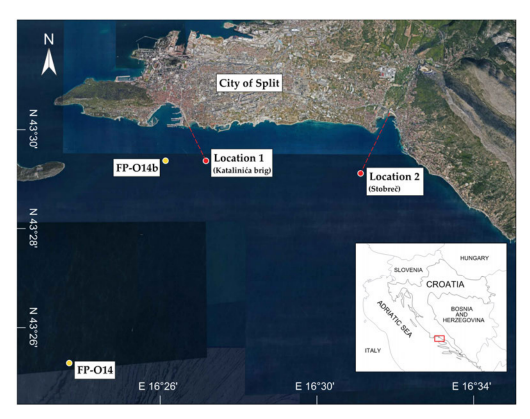


Figure 1. Locations of marine wastewater outfalls and national monitoring program stations.

The marine outfall at Location 2 (Stobreč) discharges the wastewater coming from the Stupe WWTP, where the wastewater is mechanically treated, along with grit and grease removal. The Stupe WWTP facility is designed for municipal wastewater only, with a capacity of 138.000 PE and an average flow rate equal to 30.000 m³/day. The length of the submarine outfall \varnothing 900 mm pipeline is 2.76 km, with a 200 m long diffuser section ending at a 37 m depth.

Additionally, the monitoring of different physical and chemical parameters is performed several times a year, at stations FP-O14 and FP-O14b, as shown in Figure 1, as a part of the national monitoring program.

To understand the background of the wastewater being discharged into the seawater, we provide data (Table 1) about the seasonal flowrate and available water effluent parameters for two WWTPs.

Table 1. Available data for two WWTP locations, considering the average discharge (m^3 /day) and different wastewater effluent parameter concentrations (mg/L), presented as the mean \pm standard deviation.

Location/ Parameter	Q _{SUMMER} (m³/Day)	Q _{WINTER} (m ³ /Day)	COD (mg/L)	BOD (mg/L)	TSS (mg/L)	TN (mg/L)	TP (mg/L)
Stupe	30.000	26.000	372 ± 235	199 ± 113	89 ± 59	52 ± 15	6 ± 1.5
Katalinića brig	38.000	34.000	246 ± 107	135 ± 58	69 ± 38	32 ± 5	4 ± 2

2.2. Sampling Methodology

This study was performed during the implementation of the Interreg Italy–Croatia project AdSWiM (Managed use of treated urban wastewater for the quality of the Adriatic Sea). The measurements were performed with the CTD fast profiling probe (The Sea & Sun Technology) and C3 fluorometer during sampling sessions conducted with a boat (Nautica 600) every month from February to September 2020 (with the exception of March and August due to the COVID restrictions).

Water **2021**, 13, 2016 4 of 17

CTD fast profiling probe, with an additional sensor for dissolved oxygen (model: 48M), was used for temperature, salinity, density anomaly, and dissolved oxygen measurements. The temperature is measured by a classical resistance thermometer using a platinum resistor, while the salinity is calculated from the seawater conductivity using a 7-poll cell sensor. The density anomalies were calculated automatically by the CTD/DO probe internal software using the international thermodynamic equation of seawater-2010 [52]. The dissolved oxygen sensor works on the principle of a Clark electrode. The depth is calculated from the pressure, measured by a strain gauge, and is converted to depth as 1dbar = 1 m. The turbidity, chlorophyll, and colored dissolved organic matter (CDOM) were measured using a C3 Submersible fluorometer (Turner Design). The chlorophyll measurements were performed with a blue mercury lamp, with a peak emission at 460 nm, and the fluorescence was collected at 680 nm. The CDOM was measured using a UV LED (CWL: 365 nm), with a peak emission at 350 nm and fluorescence collection at 450 nm. The turbidity was measured with the IR lamp, with a peak emission at 850 nm, and the collection of scattered light at 90°. Since the relationship between phytoplankton biomass and chlorophyll fluorescence is highly variable and depends on the physiological state and community composition [53–55], in this study, we focused on the general distribution of chlorophyll profiles in the water column concerning the stratification and effluent discharge.

The fecal pollution was estimated by the level of FIB. The FIB concentration was assessed by the membrane filtration method and the enumeration of fecal coliforms (*E. coli*) [56,57] and intestinal enterococci (ENT) [58], as indicated in the Croatian legislation (Regulation on the Quality of Marine Bathing Waters; OG 73/08). The number of viable heterotrophic bacteria was determined using the spread plate method. An automated Protos3 colony counting system (Synbiosis, UK) was used to determine the bacteriological concentration. Sampling was conducted from the boat using a Niskin sampler. The samples were collected in sterile bottles, protected from light in cool boxes, and transported to the microbiology laboratory of the Faculty of Science in Split to be processed within 4 h.

The chlorophyll concentration data at locations FP-O14 and FP-O14b were collected during the regular monitoring performed by the water management agency (Croatia Waters). The chlorophyll a concentrations were determined by the laboratory fluorometric method, with filtration through Whatman GF/C glass filters, at 0, 5, 10, 20, and 50 m depths at the FP-O14 location and at 0, 5, 10, and 43 m depths at FP-O14b.

2.3. Data Analysis

The data collection was performed by the vertical profiling of the water column down to 40 m and 35 m depths at location 1 and location 2, respectively. A moving average filter was applied to raw data sampled at a maximum spatial resolution of 3 cm to achieve spatial uniformity. The data were smoothed by a moving average filter, with a 20 cm window.

The vertical distributions of the different layers were examined by calculating the vertical gradients of the density anomalies and derived buoyancy frequency. The buoyancy frequency or Brunt-Väisälä frequency (N) represents the strength of the density stratification and, consequently, the stability of the water column. It is defined as:

$$N = \sqrt{\frac{g}{\rho_0} \frac{d\rho}{dz}} \tag{1}$$

where g is the gravitational acceleration, ρ_0 is the averaged density of the whole water profile, and $d\rho/dz$ is the water density gradient at each sampling step. To identify events when stratification occurred, data points are selected where N^2 exceeds $0.001 \, s^{-2}$ as possible pycnocline candidates [59]. The data points are then aggregated if the distance between them was less than 0.8 m, forming a possible pycnocline layer. Layers thinner than 0.8 m are neglected to exclude the impact of anomalous extremes. Aggregated layers are classified as a pycnocline and their profiles as stratified, which potentially aids in the creation of chlorophyll layers. To support the analysis of forming chlorophyll layers under

Water 2021, 13, 2016 5 of 17

the influence of marine wastewater discharge, the vertical profiles of dissolved oxygen, turbidity, CDOM, temperature, and salinity are also examined.

3. Results

The main results from six sampling cruises are presented in Figures 2 and 3 for location 1 and location 2, respectively. Each Figure contains six vertical profiles, corresponding to the sampling month with the set of parameters measured. Analysis of the buoyancy frequency showed that stratification occurred in May, June, July, and September for both locations. In February and April, the winter and early spring, lower temperatures induced vertical mixing in the water column. As a result of the strong sea surface warming, the shallowest stratification occurred in July, while the deepest occurred in September. At both locations 1 and 2, the temperature and salinity followed their seasonal pattern, with low gradients in winter and steeper gradients during stratified periods.

3.1. Location 1

3.1.1. February 2020

During the February sampling cruise, the winter mixing of the water column, induced by wind and low temperatures, allowed the upwelling of the bottom seawater rich with nutrients, causing the rise of chlorophyll concentrations from the bottom to the upper layers of the water column (Figure 2, February). The February vertical profiles shown in Figure 2 also depict a slight increase in the turbidity and CDOM around a 25 m depth, probably due to the phytoplankton activity, which can contribute to the organic matter by excreting photosynthetic products or by lysing [60]. The homogeneous vertical dissolved oxygen profile indicates a high oxygen enrichment throughout the water column, mostly as a consequence of the higher oxygen production and water column aeration during the winter weather conditions.

3.1.2. April 2020

The April vertical profiles shown in Figure 2 show a higher concentration of chlorophyll at the bottom, along with higher values of CDOM and turbidity. The observed dip in the dissolved oxygen levels near the bottom indicates the presence of a wastewater plume. To support this hypothesis, the observed high level of fecal pollution is presented in Figure 4. The high levels of *E. coli* and ENT support the dissolved oxygen dip, as bacteria consume oxygen from seawater during the process of decomposition of organic matter from the wastewater plume. The homogeneous vertical profile of dissolved oxygen in the upper part of the water column indicates that the oxygen production exceeds the consumption in that part of the water column.

3.1.3. May 2020

The stratification of the water column started in May. The chlorophyll vertical profile, shown in Figure 2, has no gradients in the upper layer, with a steady increase below the forming stratification point. The increase in the chlorophyll concentration is followed by an increase in CDOM and turbidity. Since the dissolved oxygen profile was homogeneous, we presume that the phytoplankton production of oxygen dominated over the consumption, without any observed effluent effect.

Water 2021, 13, 2016 6 of 17

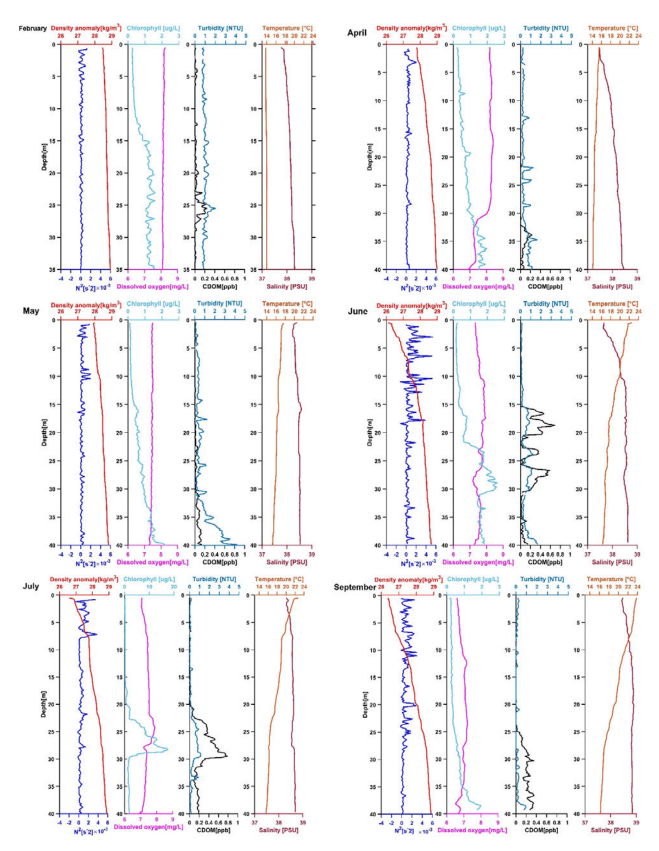


Figure 2. Vertical profiles sampled during 2020 at location 1. The measured parameters are density anomaly (red line), buoyancy frequency (blue line), chlorophyll (light blue line), dissolved oxygen (magenta line), turbidity (green line), CDOM (black line), temperature (orange line), and salinity (brown line).

Water 2021, 13, 2016 7 of 17

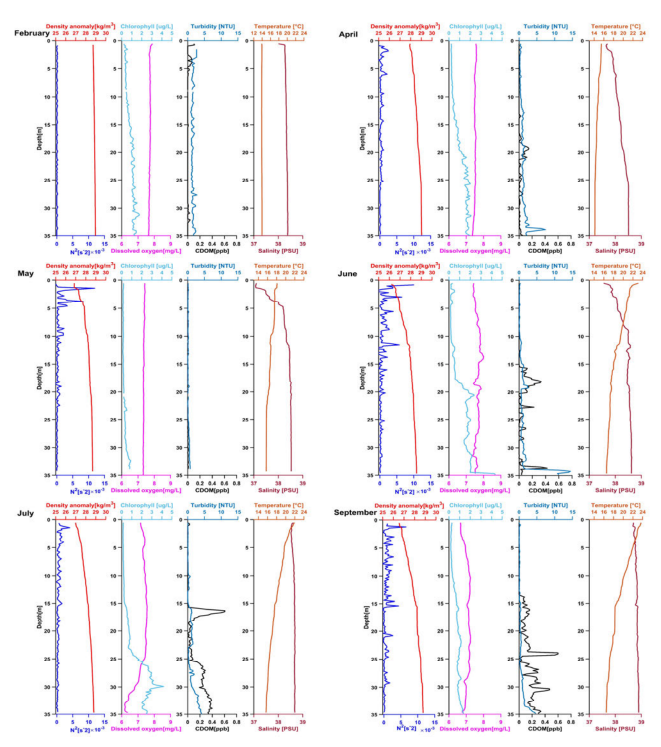


Figure 3. Vertical profiles sampled during 2020 at location 2. The measured parameters are density anomaly (red line), buoyancy frequency (blue line), chlorophyll (light blue line), dissolved oxygen (magenta line), turbidity (green line), CDOM (black line), temperature (orange line), and salinity (brown line).

Water 2021, 13, 2016 8 of 17

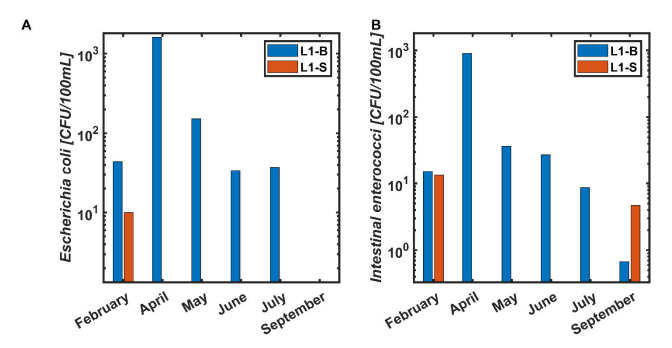


Figure 4. Microbial water quality ((**A**) *Escherichia coli*, (**B**) Intestinal enterococci) at location 1. L1-B refers to location 1 bottom (blue), while L1-S refers to location 1 surface location (orange).

3.1.4. June 2020

During the June sampling cruise (Figure 2), we observe a broad chlorophyll peak with three plateaus, starting below a depth of 15 m. The chlorophyll peaks coincide with the CDOM and turbidity peaks, indicating an effluent plume layer. Consequently, the phytoplankton biomass increases, further contributing to the organic matter concentration and turbidity rise. This increase is matched by a decrease in salinity and dissolved oxygen. The oxygen decrease is a consequence of the bacterial oxidation of organic matter, which is flushed into the seawater by the marine outfall discharge. In the June sampling cruise, the stratification is much steeper, compared to May, and it occurs at a depth of around 18 m.

3.1.5. July 2020

The July vertical profiles shown in Figure 2 show that the chlorophyll vertical profile is not comprised of several plateaus, like the one in June, but has a steep slope on the deeper side and two shoulders on the shallow side. Furthermore, it is present much deeper than the stratification layer, which occurs at a depth of 7 m. The increase in chlorophyll is matched by an increase in CDOM and turbidity values and a decrease in dissolved oxygen and salinity. The observed peaks of different parameters during the July cruise indicate the presence of an effluent plume below the pycnocline. The dissolved oxygen increases down to the 24 m depth and then begins to decline near the chlorophyll peak.

3.1.6. September 2020

The September vertical profiles shown in Figure 2 show the pycnocline presence down to the 22–23 m depth, which was the deepest pycnocline observed during our cruises. Below the pycnocline, at a 25 m depth, an increase in chlorophyll, CDOM, and turbidity is observed, together with a small decrease in dissolved oxygen at the same depth. Near the bottom of the water column, the oxidation of organic matter reduced the dissolved oxygen concentration, indicating that more oxygen was being used than produced. In this case, the deep pycnocline probably submerged the effluent plume, keeping it near the bottom of the water column.

Water 2021, 13, 2016 9 of 17

3.2. Location 2

3.2.1. February 2020

During the February sampling cruise, a similar vertical chlorophyll profile occurred at location 2 as that at location 1. The February chlorophyll profile shown in Figure 3 shows a rise in the chlorophyll concentrations from the bottom to the top of the water column. The homogeneous vertical turbidity, CDOM, salinity, and dissolved oxygen profiles indicate the water column mixing. Furthermore, the elevated levels of FIB (Figure 5) at the surface of the water column indicate the effluent rising to the surface of the water column due to the mixing process under the unstratified water column conditions.

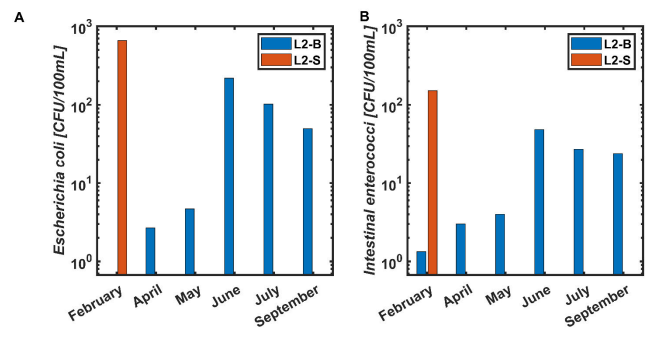


Figure 5. Microbial water quality ((**A**) *Escherichia coli*, (**B**) Intestinal enterococci) at location 2. L2-B refers to location 2 bottom (blue), while L2-S refers to location 2 surface location (orange).

3.2.2. April 2020

The April vertical profiles shown in Figure 3 indicate that the mixing of the water column creates the upwelling of the bottom water, causing a similar vertical chlorophyll profile to the one observed in February. The observed peaks in CDOM and turbidity, within the mixing process, are due to the available nutrients for primary production, resulting in the phytoplankton activity. The homogeneous dissolved oxygen vertical profile indicates a high oxygen enrichment throughout the water column, reflecting surplus phytoplankton production and an aerated water column due to the lack of stratification.

3.2.3. May 2020

Stratification of the water column at location 2 was first observed in May 2020. The observed chlorophyll, CDOM, and turbidity profiles in May 2020 (Figure 3) reveals no significant concentration and/or gradients. Such situations are mostly a consequence of the complex physical, biological, and chemical interactions, together with hydrodynamic processes, particularly due to the fact that the wastewater effluent is only periodically discharged into the sea. However, the May chlorophyll profile, shown in Figure 3, shows an increase in values near the bottom of the water column. The increase in the chlorophyll concentration is matched by a small increase in CDOM and turbidity near the bottom, without a decline in the dissolved oxygen vertical profile. Since the dissolved oxygen profile was homogeneous, we presume that the phytoplankton production of oxygen exceeded the consumption by bacteria.

Water **2021**, 13, 2016 10 of 17

3.2.4. June 2020

The June vertical profiles presented in Figure 3 show the pycnocline at a 14 m depth, after which an increase in chlorophyll, CDOM, and turbidity is observed. Higher chlorophyll concentrations below a 15 m depth, in conjunction with the CDOM and turbidity increase, indicate the presence of an effluent plume layer accumulated below the pycnocline. The presence of organic matter, together with a higher phytoplankton biomass, leads to an increase in turbidity. Furthermore, the input of organic material contributes to the reduction of available oxygen through bacteria oxidation processes, which is supported by the shape of the vertical dissolved oxygen profile. The reduced salinity at 15 to 20 m depths supports the conclusion that these changes are a direct consequence of the wastewater plume. In comparison to the other months, in June 2020 the highest chlorophyll, CDOM, and turbidity values are measured at the bottom of the water column. The concentration of 220 CFU/100 mL of *E. coli* indicates the presence of an effluent, in agreement with other observed parameters.

3.2.5. July 2020

During the July sampling cruise (Figure 3, July), pycnocline occurred at a very shallow depth of 4 m. Increased values of chlorophyll, CDOM, and turbidity are observed from 15 m to 35 m depths. The observed peaks of chlorophyll, turbidity, and CDOM, with a homogeneously dissolved oxygen profile, down to the 25 m depth, indicate the presence of phytoplankton biomass, producing dissolved oxygen, which prevailed over the consumption. A sharp CDOM peak at the 15 m depth was due to the phytoplankton activity, as no dissolved oxygen decrease was observed at that depth. The decrease in the dissolved oxygen below the 25 m depth indicates that the plume effluent had a more significant effect in this layer. The oxygen consumption during the organic matter decomposition at this depth prevailed over the production, causing an increase of CDOM and turbidity in the bottom layer. The observed concentration of 103 CFU/100 mL for *E. coli* indicates an intrusion of fecal material by the marine outfall. Thus, the July profile showed the characteristics of two competing processes. One is a higher oxygen production from the phytoplankton activity and the other is a higher oxygen consumption from the oxidation of organic material entering the marine system by the wastewater discharge.

3.2.6. September 2020

September vertical profiles in Figure 3 show a homogeneous chlorophyll profile within the upper layer up to the pycnocline at a 15 m depth. Below the stratification, the chlorophyll concentration is increasing towards the bottom of the water column. The organic matter peaks were observed below the pycnocline, while the turbidity increases only near the bottom. As the vertical dissolved oxygen profile shows little variability, we presume that the organic matter reflects phytoplankton activity, indicating that the production dominated over the consumption.

4. Discussion

The majority of the vertical chlorophyll profile studies conducted in coastal region and estuaries are based on laboratory measurements or satellite techniques, lacking direct in situ measurements, especially the influence of marine outfalls. Therefore, this study aimed to identify the chlorophyll vertical profiles under the influence of the water column stratification status and the presence of an effluent plume. Based on the measurement results, we classify the chlorophyll profiles into three types (Table 2). The profiles observed in February and April 2020, with the observed rise of the chlorophyll concentrations from the bottom to the top of the water column, are classified as the Upper Chlorophyll Profile (UCP). The other two profile types are divided between the sampling sessions when stratification of the water column was observed, and they differ by the presence of an effluent plume. In May at both locations and in September at location 2, the vertical chlorophyll profile was characterized by a higher concentration near the bottom, without

Water 2021, 13, 2016 11 of 17

a decrease in dissolved oxygen. These profiles are classified as the Bottom Chlorophyll Profile (BCP). The third profile type is characterized by peaks of chlorophyll coinciding with CDOM and turbidity peaks and a matching decrease in salinity and dissolved oxygen, which is a consequence of an effluent plume intrusion. These profiles are classified as the Effluent Chlorophyll Profile (ECP). A summary of this classification is given in Table 2.

		Location 1		Location 2				
Month	Stratification	Effluent	Type	Stratification	Effluent	Type		
February	Х	Х	UCP	Х	Х	UCP		
April	X		UCP-ECP	X	Х	UCP		
May		Х	ВСР	\checkmark	Х	ВСР		
June			ECP		√	ECP		

ECP

ECP

ECP

BCP

χ

Table 2. Vertical profile classification summary.

July

September

In unstratified conditions, observed in February and April 2020, at both locations, the upwelling of the bottom, nutrient-rich waters induced a rise in the chlorophyll concentrations from the bottom to the top of the water column, which we classified as UCP. This rise is supported by the CDOM and turbidity vertical profiles, which show only small variations along the vertical column. The bottom water upwelling also resulted in a higher FIB level at the surface, indicating a rise in fecal pollution due to marine outfalls. In unstratified conditions, the dissolved oxygen vertical profiles were homogeneous. This is due to vertical mixing, which allows air-sea gas exchange with the bottom waters and the influence of external forces, such as wind [61]. An exception occurred in April 2020 at location 1 due to an effluent plume intrusion, which decreased the dissolved oxygen values. These conditions induce the characteristics of the chlorophyll profile of both the ECP and UCP (Figure 2). The oxygenation of the water column down to a 30 m depth was due to the primary production of phytoplankton, but it was also due to the unstratified water column, which caused vertical mixing and aeration of the entire water column. At the bottom of the water column, the dissolved oxygen concentration and chlorophyll are inversely correlated, suggesting that the consumption of oxygen dominates over its production. Similar observations have been made in many estuaries with nutrient inputs from rivers [62–67]. Marine outfalls discharge nutrients into the sea, resulting in similar biological processes occurring as those when rivers discharge nutrients into the estuary.

From May to September, stratification occurred at both locations, and the observed profiles are classified as BCP (May 2020, September 2020) and ECP (June 2020, July 2020, September 2020) (Table 2). BCPs are characterized by a higher chlorophyll concentration near the bottom of the water column. This increase is likely due to the phytoplankton activity and resuspension of the bottom sediment, where dead plankton and other sources of organic matter fertilizer, unconsumed food, and fecal matter settle. In May 2020, despite the water column stratification, the vertical distribution of dissolved oxygen was homogeneous, similar to a phenomenon observed in some estuaries, where, under stratified conditions and higher temperatures, hypoxia occurred [68–70]. In our case, hypoxia was never observed, even in summer, but a higher dissolved oxygen variability was present.

To confirm the occurrence of UCP and BCP types in the absence of an effluent plume, we used data from measurements conducted by the Croatian water management agency in the period from 2012–2019, sampled at two locations (FP-O14 and FP-O14b). Both stations are used to represent coastal seawater unaffected by marine outfalls. At both of these locations, during months when no stratification is present, the vertical chlorophyll profile shows upwelling, indicating the existence of UCP (Figures 6 and 7). Under stratified conditions, narrower ranges of the chlorophyll concentrations are observed in the upper part, with the highest values measured at the bottom, which are classified as

Water 2021, 13, 2016 12 of 17

BCP (Figures 6 and 7). These data show the existence of UCP and BCP profiles in coastal, unpolluted waters, which follow a seasonal cycle of sea warming and the formation of stratified conditions, and differentiate the naturally occurring chlorophyll profiles from the ones induced by anthropogenic influence.

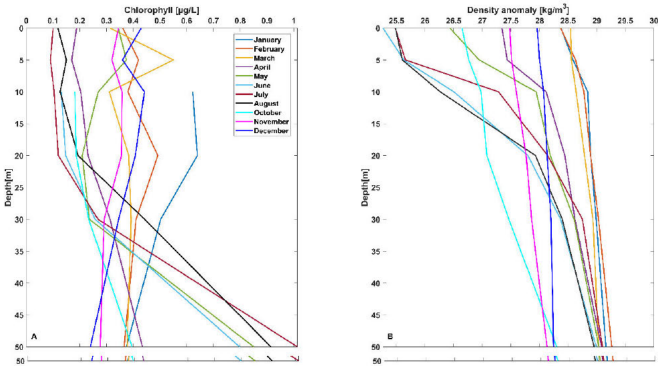


Figure 6. Geometric mean vertical chlorophyll (A) and density anomaly (B) concentrations observed during the 2012–2019 period at the measuring station, FP-O14.

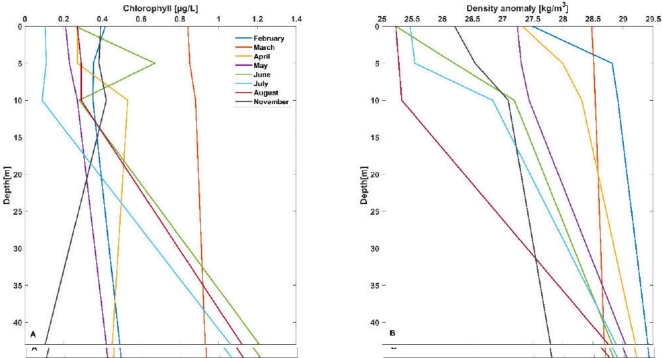


Figure 7. Geometric mean vertical chlorophyll (A) and density anomaly (B) concentrations observed during the 2012–2019 period at the measuring station, FP-O14b.

Water 2021, 13, 2016 13 of 17

The vertical profiles of CDOM, turbidity, dissolved oxygen, and salinity further highlight the difference between ECP and BCP. The nutrient enrichment by marine outfalls induces phytoplankton biomass growth and increases the chlorophyll concentration. The highest chlorophyll concentration observed was at location 1 in July 2020. Such a surge in the chlorophyll concentration could be due to the accumulation of living algae, supported by raw sewage, containing higher amounts of nutrients, especially nitrogen and phosphorus. Besides nutrients, the marine outfalls also discharge organic matter into the coastal waters, resulting in increased organic matter concentrations and turbidity. The increased turbidity levels are the result of the input of organic matter from the wastewater discharge, creating a good condition for the presence of fecal bacteria. The effluent intrusion creates conditions where the oxygen consumption, during the bacterial oxidation of organic matter and decomposition of dead algae and plants, exceeds the production of dissolved oxygen by photosynthesis. The observed decrease in the dissolved oxygen profiles furthermore separates the BCP and ECP types. The gradient of dissolved oxygen with depth (higher concentration above the pycnocline) indicates that stratification may play a role in lowering the oxygen concentration by reducing the gas exchange between the air and sea surface layer [61]. While stratification can influence the dissolved oxygen profile, the homogeneously dissolved oxygen profiles observed in May suggest that the highest variability of dissolved oxygen concentration is due to the influence of a wastewater effluent plume and not stratification. The variations in salinity also supported our hypothesis about the nature of ECP. Similar changes in salinity were previously observed for other submarine outfall sites and provide further evidence of a freshwater input through the outfall system [71–74].

Due to the fact that both WWTPs are discharging a municipal wastewater effluent, the seasonal variations are due to the variations in the residential population of the Split municipal area, which increases during the summer months. Consequently, such changes can influence the wastewater volume. Thus, the summer discharges are approximately 12% higher than those during the winter period, as presented in Table 1. This is also shown in Figures 2 and 3 in terms of an increased concentration and/or gradients of measured parameters. While the discharged wastewater and the seawater start mixing quickly and reaching equilibrium with the surrounding seawater, the salinity profiles may serve as additional information, aiding in distinguishing the chlorophyll profiles. During the observed stratification, the effluent plume was held below the pycnocline, which is generally ecologically favorable, as it maintains a higher quality of the surface waters.

In addition, the circulation in the wider area surrounding the two studied outfalls has a typical barocline current pattern, with relatively larger surface layer velocities due to the regular wind intensities. Furthermore, during the unstratified conditions, the bottom upwelling is intensified, creating the potential for effluent plume surfacing. In such cases, it is expected that surface currents will further contribute to the mixing and spreading of chlorophyl.

5. Conclusions

To study the wastewater impact on the chlorophyll vertical distribution, measurements were performed in the eastern central Adriatic Sea at two locations of marine outfalls, near the city of Split in Croatia. We describe the three vertical chlorophyll profile types, based on the stratification conditions and the effluent plume presence. The water column mixing induced the upwelling of the bottom water, generating UCP and enabling the rise of the effluent towards the surface. In the stratified conditions, categorized as the ECP type, the pycnocline prevents the plume from rising, keeping it in the lower part of the water column, while the BCP type of the profile represents a case when the water column is stratified, but no effluent plume is present. Our categorization of the different vertical chlorophyll profiles is supported by data collected by the Croatian water agency at the locations unaffected by the marine outfalls. All three observed chlorophyll profile types are the consequence of the interplay between the physical, biological, and chemical processes. To distinguish these processes and confidently identify the presence of the effluent plume

Water 2021, 13, 2016 14 of 17

we use the FIB measurements, along with dissolved oxygen gradients, to examine the conditions when the oxygen consumption exceeds the production, indicating organic matter oxidation. This, together with the salinity, CDOM, and turbidity measurements, allows us to separate the wastewater influence from the natural processes. We confirm the importance of the stratification in submerging the effluent below the pycnocline, which is highlighted in February 2020, when column mixing allowed the FIB to rise to the surface of the water column. To our knowledge, this is a first study involving in situ vertical measurements coupled with the submarine outfall, analyzing its influence on the vertical chlorophyll distribution.

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3.2. Microbiome and antibiotic resistance profiling in submarine effluent-receiving coastal waters in Croatia

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Wastewater treatment plant (WWTP) effluents are an important source of both commensal and pathogenic bacteria and their antibiotic resistance genes (ARGs) into water bodies. This research, for the first time, analyse the impact of partially treated submarine effluents on the water quality of two distinct water columns. Its aim is to comprehensively investigate and understand the water column microbiome and resistome using 16S rRNA amplicon sequencing. This study highlights the adverse effects of submarine effluent discharge, which enriches bacterial communities with non-native taxa carrying various ARGs, potentially increasing the spread of antibiotic resistance in nature.

Analysis of the coastal marine microbiome and resistome contributes to a better understanding of the impact of submarine outfalls on water column quality, in terms of the presence of different allochthonous bacterial species and human pathogens as potential carriers of a variety of ARGs.

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Microbiome and antibiotic resistance profiling in submarine effluent-receiving coastal waters in Croatia[★]

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ABSTRACT

Wastewater treatment plant (WWTP) effluents are pointed as hotspots for the introduction of both commensal and pathogenic bacteria as well as their antibiotic resistance genes (ARGs) in receiving water bodies. For the first time, the effect of partially treated submarine effluents was explored at the bottom and surface of the water column to provide a comprehensive overview of the structure of the microbiome and associated AR, and to assess environmental factors leading to their alteration. Seawater samples were collected over a 5-month period from submarine outfalls in central Adriatic Sea, Croatia. 16S rRNA amplicon sequencing was used to establish taxonomic and resistome profiles of the bacterial communities. The community differences observed between the two discharge areas, especially in the abundance of Proteobacteria and Firmicutes, could be due to the origin of wastewaters treated in WWTPs and the limiting environmental conditions such as temperature and nutrients. PICRUSt2 analysis inferred the total content of ARGs in the studied microbiomes and showed the highest abundance of resistance genes encoding multidrug efflux pumps, such as MexAB-OprM, AcrEF-TolC and MdtEF-TolC, followed by the modified peptidoglycan precursors, transporter genes encoding tetracycline, macrolide and phenical resistance, and the bla operon conferring β -lactam resistance. A number of pathogenic genera introduced by effluents, including Acinetobacter, Arcobacter, Bacteroides, Escherichia-Shigella, Klebsiella, Pseudomonas, and Salmonella, were predicted to account for the majority of efflux pump-driven multidrug resistance, while Acinetobacter, Salmonella, Bacteroides and Pseudomonas were also shown to be the predominant carriers of nonefflux ARGs conferring resistance to most of nine antibiotic classes. Taken together, we evidenced the negative impact of submarine discharges of treated effluents via alteration of physico-chemical characteristics of the water column and enrichment of bacterial community with nonindigenous taxa carrying an arsenal of ARGs, which could contribute to the further propagation of the AR in the natural environment.

1. Introduction

In many coastal areas, submarine sewage outfalls are used to dispose of partially or fully treated wastewater. Although such disposal is environmentally acceptable, it can also be a significant factor of pollutants' influent and further spread to the pristine marine environment. Namely, the effluents commonly contain microbiological and chemical pollutants (Lapointe et al., 2011; Roth et al., 2016; Vidal-Dorsch et al., 2012; Wang et al., 2014; Yang, 2000) which can impact the biodiversity of coastal marine microbiota (Mittal et al., 2019; Zheng

Abbreviations: Wastewater treatment plant, (WWTP); antibiotic resistance gene, (ARG); antibiotic resistance, (AR); antibiotic-resistant bacteria, (ARB); fecal indicator bacteria, (FIB); temperature, (T); salinity, (SAL); density anomaly, (SIGMA); dissolved oxygen, (DO); chlorophyll a, (CHL); turbidity, (TUR); nitrite nitrogen, (NO2–N); nitrate nitrogen, (NO3–N); ammonia nitrogen, (NH3–N); total nitrogen, (TN); total phosphate, (TP); silicate ions, (SiO₄³); colored dissolved organic matter, (CDOM); imipenem, (IMP); colistin, (CL); operational taxonomic units, (OTUs); oxacillinase, (OXA); (CAMP), cationic antimicrobial peptide.

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et al., 2021; Zheng et al., 2019) and alter environmental parameters, including pH (Shen et al., 2013; Yannarell and Triplett, 2005), salinity (Hou et al., 2017), temperature (Zheng et al., 2019) and level of nutrients (Lu et al., 2020; Wang et al., 2018).

In addition to microbial diversity, significant changes in gene abundances, especially virulence and antibiotic resistance genes (ARGs) have been identified in anthropogenically impacted marine and freshwater environments (Bondarczuk and Piotrowska-Seget, 2019; Fresia et al., 2018). Antibiotic resistance (AR) is a particularly challenging problem in these ecosystems, being the final recipient of a municipal, hospital, industrial, agricultural, and aquaculture wastewaters (Rizzo et al., 2013). In this regard, wastewater treatment plants (WWTPs) are considered as major sources of ARGs and antibiotic-resistant bacteria (ARB) with sewage effluents acting as hot spots for their further dissemination to the environment (Griffin et al., 2020; Kotlarska et al., 2015; Proia et al., 2018; Su et al., 2020). WWTP effluents in coastal waters in China were found to be enriched with genes encoding for resistance to multiple classes of antibiotics (Chen et al., 2020; Su et al., 2020) that positively correlated with nitrate, active phosphorus and pH (Chen et al., 2020).

Molecular biology and computational biology tools are now being widely used for investigating microbial communities (Dinsdale et al., 2008; Fresia et al., 2018), while PICRUSt2 (Douglas et al., 2020) annotates the functional features, among which the ARGs (Šamanić et al., 2021), expanding the possibility for a comprehensive study of the vast environmental resistance reservoirs associated to wastewaters (Cai et al., 2014; García-Aljaro et al., 2019; Narciso-da-Rocha et al., 2018; Port et al., 2012).

Importantly, a better understanding of the microbial wastewater communities can provide an important framework for the more effective design and operation of novel and robust wastewater treatment systems. Based on the available literature, this study is the first to provide a comprehensive analysis of the bottom and the surface of the water body receiving the effluents from the submarine outfalls with respect to the microbiome diversity and AR prediction, in combination with the vertical profiling of the water column.

The objectives of this study were to (i) determine the microbial communities at the bottom and at the surface of the water column adjacent to the effluent plume discharge; (ii) determine if there is a difference in microbial composition between the two submarine outfall discharges, particularly in regards to human pathogenic bacteria (iii) functionally predict the repertoire of ARGs in bacterial communities based on taxonomic outlining of effluent and surface water samples (iv) assess the potential of pathogenic bacterial genera as reservoirs of AR in analyzed marine microbiomes.

2. Materials and methods

2.1. Description of the study sites

This study focused on the highly urbanized central part of the eastern Adriatic coast in Croatia (Fig. S1.), covering the municipalities of Split, Solin and Stobreč with a population of >200,000. Two WWTPs, Katalinića brig and Stupe-Stobreč, are processing wastewaters from the wider Split area and discharging it through submarine outlets into the Brač and Split Channels. While Stobreč WWTP processes only municipal wastewater, Katalinića brig WWTP combines municipal and rainfallrunoff wastewater. Both WWTPs have only mechanical treatments, including coarse and fine screens, while Stobreč WWTP also provides oil and sand removal. The Stobreč WWTP is designed for municipal wastewater only, with capacity of 138.000 population equivalent (PE) and an average flow rate equal to 30,000 m³/day, while Katalinića brig WWTP is designed for 122,000 PE with an averaged flow rate of 35,000 m³/day. Another important feature of the Katalinića Brig WWTP is that it also collects the wastewater from University Hospital Centre Split, a leading medical center in southern Croatia with 1400 beds. After

treatment, the Stobreč WWTP is discharging the effluent by gravity through the submarine outfall of 2.76 km in length with a 200 m diffuser section at depth of 37 m (43°28′53.6 "N 16°31′04.3 "E). The Katalinića Brig WWTP is discharging the treated wastewater by pumping through the submarine outfall of 1.3 km in length at depth of 42 m (43°29′22.7 "N 16°27′11.2 "E), respectively.

2.2. Sampling

Samplings were conducted once a month during campaigns in February, May, June, July, and September 2020. At each site, three 500-mL water samples were taken consecutively as a representative grab sample set for further analysis. At Katalinića Brig, seawater was collected at the surface (abbreviated as KS) and at the sea bottom adjacent to the submarine outfall (abbreviated as KB). Same approach was carried out at the Stupe submarine outfall, abbreviated as SS (Stobreč surface) and SB (Stobreč bottom). Sampling was done with the boat (Nautica 600, Croatia) using a Niskin sampler. Grab samples were collected in sterile bottles, protected from light in cool boxes, transported to the laboratory within 4h at 4 °C and processed immediately. Seawater samples for chemical analyses were transported within 8h at 4 °C to the laboratory of Public Health Institute in Zadar, Croatia, and then stored at 80 °C for further analyses.

2.3. Physico-chemical analysis

Physical parameters were measured during field samplings. Sea & Sun Technology CTD fast profiling probe with an additional sensor for dissolved oxygen (DO) (model: 48M) was used for temperature (T), salinity (SAL), and density anomaly (SIGMA). Data collection was performed by downward profiling of water column up to 40m depth at Katalinića Brig station and up to 35m depth at Stupe station. SIGMA was calculated automatically by CTD/DO probe software using an international thermodynamic equation of seawater-2010 (IOC et al., 2010). Chlorophyll a (CHL) was determined by a standard method for in vivo determination of CHL in marine waters (Arar and Collins, 1997). Turbidity (TUR) (ISO 7027-1), nitrite nitrogen (NO2-N) and nitrate nitrogen (NO3-N) (Baird and Bridgewater, 2017), ammonia nitrogen (NH3-N) (Ivancic and Degobbis, 1984), total nitrogen (TN) (ISO 12260), total phosphate (TP) (ISO 6878), and silicate ions (SiO 4³⁻) (Grasshoff et al., 1998) were analyzed by laboratory standard methods, listed for each parameter in brackets. Colored dissolved organic matter (CDOM) was measured by a C3 Submersible fluorometer (Turner Design), using a UV LED, with peak emission at 350 nm and fluorescence collection at

2.4. Bacterial enumeration

The level of fecal pollution was assessed by the membrane filtration method and enumeration of FIB, including the fecal coliforms (E. coli) (ISO 9308-1; Jozić et al., 2018) and intestinal enterococci (ISO 7899-2), as indicated in the Croatian legislation (Regulation on the Quality of Marine Bathing Waters; OG 73/08). The number of viable heterotrophic bacteria was determined using the spread plate method. Aliquots of 0.1 mL were plated on Marine agar (MA; BD Difco, USA) and the plates were incubated at room temperature for up to one week to ensure the growth of the slow-growing bacteria. To determine the proportion (in %) of imipenem (IMP)- and colistin (CL)-resistant bacteria, aliquots from 1 to 50 mL of each water sample were filtered through a 0.22 μm pore-size MCE membranes and the filters were placed on MA (BD Difco) supplemented with IMP (4 $\mu g/mL$) or CL (8 $\mu g/mL$) and without antibiotics. The two antibiotics were chosen regarding their global relevance in treatment of infections caused by emerging multidrug- and carbapenem-resistant pathogens (WHO, 2017). The concentration of IMP was selected according to international guidelines (CLSI, 2020). On the other hand, a two-fold higher concentration of CL than the MIC breakpoint was used due to poor diffusion of this drug in solid media (Turlej-Rogacka et al., 2018). An automated Protos3 colony counting system (Synbiosis, UK) was used in all experiments. All plate counts were performed in triplicate and the number of bacteria was expressed as mean count (CFU/100 mL) \pm standard deviation (SD).

2.5. DNA extraction and illumina-based 16S rRNA amplification

One liter of each water sample was filtered through 0.22 µm poresize MCE membrane filters and DNeasy PowerWater Kit (Qiagen, Germany) was used to extract genomic DNA. The concentration and quality of DNA were analyzed using a NanoDrop® Spectrophotometer 1000 (Thermo Scientific, USA). DNA samples were sent to a Novogene Europe (Cambridge, UK) for 16S amplicon metagenome sequencing. A 470 bp hypervariable region V3-V4 of the 16S rRNA gene was amplified using standard primers 341F and 806R and Phusion® High-Fidelity PCR Master Mix (New England Biolabs, UK). PCR products were purified, libraries generated using NEBNext® UltraTM DNA Library Prep Kit (Illumina, UK) and analyzed on a NovaSeq platform (Illumina). 250 bp paired-end raw reads were merged using FLASH (V1.2.7) and highquality clean tags were obtained after quality filtering (Caporaso et al., 2010) using QIIME (V1.7.0). Sequences were analyzed using Uparse (v7.0.1001) and those with \geq 97% similarity were assigned to the same Operational Taxonomic Units (OTUs). Representative sequences were compared with SILVA Database (Quast et al., 2013) for species annotation at each taxonomic rank (threshold: 0.8-1) using MUSCLE (White et al., 2009).

2.6. PICRUSt functional prediction

The PICRUSt2 (Douglas et al., 2020) was used for functional gene annotation of the 16S rRNA amplicon sequences to assess the molecular AR mechanisms, including the β -lactam, aminoglycoside, tetracycline, macrolide, phenicol, fosfomycin, sulfonamide, trimethoprim, rifamycin, quinolone and vancomycin resistance genes, as well as vancomycin, multidrug and cationic antimicrobial peptide (CAMP) resistance modules. Appropriate gene sets and gene variants were analyzed using the KEGG Brite Antimicrobial Resistance Genes database (KO01504) (Šamanić et al., 2021). Heatmap was generated to depict the relationship of the most common pathogenic genera in microbiomes (relative abundance expressed as square roots of the percentage values; threshold: 0.73) with resistance gene sets and variants by using reads from trimmed OTUs table and custom Python 3.9 scripts. Graphical design was done with Inkscape 1.0.1.

2.7. Data analysis

Relative abundance of bacterial taxa was calculated by dividing an absolute abundance with a total abundance of species at each taxonomic rank. Alpha diversity indices (Chao1, Shannon, Simpson, ACE, Good's coverage) were calculated using QIIME (v1.7.0) and plotted using MATLAB R2021a. Beta diversity was calculated on both weighted and unweighted Unifrac using QIIME software (v1.7.0). The PCoA analysis was displayed using the WGCNA, stat and ggplot2 packages in R software (v15.3). Analysis of similarities (ANOSIM) and multi-response permutation procedure (MRPP) analysis were performed by R software (Vegan package) to evaluate the significance of variations between and within the structure of bacterial communities from two WWTP submarine effluents. To determine the significantly different OTUs, the linear discriminant analysis (LDA) effect size (LEfSe) algorithm was conducted (Segata et al., 2011) using a threshold greater than 2.0 for the logarithmic LDA scores. Metastat was calculated by R software and p-value using a permutation test (p < 0.05 was statistically significant). Pearson correlations analysis and figures were done in MATLAB R2021a.

3. Results

3.1. Environmental parameters and bacterial counts at the two effluent-receiving area

The physical and chemical characteristics of the bottom and surface water layers were analyzed at both submarine outfalls (Fig. 1, Table S1). The T and SAL values showed an opposite pattern as colder and saltier water sinks. Unstratified water column prevailed during February, while formation of stratification started in May and followed seasonal cycle, with shallowest pycnocline detected in July and the deepest in September (Fig. S2.). The water column was well oxygenated at both sites based on dissolved oxygen levels. Higher levels of CHL, TUR, and CDOM were found at the bottom, at both stations, while an opposite pattern was observed for TN. The highest NO2–N and NO3–N concentrations were found at SB. The highest abundance of *E. coli* (660 CFU/100 mL) and fecal enterococci (151 CFU/100 mL) was observed at SS during sampling in February (Table 1). Overall, the median values of *E. coli* (50 CFU/100 mL) and enterococci (24 CFU/100 mL) counts were highest in SB and were followed by KB.

3.2. 16S rRNA sequencing analysis

The total number of filtered reads generated on the Illumina platform was 1,472,326. After quality filtering, a total of 967,900 effective tags, with a mean length of 411 bp, were clustered into 15,200 OTUs. On average, 760 OTUs were identified per sample, with 767, 451, 666 and 1156 OTUs detected in SB, SS, KB and KS samples, respectively. A total of 332 OTUs were common to samples from the two sites (Fig. S3). As expected, the highest overlap was found between the two bottom locations, which included 516 OTUs in common. On contrary, two surface locations shared the least OTUs (n 452) (Fig. S3).

3.3. Composition of bacterial community at two WWTP submarine effluents

A total of 37 bacterial phyla were identified at two effluent sites. The highest number of phyla was observed in SB (n 29), followed by KS, KB, and SS with 26, 25, and 24 identified phyla, respectively.

Differences between the two submarine discharge areas were noticeable at the phylum and class level (Fig. 2). The most abundant phylum at Stobreč effluent site was *Firmicutes* with the relative abundance of 45.64% in the bottom (SB) and 50.51% in surface (SS) samples and was followed by *Proteobacteria* (30.33% and 23.40%). *Cyanobacteria, Bacteroidetes* and *Actinobacteria* were present in a range from 3.53% to 10.97% in SB samples and from 5.24% to 6.45% in SS samples, respectively. On the other hand, *Proteobacteria* was the predominant phylum at the Katalinića brig effluent site with the relative abundance of 44.04% and 59.30% in KB and KS samples. These were followed by *Firmicutes* (32.90% and 15.54%), *Cyanobacteria* (6.33% and 14.30%), *Bacteroidetes* (<5%) and *Actinobacteria* (<3%).

The comparison of the community diversity at the class level was more informative, evidencing a distinctive pattern of each sample. A total of 71 bacterial classes were identified in this study, with *Clostridia*, *Alphaproteobacteria* and *Gammaproteobacteria* being the most abundant ones (Fig. 2). These were differently distributed between the samples. *Clostridia* was found to be the most abundant class in SB, SS and KB, while it was the fourth most prevalent in KS. *Gammaproteobacteria* was the second most abundant class in KS, KB and SS, and, depending on the sample, were followed by *Oxyphotobacteria*, *Alphaproteobacteria* and *Bacilli* (Fig. 2).

The relative abundance of bacterial taxa at the order, family, and genus levels is shown in Fig. S4. *Clostridiales* was the most abundant order except in KS, where SAR 11 clade of marine autochthonous bacteria occupied this position. Consequently, the human gut-related bacteria of the *Lachnospiraceae* of the order *Clostridiales* were the most

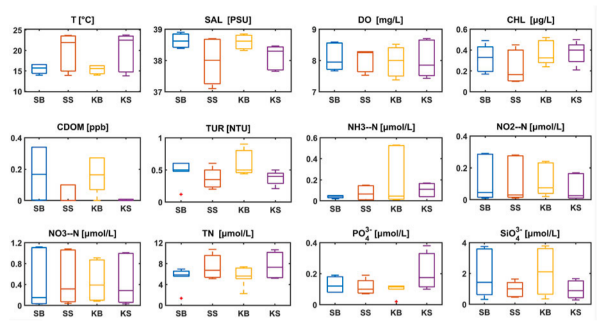


Fig. 1. Boxplots of environmental parameters at effluent discharge of Stobreč (bottom and surface, SB and SS) and Katalinića Brig (bottom and surface, KB and KS) WWTPs. Central line indicates the median value, and the bottom and top edges of the box indicate the 25th and 75th percentiles. The outliers are plotted individually using the '+' symbol.

Table 1 Total counts of FIB (CFU \pm SD/100 mL), total heterotrophic bacteria (CFU \pm SD/1 mL) and estimated percentage of the viable heterotrophic bacteria resistant (%R) to IMP and CL.

Bacteria	Site	February	May	June	July	September
E. coli	SB	0.7 ± 0.6	4.67 ± 2.5	219 ± 6.7	102.7 ± 10.7	50 ± 7
	SS	660 ± 14.7	0	0	0	0
	KB	44 ± 4.4	152 ± 10.8	34 ± 4.6	37.3 ± 8.9	1.3 ± 0.6
	KS	10 ± 4.6	0	0	0	0
Intestinal enterococci	SB	1.3 ± 0.6	4 ± 1	48.7 ± 4.2	27.3 ± 3.2	24 ± 4.6
	SS	151.6 ± 3.2	0	0	0.7 ± 0.6	0
	KB	15.3 ± 2.5	36.6 ± 3.2	27.3 ± 4.0	8.7 ± 0.6	0.7 ± 0.6
	KS	13.3 ± 2.3	6 ± 1	0	0	$\textbf{4.7} \pm \textbf{3.2}$
Total viable heterotrophic bacteria	SB	$\overline{163.3\pm1.2}$	776.67 ± 16.2	3017 ± 47.5	2577 ± 12.7	1071 ± 59.8
	SS	630 ± 17.5	1300 ± 30.3	1450 ± 5.57	2140 ± 63.3	1227 ± 37.5
	KB	2090 ± 47.5	2993 ± 27.3	1407 ± 48.5	1607 ± 68.1	1016 ± 12.0
	KS	456.6 ± 8.4	966.7 ± 5.8	2950 ± 21.9	2940 ± 16.1	1596 ± 17.6
% R-IMP	SB	0.43	0.86	1.50	0.47	15.69
	SS	0.39	2.55	2.21	0.23	29.34
	KB	0.10	1.34	2.84	0.26	35.43
	KS	0.45	2.75	7.81	0.37	22.56
% R-CL	SB	14.76	22.66	9.41	1.67	15.97
	SS	5.65	6.09	15.86	4.86	24.45
	KB	7.52	46.78	15.78	4.79	42.81
	KS	7.07	20.69	11.28	6.94	38.85

abundant family, accounting for between 26.1% and 18.6% of the bacterial community in SB, SS and KB. *Agathobacter* (7.1%), *Lactobacillus* (8.1%) and *Salmonella* (6.5%) were the most abundant genera in SB, SS and KB. On the other hand, *Cyanobiaceae* (10.7%) of the order *Synechococcales* (Oxyphotobacteria) were the most frequently presented family in KS, with *Synechococcus* CC9902 being the most abundant bacterial genus (8%). It is noteworthy that genera with a relative abundance of less than 2% accounted for more than 55% of the bacterial community in the samples from Katalinića brig (KS and KB). On the other hand, this percentage was about 45% in the samples from the Stobreč effluent site.

Furthermore, Fig. S5 shows a heatmap of the 35 most abundant genera in each sample, providing information on abundance and

clustering them by origin. The most abundant genera found in samples from the seafloor near both submarine outfalls (KB and SB) are rarely present in surface samples (KS and SS) and *vice versa*.

The microbial communities from the Katalinića brig site (KB and KS) showed higher richness and microbial diversity compared to the Stobreč outfall samples (Table 2, Fig. S6), as indicated by a higher number of genera with a relative abundance of <2%, as well as other less abundant genera (Fig. S4). Community diversity estimated by the Shannon's and Simpson's indices was highest in the surface samples (KS > SS > KB > SB). KS also had the highest community richness among surface and bottom samples and was significantly different (p < 0.05; Wilcox test) from SS.

According to the ANOSIM and MRPP analyses (Table S2), the

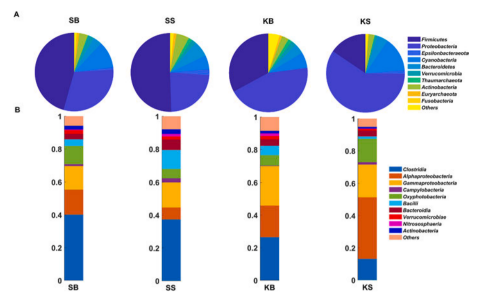


Fig. 2. Relative abundance of the ten most abundant taxa at phylum (A) and class (B) levels comprising the bacterial communities from the Stobreč (bottom and surface, SB and SS) and Katalinića Brig (bottom and surface, KB and KS) WWTPs effluent sites.

Table 2Alpha diversity indices showing the richness (observed species, Chao1 indices), diversity (Shannon and Simpson indices) and evenness of bacterial communities at two effluent-receiving sites over five-month period.

Indices	SB	SS	KB	KS
Observed species	307	285 ^a	308	374 ^b
Chao1	474.77	387.65 ^a	483.32	687.73 ^b
Shannon	5.95	6.15	5.89	6.29
Simpson	0.96	0.96	0.95	0.97
Evenness .	0.94	0.98	0.93	0.94
Pathogenic genera ^c	17.78	32.45	22.65	9.50

- ^a Significantly different (*p*-value <0.05) by Wilcox-test.
- ^b Significantly different (*p*-value <0.05) by Wilcox-test.

variations in bacterial community structure between KS and the other 3 groups were greater than within these groups, being statistically significant in the case of KS and SS (p < 0.05). Of note, the bacterial community at KS differed from the other samples already at the class level (Alphaproteobacteria > Gammaproteobacteria > Oxyphotobacteria > Clostridia).

3.4. Spatial variations of microbial community composition

Beta diversity was presented by the weighted UniFrac PCoA biplot based on Bray-Curtis distance (Fig. 3). SS and SB microbiomes plotted more closely, while differences were more evident between KS and KB. As expected, KS was the most distant from SS, as supported by Venn diagrams (Fig. S3). The monthly surface samples clustered very tightly except for those obtained in February (KS.2 and SS.2) that were plotted separately from the rest of the pool. For SS, this discrepancy is mainly due to the reduction of *Clostridia* and the higher abundance of *Alphaproteobacteria*, *Gammaproteobacteria*, *Campylobacteria* and *Oxyphotobacteria*, while for KS this is related to the increase of *Clostridia* and the lower abundance of *Alphaproteobacteria* and *Gammaproteobacteria*.

Significant differences in the composition of bacterial community between KS and SS were further confirmed by ANOSIM (R $\,$ 0.412, p 0.032) and MRPP (p $\,$ 0.013) tests (Table S2). According to LEfSE analysis, the classes with statistically significant difference (p < 0.05) were *Bacilli* in SS and *Alphaproteobacteria* in KS (Fig. 4., Fig. S7). In this

regard, the histogram of LDA scores showed that KS was much more enriched in *Rhodospirillales, Rhodobacterales* and *Puniceispirillales* of the *Alphaproteobacteria* class and these were followed by *Oceanospirillales* and *Alteromonadales* of the *Gammaproteobacteria* class, when compared to SS. On the other hand, SS was characterized by a higher abundance of *Lactobacillales* from the class of *Bacilli*, following by *Enterobacteriales* and *Bacteroidales* (Fig. 4).

The weighted UniFrac distance boxplot of beta diversity indices (Fig. S8) showed the differences in microbial community structure from bottom and surface samples collected at both effluent sites, and these were statistically significant (p < 0.05, Wilcox test) in the case of KS and KB. In support of the UniFrac analysis, LEfSE analysis showed that KS was more enriched in *Alphaproteobacteria* (o_Puniceispirillales; f_SAR116_clade), Gammaproteobacteria (f_Alteromonadaceae; g_Glaciecola) and Oxyproteobacteria (g_Synechococcus_CC9902), when compared to KB (Fig. S9a). The bacterial taxon with the highest differential abundance in KS was Synechococcus_CC9902 (LDA score [log10]>4). Between SS and SB (Fig. S9b), significantly different abundances occurred only for members of the order Lactobacillales, which were more abundant in SS (LDA score [log10]>4).

3.5. Influence of environmental factors on structure of the bacterial community

The dependence of the bacterial classes in studied samples and environmental conditions were investigated using CCA analysis (Fig. S10). The abundance of Clostridia was mainly influenced by CDOM. Furthermore, Alphaproteobacteria were positively correlated with PO₄³ and negatively correlated with CHL. The abundance of Bacilli, Bacteroidia, Nitrososphaeria, and Termoplasmata was influenced by nitrogen nutrients (NO3-N, NO2-N and NH3-N). This was confirmed by Pearson correlation analysis in which Bacilli (r 0.50; p < 0.05) and Nitrososphaeria (r 0.55; p < 0.05) were positively correlated with NO2–N. Bacilli, Nitrososphaeria and Termoplasmata correlated positively and significantly correlated with NO3-N (r 0.55; r 0.48; r Bacteroidia (r 0.58; p < 0.05) and Termoplasmata (r 0.44; p < 0.05) correlated significantly with NH3-N. CDOM had influence on abundance of Clostridia (r 0.60; p < 0.05), while Alphaproteobacteria correlated positively with PO_4^{3-} (r 0.51; p < 0.05) and negatively with CHL (r 0.46; p < 0.05).

Pearson correlation analysis (Fig. 4) showed that *Lactobacillales* of the class *Bacilli* correlated negatively with temperature, while a positive

^c Abundance (%) was defined as the ratio of the read count of the genus per total number of read counts in the sample.

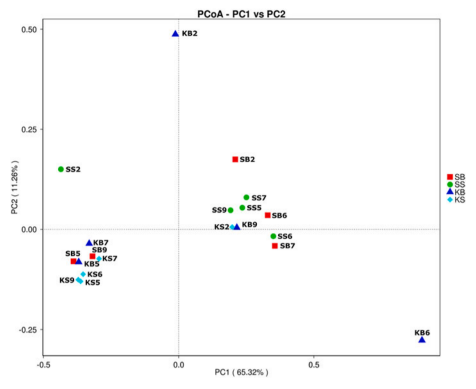


Fig. 3. Beta diversity of the bacterial communities at studied WWTPs effluent sites. Principal coordinates analysis (PcoA) plots were constructed applying the weighted Unifrac distance matrix on the CSS normalized OTU table data.

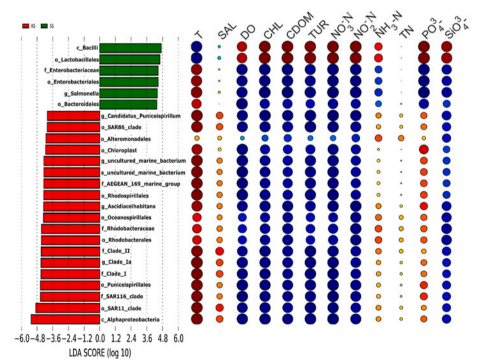


Fig. 4. Pearson correlation matrix of microbial taxa in KS and SS of LDA values higher than 2.0 with environmental factors. The circles indicate positive (red) and negative (blue) correlation, while circle size indicates the size of the correlation. T, temperature; SAL, salinity; DO, dissolved oxygen; CDOM, colored dissolved organic matter; NO $_3$, nitrate anion; NO $_2$, nitrite anion; NH $_4$ ⁺, ammonium cation; TN, total nitrogen; PO $_4$ ³, phosphate anion. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

correlation was found with most parameters, including CDOM, TUR, CHL, nitrites and nitrates. The opposite pattern of *Lactobacillales* was observed for the order *Enterobacteriales* of the class *Gammaproteobacteria* and the rest of the microbes persisting at both surface sites with environmental parameters. Only the orders *Enterobacteriales* and *Bacteroidales* showed a negative correlation with NH3–N and ${\rm PO_4}^{3-}$, while the rest of the microbes from both stations correlated positively.

3.6. Bacterial genera associated with human pathogenesis

Of the total of 628 identified genera, 74 include species associated with disease in humans. The relative abundance of these taxa per microbiome is shown in Table 2 and was the highest in SS and KB (32.45 and 22.65%). Of these, Acinetobacter, Salmonella, Escherichia-Shigella, Paenibacillus, Bacteroides, Lactobacillus, Bifidobacterium and Arcobacter,

were among the most prevalent pathogenic genera, and were abundant by >2% in at least one of the analyzed bacterial communities (Fig. 5).

3.7. Resistome prediction

PICRUSt2 inferred ARGs content in studied microbiomes (Fig. 6, Table S3). We detected the highest abundance of genes encoding for multidrug resistance efflux pumps, particularly the MexAB-OprM (max 20.9% in KS, avg 15.3%), AcrEF-TolC (max 10.4% in KS, avg 8%) and MdtEF-TolC (max 10.4% in KS, avg 6.8%). This was followed by the genes encoding for modified peptidoglycan precursors in Gram-positive bacteria, D-alanyl-D-lactate (vanA, vanB, vanD and vanM) and D-alanyl-D-serine (vanC, vanE, vanG, and vanL) ligase responsible for resistance to vancomycin and teicoplanin. These were primarily associated with microbiome from Strobreč WWTP, with 16% and 8.3% of relative abundances in SB (avgs 12.6% and 6.2%). This was followed by transporter genes encoding for tetracycline (including tet-like genes), macrolide and phenicol resistance (avg 6.8%), genes of bla operon (including regulator genes blaR1 and blaI, and β-lactamase genes blaZ and penP) conferring β-lactam resistance (avg 5.9%), acetyltransferases (avg 3.2%) mediating phenical resistance and the genes related to the repression of OprD porin (avg 3%) leading to imipenem resistance.

When comparing the ARGs content, SS was significantly enriched with genes related to cationic antimicrobial peptide (CAMP) resistance, such as dltABCD operon (abundance 5.6%, avg 3.1%, p < 0.05). In addition, genes encoding for OXA-type of class D β-lactamases were enriched in SS in comparison to SB and KS (0.25%, avg 0.18%). In addition, SS microbiome displayed a significant increase of genes encoding for multidrug resistance efflux pumps AcrEF-TolC, MdtEF-TolC and MexEF-OprN when compared to KS. Fig. 7 and Table S4. show more specifically in which extent the most abundant pathogenic genera contribute to the resistance to nine classes of antimicrobial drugs as well as to multiple classes. For instance, genus Klebsiella greatly contributed to aminoglycoside, CAMP, macrolide, phenicol, quinolone, β-lactam and multidrug resistance in the KB microbiome. On the other hand, a number of genera, among which Acinetobacter, Arcobacter, Bacteroides, Escherichia-Shigella, Pseudomonas and Salmonella, account for the most of multidrug resistance genotypes in analyzed microbiomes regardless of their origin. Overall, Acinetobacter, Bacteroides and Pseudomonas are the genera that contributed to the resistance to the most of antibiotic classes in analyzed bacterial communities (Fig. 7.).

4. Discussion

The main objective of this study was to provide a comprehensive overview of the microbiome structure and associated AR in treated effluent from submarine outfalls in the central Adriatic Sea in Croatia, as well as to assess environmental factors leading to microbiome variation at the bottom and the surface of the effluent-receiving water column.

4.1. Microbial counts

Higher counts of FIB in most of the samples collected from the bottom compared to those from the surface indicate that bottom sites are influenced by fecal pollution from the submarine outfalls. In addition, due to a lockdown caused by the COVID-19 pandemic, sampling was mainly conducted in the warmer months (May, June, July and September) characterized by a stratified water column. A significant decrease and complete absence of FIB in most surface samples suggest that the stratification of the water column was strong enough to maintain discharge below the pycnocline. On the other hand, the lack of stratification could explain the higher level of FIB as well as organic matter, TN, and PO_4^{3-} detected in surface samples in February at both stations, especially in SS. Nevertheless, gulls as sources of point pollution could have also contributed to the higher FIB counts in these samples (Araujo et al., 2014).

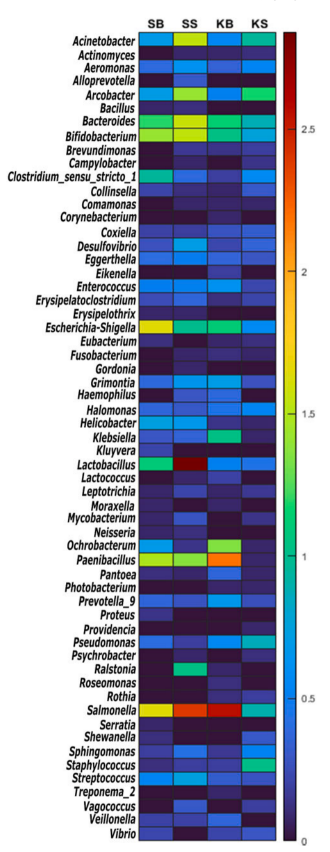


Fig. 5. Heatmap of the 74 pathogenic genera in water samples from Stobreč bottom (SB), Stobreč surface (SS), Katalinića Brig bottom (KB), and Katalinića Brig surface (KS), showing variations in relative abundance (expressed as square roots of the percentage values).

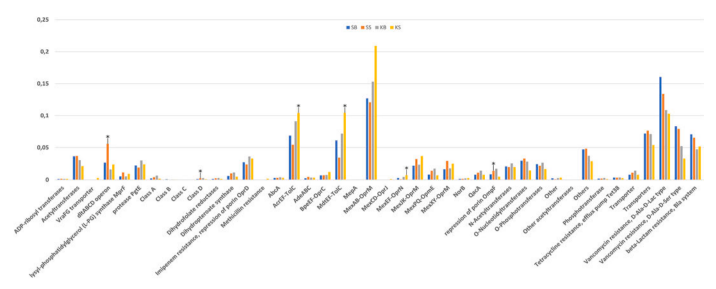


Fig. 6. Repertoire of antibiotic resistance mechanisms associated with the effluent-impacted microbiomes as predicted by PICRUSt tool, representing the average of the relative abundances of each gene set and gene variant available in the KEGG Brite Antimicrobial Resistance Genes database. Significant difference (in term of increase) in abundance between the studied microbiomes is denoted by * for p < 0.05 and **p < 0.01. Supporting information is available in Table S1. Sampling sites: Stobreč (bottom and surface, SB and SS) and Katalinića Brig (bottom and surface, KB and KS) WWTPs discharge sites.

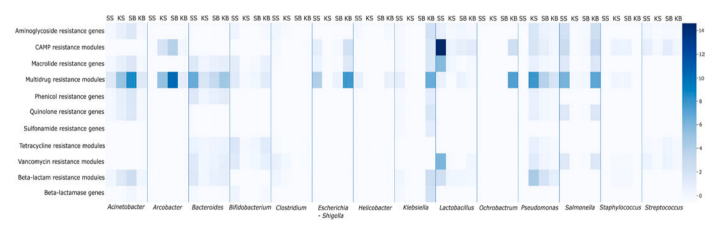


Fig. 7. Heatmap depicting the relative abundance of 14 most common pathogenic genera (expressed as square roots of the percentage values) in studied microbiomes with regards to their reference to KEGG Brite Antimicrobial Resistance Genes database.

Higher bacterial community diversity detected at the surface than at the bottom of the water column at both locations provides additional evidence that the microbiome in the bottom layer is under more anthropogenic pressure, particularly regarding human commensal bacteria and nutrients, resulting in a less divergent microbial profile. In this regard, KS was found to be under the least anthropogenic influence (as for human commensals, environmental parameters, and nutrients) and had the highest community diversity, which also corresponded to the highest richness as observed by alpha indices.

4.2. Most prevalent bacterial taxa at two effluent-receiving marine areas

Two WWTPs in the central Adriatic Sea process diverse types of wastewaters. Notably, Katalinića Brig WWTP treats municipal, hospital and rainfall-runoff wastewater, while Stobreč WWTP processes only the municipal wastewater. This could be one of the factors leading to a difference in microbial community structure between those two locations, with *Firmicutes* being the most abundant phyla near Stobreč wastewater submarine outfall (SB and SS) and *Proteobacteria* near Katalinića Brig submarine outfall (KB and KS). At both locations, bottom and surface microbiomes consist of typical phyla found in WWTPs effluents worldwide (Gai et al., 2014; García-Aljaro et al., 2019;

Narciso-da-Rocha et al., 2018; Port et al., 2012) and in adjacent freshwater (Bondarczuk and Piotrowska-Seget, 2019) or seawater (Zheng et al., 2019). Notably, the commonly detected phyla in treated wastewater include Firmicutes, Proteobacteria, Actinobacteria and Bacteroidetes with classes such as Clostridia, Bacilli, Bacteroidia, Alpha- and Gammaproteobacteria, reflecting the profile of the human gut microbiome. As expected, Clostridia, whose members are commonly found in human and animal feces (Cook et al., 2010; Liu et al., 2008), were the most abundant class in SB, SS, and KB. Besides Firmicutes, the most dominant phyla in this study were *Proteobacteria* and the class *Bacteroidia*, which prevail in the human and animal gastrointestinal tract and are thus highly abundant in sewage (Su et al., 2017). On the other hand, the presence of Actinobacteria, which were among the four most abundant phyla in this study, in coastal marine areas suggests an influence of terrestrial or freshwater runoff (Kelly and Chistoserdov, 2001; Port et al., 2012). The predominant phylum in such sewer systems is mostly Proteobacteria (Hu et al., 2017; Newton et al., 2013), Moreover, a handful of environmental studies point to Firmicutes as the dominant phyla in various WWTPs discharges (McLellan et al., 2010; Port et al., 2012). In accordance, Firmicutes were among the two most abundant phyla at surveyed sites, and were mainly represented by the families Lachnospiraceae, Ruminococcaceae, Lactobacillaceae and Paenibacillaceae. Both Lachnospiraceae and *Ruminococcaceae* are commonly associated with the human and animal gastrointestinal tracts and have been previously proposed as microbial source detection markers (Newton et al., 2013, 2011).

Proteobacteria were in this study mainly represented by Enterobacteriaceae, Rhodobacteraceae, Saccharospirillaceae, SAR116_clade and Clade_I. Enterobacteriaceae constitute an important part of the human gut (Pitout, 2008) and were found to be the most prevalent in KB, which could be related to the fact that hospital wastewater is processed in this particular WWTP, although University Hospital Split had a samewhat decrease in a number of hospitalized patients in 2020 when compared to a pre-pandemic 2019 (Croatian Institute of Public Health, 2020, 2021). It is noteworthy that hospital wastewater is a significant source of multidrug-resistant Enterobacteriaceae (Chagas et al., 2011; Galvin et al., 2010; Kotlarska et al., 2015; Picao et al., 2013). On the other hand, autochthonous marine Rhodobacteraceae and SAR116 clade which represent ubiquitous lineages of marine bacterioplankton (Giovannoni and Rappé, 2000; Garrity et al., 2005) contributed significantly to the higher diversity of KS microbiome. Regarding the less abundant bacterial taxa (Fig. 2), most of them were human gut commensals, whereas Deltaproteobacteria were found to be related with antibiotic polluted sites (Guan et al., 2018; Xiong et al., 2015).

4.3. Environmental influences on bacterial community structure

T, CDOM and nitrogen nutrients were shown to be the most determining factors related to the changes in the microbial community. The positive correlation of Alphaproteobacteria, Gammaproteobacteria and Bacilli with nutrients and CHL is a consequence of enrichment of water column with nutrient sources from the wastewater, which is in accordance with previous studies (Fierer et al., 2007; Wang et al., 2018; Zoppini et al., 2010). Clostridia indicate sewage pollution (Hill et al., 1993) and correlate with CDOM which is used for wastewater mapping in coastal waters (Nezlin et al., 2020; Rogowski et al., 2012). T and SAL were limiting factors for microbial growth (Miller et al., 2009; Wang et al., 2016). In our study, SAL was not the predominant factor, as chemical, rather than physical parameters, influenced the structure of microbial community, probably due to the loads of nutrients. Zheng et al. (2019) found that nutrient concentration was not significantly related to microbial community, which is contrary to our study but in agreement with some other studies (Lu et al., 2020; Wang et al., 2018). The observed differences could be related to the composition of influent, wastewater treatment stages used in different WWTPs, as well as the proximity of the marine outfalls and surveyed sites.

4.4. Spatial and vertical variations of microbiome composition

PCoA did not identify a clear pattern of grouping the microbiomes according to the site. Clustering of the surface microbiomes, at both marine outfall locations, was tighter when compared to the bottom microbiomes. This could be because the bottom of the water column is under intrusion of the wastewater plume as the wastewater discharge is discrete rather than continuous. The WWTP operates by pumping the effluent approximately every 45 min when the compensation basin is filled. In addition, the difference in the composition of the wastewater and the time lag between the plume intrusion and field sampling could also contribute to a higher heterogeneity of the monthly sampled bottom microbiomes. Surface microbiomes, besides KS.2 and SS.2, clustered tightly at the monthly level, but separately regarding the effluent site which proves our presumption about the lower effluent impact on surface microbial communities. During February, the vertical mixing of the water column may have attributed to such variations in microbial structure at surface locations. Additionally, FIB enumeration and the values of some environmental parameters support our presumptions about effluent rising to the surface due to unstratified conditions, which further induced changes in surface microbiome structure.

LEfSE analysis of surface microbiomes revealed a higher

anthropogenic impact in case of SS than KS. Members of classes *Bacilli* and *Gammaproteobacteria* were found to be significant taxonomic biomarkers in SS and *Alphaproteobacteria* in KS. *Alphaproteobacteria* members are morphologically, physiologically and metabolically diverse and adapted to different habitats (Rathsack et al., 2011; Williams et al., 2007), participating in several important metabolic processes such as photosynthesis, nitrogen fixation and ammonia oxidation (Campagne et al., 2012). Thus, the significance of *Alphaproteobacteria* in KS points to this site as the most diverse and under the lowest anthropogenic impact. On contrary, SS was more abundant with members of clinically important bacteria, such as *Gammaproteobacteria* and *Bacilli*, which are more associated with the anthropogenic impact. It is also important to note that KS bacterial community correlated positively with SAL, unlike SS, probably due to a larger abundance of autochthonous marine bacteria in KS.

LEfSE analysis revealed more differences in microbial composition between the bottom and surface microbiomes at the Katalinića Brig than at the Stobreč submarine outfall. These results provide additional evidence of a higher anthropogenic impact (as for nutrients, environmental parameters and human commensal bacteria) in SS than in KS. Moreover, KS biomarkers in comparison to KB were bacterial genera which commonly persist in surface coastal waters, such as SAR_116 clade (Treusch et al., 2009) and *Synechococcus* sp. (Agawin et al., 2003; Kim et al., 2018), evidencing higher variability of KS microbiome and lower impact of human gut-associated bacteria.

4.5. PICRUSt prediction of AR determinants

Annotation of microbiome-specific AR profiles based on 16S rRNA data revealed differences among the studied sites. KB was found to have the highest average incidence of CL and IMP resistance over a 5-month sampling period compared to other sites (Table 1). This may be related to the overabundance of multidrug efflux-related genes indicated by PICRUSt as predominant AR markers (Fig. 6). Notably, KB was found to have the highest relative abundance of chromosomally encoded tripartite RND efflux systems such as AcrAB-TolC, which contributes to the resilience of E. coli to nine classes of antibiotics, including carbapenems (Kobylka et al., 2020), as well as the MexAB-OprM, whose upregulation in Pseudomonas aeruginosa is generally associated with IMP resistance (Moubareck et al., 2019) and also tolerance to CL (Pamp et al., 2008). Moreover, genes encoding oxacillinases (OXA) of the class D β-lactamases were enriched in SS compared to SB, which was supported by the increased levels of IMP resistance (Table 1) as well as the highest detected abundance of Acinetobacter (Suppl Fig. S3) in this bacterial community. Various Acinetobacter spp. intrinsically possess class D enzymes (Evans and Amyes, 2014) with some, such as OXA-23 and OXA-58, being detected in Enterobacteriaceae and P. aeruginosa are spread globally (Munita and Arias, 2016).

Moreover, various emerging pathogenic genera were introduced by effluents and found at the bottom and surface of the water column, indicating anthropogenic effluent impact on the indigenous bacterial community. Introduced bacteria were found to carry characteristic AR signature (Fig. 7) and enrich the marine resistome, pointing to the potential risk for further inter- and intra-species transmission of AR determinants (Maravić et al., 2013, 2014; Rizzo et al., 2013; Zhuang et al., 2021).

5. Conclusions

Our study evidenced the impact of submarine discharges of inadequately treated wastewater on seawater quality at the bottom and surface of effluent-receiving waters of the central Adriatic. Despite stratified conditions during most sampling, the water column surface was affected by the effluent plume, resulting in alteration of physicochemical properties and enrichment of the marine bacterial community with nonindigenous and pathogenic bacteria carrying an arsenal of AR determinants. An integrated approach annotating the ARGs repertoire based on the taxonomic profile of the whole bacterial community enables a powerful basis for a comprehensive investigation of the extent of ARGs and bacterial pollution in the environment without using time-consuming and often limiting culture-based techniques, providing an important framework for the design and operation of more effective wastewater management.

Credit author statement

MK: Investigation, Formal analysis, writing of original draft, Visualization, Data curation; HK, MDŽ; , IŠ; : Validation, Methodology, Formal analysis; RA: Conceptualization; Funding acquisition; AM: Conceptualization, Formal analysis, writing of original draft, Visualization, Supervision, Data curation, Funding acquisition. All authors read and approved the final manuscript.

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Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.envpol.2021.118282.

Availability of data

The raw data obtained by Illumina-based amplicon sequencing of 16S rRNA gene have been deposited in the European Nucleotide Archive (ENA) at EMBL-EBI under accession number PRJEB45742.

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3.3. Submarine outfalls of treated wastewater effluents are sources of extensively- and multidrug-resistant KPC- and OXA-48-producing *Enterobacteriaceae* in coastal marine environment

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This study addresses the global health threat posed by carbapenemase-producing Enterobacteriaceae (CPE) present in the environment and introduced through submarine outfalls. Of fifteen KPC-producing *Enterobacteriaceae*, eight isolates were analysed by whole genome sequencing (WGS) and showed resistance to 19 antimicrobial classes, virulence genes and plasmid replicons. Several isolates carried 43-90 antibiotic resistance genes, while four carried the carbapenemase genes *bla*_{KPC-2} and *bla*_{OXA-48}. Remarkably, *bla*_{KPC-2} was identified on ~40kb IncP6 plasmids, which is a first European report.

This study highlights the occurrence of XDR and potentially virulent KPC -producing *E. coli* in coastal waters and highlights the risk of an infectious threat from submarine outfalls. Furthermore, all these strains were found only in submarine outfalls receiving hospital effluent from associated WWTP. This underlines the influence of hospital effluents on the further spread of AR determinants in the environment.



Submarine Outfalls of Treated Wastewater Effluents are Sources of Extensively- and Multidrug-Resistant KPC- and OXA-48-Producing Enterobacteriaceae in Coastal Marine Environment

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The rapid and ongoing spread of carbapenemase-producing Enterobacteriaceae has led to a global health threat. However, a limited number of studies have addressed this problem in the marine environment. We investigated their emergence in the coastal waters of the central Adriatic Sea (Croatia), which are recipients of submarine effluents from two wastewater treatment plants. Fifteen KPC-producing Enterobacteriaceae (nine Escherichia coli, four Klebsiella pneumoniae and two Citrobacter freundii) were recovered, and susceptibility testing to 14 antimicrobials from 10 classes showed that four isolates were extensively drug resistant (XDR) and two were resistant to colistin. After ERIC and BOX-PCR typing, eight isolates were selected for whole genome sequencing. The E. coli isolates belonged to serotype O21:H27 and sequence type (ST) 2795, while K. pneumoniae isolates were assigned to STs 37 and 534. Large-scale genome analysis revealed an arsenal of 137 genes conferring resistance to 19 antimicrobial drug classes, 35 genes associated with virulence, and 20 plasmid replicons. The isolates simultaneously carried 43-90 genes encoding for antibiotic resistance, while four isolates co-harbored carbapenemase genes blakec.2 and bla_{OXA-48}. The bla_{OXA-48} was associated with IncL-type plasmids in E. coli and K. pneumoniae. Importantly, the blakec-2 in four E. coli isolates was located on ~40 kb IncP6 broad-host-range plasmids which recently emerged as blakpc-2 vesicles, providing first report of these blakpc-2-bearing resistance plasmids circulating in E. coli in Europe. This study also represents the first evidence of XDR and potentially virulent strains of KPC-producing E. coli in coastal waters and the co-occurrence of bla_{KPC-2} and bla_{OXA-48} carbapenemase genes in this species. The leakage of these strains through submarine effluents into coastal waters is of concern, indicating a reservoir of this infectious threat in the marine environment.

Keywords: carbapenemase-producing Enterobacterlaceae, marine environment, coastal waters, Croatia, KPC, OXA-48

1

INTRODUCTION

Antibiotic resistance is one of the greatest threats to global health nowadays, leading to the higher mortality rates and increased economic costs (Pulingam et al., 2021). The natural environment has been recognized as one of its major reservoirs (Amarasiri et al., 2020), as antibiotic-resistant human pathogens have been detected in coastal marine areas (Šamanić et al., 2021), rivers (Ekwanzala et al., 2020), lakes (Su et al., 2020) and shellfish (Maravić et al., 2013). Effluents from the wastewater treatment plants (WWTPs) have been evidenced as particularly significant routes for dissemination of antibiotic resistance in the natural environment (Ekwanzala et al., 2020), especially the hospital effluents through which the emerging opportunistic pathogens directly enter from the hospital to the aquatic environment (Ekwanzala et al., 2019).

In recent decades, the rapid spread of Gram-negative bacteria resistant to the most potent β-lactam antibiotics, the carbapenems, and the continuous emergence of new resistant strains have raised the global alarm. In 2017, the World Health Organization defined priority categories for emerging multidrug-resistant pathogens for which new antimicrobials are urgently needed, with carbapenem-resistant Enterobacteriaceae (CRE) identified as critical (WHO, 2017). Carbapenem resistance in these bacteria arises mainly from the production of carbapenemases, of which KPC, SME, IMI and NMC belonging to Ambler class A, IMP, VIM and NDM metallo-β-lactamases (MBLs) belonging to class B, and OXA-48 and its derivatives belonging to class D have been detected worldwide (Bonomo et al., 2018; Brolund et al., 2019). In Croatia, CRE isolates are being increasingly reported in hospitals, mainly due to the rapid spread of KPC-producing Enterobacteriaceae first in the northwest (Jelić et al., 2016) and later in the southern coastal regions (Bedenić et al., 2021).

Considering the importance of CRE for public health and the One Health approach, we aimed to study for the first time their occurrence in the coastal waters of the eastern Adriatic, focusing on the area influenced by the submarine sewage outlets of two WWTPs, which could serve as potential routes for the introduction of these bacteria into the coastal marine environment. The isolated CRE were analyzed by PCR for the presence of carbapenemaseencoding genes, after which eight KPC-2-producing Enterobacteriaceae isolates were subjected to high-throughput DNA sequencing. We then performed a detailed search of the obtained genome sequences, focusing on the antibiotic resistance genes (ARGs), virulence factors and plasmid replicons. The isolates were also assigned to sequence types (STs) using the multi-locus sequence typing (MLST) scheme and their serotype was determined. This study led to the first identification of potentially virulent CRE in the marine environment in Croatia, evidencing a transmission route through submarine outfalls and a new reservoir of these opportunistic pathogens in Croatia outside hospital settings.

MATERIALS AND METHODS

Sampling

Fifteen Enterobacteriaceae isolates were recovered in June, July, and September 2020 as part of the project aimed to study the impact of treated submarine effluents in the coastal waters of the central Adriatic Sea in Croatia. Details of the sampling procedure and locations have been described previously (Kvesić et al., 2021). Briefly, the study focused on submarine effluents from the two WWTPs, the Katalinića brig and the Stupe-Stobreč, which mechanically treat wastewater from the wider Split area at an average flow rate of 35,000 and 30,000 m³/day, respectively, and discharge it through submarine outfalls into the coastal waters of the Brač and Split channels (Figure 1). While the Stobreč WWTP processes only municipal wastewater, the Katalinića brig WWTP treats municipal wastewater and stormwater runoff. The Katalinića brig WWTP also collects wastewater from the University Hospital Centre Split, the largest medical center in southern Croatia with 1,400 beds serving a population of approximately 500,000, which increases sharply in the summer months during the tourist season. The submarine outfalls of the Katalinića brig (43°29'22.7 "N, 16°27'11.2 "E) and the Stobreč WWTP (43°28'53.6 "N 16°31'04.3 "E) are located at a depth of 42 and 37 m, respectively.

Water samples were collected from the boat using a Niskin sampler, transferred to sterile 1L bottles, protected from light, and transported to the laboratory for further analysis within 4h at $4^{\circ}C$.

Bacterial Identification and Antibiotic Susceptibility Testing

One hundred milliliters of the water samples were filtered through 0.2 µm pore size MCE membrane filters (GE Healthcare, United Kingdom), which were then placed on CHROMID® Carba agar (bioMérieux, France) and incubated for 48 h. This chromogenic medium selects for the growth of CRE and allows the typical pink to the burgundy appearance of Escherichia coli colonies and blue-green to blue-grey of Klebsiella, Enterobacter, Serratia and Citrobacter spp. Incubation was carried out at 42°C to suppress the growth of autochthonous environmental species that are unable to grow under mesophilic conditions. Based on colony morphology, all putative Enterobacteriaceae isolates were cultivated in pure culture on MacConkey agar (Biolife, Italy) at 37°C for 18h and identified to species level using MALDI-TOF MS (Microflex LT mass spectrometer and MALDI Biotyper 4.1.80, Bruker Daltonics, Germany).

The isolates were tested for susceptibility to 14 antibiotics using Etest strips (AB Biodisk, Sweden) except for colistin (CL) whose susceptibility was tested using the broth microdilution method. The tests were performed, and the minimum inhibitory concentrations (MICs) were interpreted based on the European Committee on Antimicrobial Susceptibility Testing (EUCAST) guidelines (EUCAST, 2020). The MIC value of CL was recorded as the lowest concentration showing no visually detectable bacterial growth in the 96-well microtiter plates and was the



FIGURE 1 | Sampling sites (represented by squares) at the submarine outfalls of the two WWTPs, central Adriatic Sea, Croatia.

consensus value of the experiment performed in triplicate. The antibiotics tested (except CL) and their maximum concentrations were as follows: piperacillin/tazobactam (TZP, 256 µg), piperacillin (PIP, 256 µg), ceftazidime (CAZ, 256 µg), cefotaxime (CTX, 32 μg), cefepime (FEP, 256 μg), aztreonam (ATM, 256 μg), imipenem (IMP, 32 µg), meropenem (MER, 32 µg), ertapenem (ETP, 32 μg), ciprofloxacin (CIP, 32 μg), gentamicin (GEN, 256 µg), tetracycline (TET, 256 µg), and trimethoprimsulfamethoxazole (SXT, 1/19 µg). E. coli ATCC 25922 was used as a control. According to Magiorakos et al. (2012), multidrugresistant (MDR) phenotype was defined as acquired non-susceptibility to at least one antibiotic from three or more classes, while the extensively drug-resistant (XDR) phenotype was designated as non-susceptibility to at least one agent from all but two or fewer antibiotic classes (i.e., remaining susceptible to only one or two categories).

Isolates were further tested for the presence of class C AmpC β -lactamases with AmpC Etest (AB Biodisk) and carbapenemases with MBL Etest (AB Biodisk) and Rapidec Carba NP test (bioMérieux) according to the manufacturer's instructions. The production of extended-spectrum β -lactamases (ESBLs) was tested using clavulanic acid (CLA) combination discs. The

phenotype consistent with the production of ESBLs was defined by an increase in zone diameter of $\geq 5\,\mathrm{mm}$ for CAZ and/or CTX in combination with CLA compared to its zone when tested alone (EUCAST, 2013).

PCR Screening for Carbapenemase Encoding Genes and *mcr-1* Gene

Genomic DNA was extracted using the NucleoSpin Microbial DNA kit (Macherey-Nagel, United Kingdom) and the concentration and quality of DNA were analyzed using the NanoDrop® Spectrophotometer 1000 (Thermo Scientific, United States). Multiplex PCR assays were performed to screen for the presence of carbapenemase genes encoding class A KPC and class B IMP, VIM and NDM using the primers and PCR conditions described previously (Poirel et al., 2011). Isolates were screened by standard PCR for the presence of the *mcr-1* gene, which encodes plasmid-mediated colistin resistance (Liu et al., 2016). The amplified fragments were separated on a 1% (w/v) agarose gel, purified using the ReliaPrepTM DNA Clean-Up and Concentration System (Promega, United States) and subjected to Sanger sequencing of both strands in Macrogen Europe

service (Netherlands). The obtained nucleotide sequences were compared with the homologous sequences from the GenBank database using the BLASTn algorithm.¹

Molecular Typing of Bacterial DNA

To exclude the possibility of clonal relatedness between the isolates of the same species, enterobacterial repetitive intergenic consensus (ERIC) and BOX PCR analyses were performed using the primers and conditions previously described (Araújo et al., 2014).

High-Throughput DNA Sequencing and Computational Data Analysis

Genomic DNA was sent to Novogene (Cambridge, United Kingdom) for whole genome sequencing (WGS) and bioinformatics analysis of the raw sequencing data. DNA libraries were prepared using the NEBNext® DNA Library Prep Kit (Illumina, United States) and, after a quality check, were subjected to pair-end sequencing on the Illumina NovaSeq 6000 platform with a read length of 150 bp at each end. The obtained reads were subjected to further quality control. Then, the clean reads were mapped to the reference genomes to detect and annotate single nucleotide polymorphism (SNP), structural variants (SV) and copy number variation (CNV) according to the mapping results. FASTQ files containing clean sequences were further analyzed using tools available at the Center for Genomic Epidemiology,2 including multi-locus sequence typing (MLST) with MLST 2.0, sequence type (ST) with SerotypeFinder 2.0, presence of virulence genes (VirulenceFinder 2.0), resistance genes (ResFinder 4.1), and plasmid replicons (PlasmidFinder 2.1). In addition, antibiotic resistance and virulence profiling was performed using the ARESdb cloud platform introduced by Ferreira et al. (2020).3 This involved searching for marker sequences with coverage of >60% and identity of >90% to those cataloged in ARESdb (Ferreira et al., 2021).

To link carbapenemase KPC-2 and OXA-48 encoding genes to specific Inc. plasmid groups, plasmids were reconstructed with SPAdes v3.13.1 from trimmed (trimmomatic v0.39) sequencing reads, after which the replicon types and resistance markers were cross referenced to the de novo assemblies (Kudirkiene et al., 2018). Plasmid Finder v2.1 determined replicon types from the WGS assemblies generated by SPAdes. Because plasmid reconstruction from short-read sequencing is challenging and assembly typically results in many fragmented contigs per genome of unclear origin, plasmidSPAdes tool was used to identify as much as plasmid contigs. The algorithm in plasmidSPAdes predicted which contigs belong to plasmid DNA and assigned those contigs into components. Components containing specific plasmid replicons and their combinations from a selected strain were further used to search against NCBI nr database using BlastN for the most similar plasmids. De novo assembled plasmids were aligned against ARESdb

¹www.ncbi.nlm.nih.gov ²http://www.genomicepidemiology.org ³https://ares-genetics.cloud with thresholds set at >60% query coverage and >90% alignment identity to detect resistance markers. Further analysis of the generated assemblies was conducted using the Proksee server to create circular alignments of the reads to the reference plasmids available in the NCBI database.⁴

RESULTS

This study investigated the emergence and antibiotic resistance of CRE in submarine effluent-receiving coastal waters of central Adriatic to contribute to the global surveillance of these opportunistic pathogens outside of hospital settings.

Strain Isolation and Antibiotic Susceptibility Pattern

Twenty-two isolates that exhibited characteristic pink or blue-green colony morphology were recovered on selective CHROMID® Carba agar (bioMérieux). By MALDI-TOF MS, seven isolates were identified as Enterococcus faecium and excluded from further investigation. The remaining 15 isolates belonged to the Enterobacteriaceae family (nine E. coli, four Klebsiella pneumoniae and two Citrobacter freundii) and were obtained from water samples collected in June, July, and September 2020 near the submarine outfall of the Katalinića Brig WWTP. All but one isolate showed resistance to at least one carbapenem antibiotic. Among them, four isolates (one E. coli and three K. pneumoniae) were designated extensively drug resistant (XDR), while nine isolates (seven E. coli, one K. pneumoniae and one C. freundii) were multidrug resistant (MDR). Although two K. pneumoniae isolates (C1 and C2) were resistant to colistin, the mcr-1 gene was not detected by PCR. The detailed antibiotic resistance profiles of the Enterobacteriaceae isolates are shown in Table 1.

The Rapidec Carba NP test indicated carbapenemase production in all 15 isolates. PCR screening and Sanger sequencing further confirmed the presence of the carbapenemase gene $bla_{\rm KPC-2}$ in all but one *C. freundii* isolate (CF2), which carried $bla_{\rm KPC-29}$. This carbapenemase gene is derived from the ancestral allele $bla_{\rm KPC-3}$, and its expression does not affect the activity of carbapenems (Hobson et al., 2020). However, the CF2 isolate remained sensitive to all beta-lactam antibiotics tested, including cephalosporins (**Table 1**), casting doubt on the full expression of this gene in this isolate.

Based on the ERIC and BOX profiles (Supplemental Figure S1), eight isolates (four *E. coli*, three *K. pneumoniae*, and one *C. freundii*) that exhibited the most diverse profiles were subjected to WGS.

The draft genome sizes of the isolates ranged from 5.1 to 5.8 Mb, with diverse sizes of N50, and numbers of coding sequences and contigs (**Table 2**). A total of 137 genes were identified mediating intrinsic or acquired resistance to 19 antimicrobial drug classes, including penicillins, cephamycins,

⁴https://beta.proksee.ca/

CRE in Coastal Marine Environment

 TABLE 1 | Antibiotic resistance profiles of 15 KPC-producing Enterobacteriaceae isolates recovered in this study^a.

Isolate no.	M18	M12	M13	M14	M15	M16	M17	M19	M20	5a	M 11	C2	C1	CF1	CF2
Species	Escherichia coli	Klebsiella pneumoniae	Klebsiella pneumoniae	Klebsiella pneumoniae	Klebsiella pneumoniae	Citrobacter freundii	Citrobacter freundii								
Isolation date ^b	06/2020	06/2020	06/2020	06/2020	06/2020	06/2020	06/2020	06/2020	06/2020	06/2020	06/2020	07/2020	07/2020	09/2020	09/2020
KPC type	KPC-2	KPC-2	KPC-2	KPC-2	KPC-2	KPC-29									
PIP	1.5	>256	>256	>256	>256	>256	>256	>256	>256	>256	>256	>256	>256	64	2
PIP/TZB	0.75	32	>256	>256	256	192	>256	>256	256	32	32	192	>256	8	1
CTX	0.047	16	16	>32	>32	>32	12	24	16	>32	>32	24	>32	32	0.5
CAZ	0.19	8	8	8	8	3	8	4	12	12	8	8	64	32	0.38
FEP	0.032	1.5	32	48	256	4	32	8	8	32	16	32	256	1.5	0.094
ATM	0.094	1.5	96	>256	>256	64	256	6	128	8	12	1,5	>256	6	0.125
IPM	0.19	>32	32	>32	24	8	32	8	8	8	8	>32	>32	0.19	0.19
MER	8	>32	>32	>32	>32	12	8	16	>32	32	6	>32	32	1	0.5
ETP	0.004	32	32	>32	8	8	32	24	32	>32	32	>32	>32	0.006	0.004
CIP	0.012	6	4	4	0.75	12	6	4	12	6	6	4	6	0.75	0.012
GEN	0.5	16	1	1	0.5	2	1	2	3	8	8	8	1	0.5	0.5
TET	4	16	2	3	32	3	3	2	12	12	>256	16	4	4	4
SXT	0.064	0.5	32	>32	0.75	0.19	>32	0.38	32	0.25	0.25	0.5	0.038	0.064	0.038
CL	0.125	2	0.125	0.25	0.125	0.5	0.125	0.0625	1	0.125	0.0625	32	16	0.125	0.0625
Resistance phenotype		MDR	XDR	XDR	XDR	XDR	MDR	MDR							
AmpC Etest	neg	pos	neg	neg	neg	neg	pos	neg							
Rapidec Carba NP test	pos	pos	pos	pos	pos	pos									
ESBL test	neg	pos	neg	pos	pos	neg	neg	neg	pos	pos	pos	pos	pos	neg	neg

TZP, piperacillin/tazobactam; PIP, piperacillin; CAZ, ceftazidime; CTX, cefotaxime; FEP, cefepime; ATM, aztreonam; IMP, imipenem; MER, meropenem; ETP, ertapenem; CIP, ciprofloxacin; GEN, gentamicin; TET, tetracycline; SXT, trimethoprim-sulfamethoxazole; CL, colistin; MDR, multidrug-resistant; XDR, extensively drug-resistant; neg, negative; and pos, positive.

^aResistance phenotype is indicated by shading according to EUCAST (2020) except for TET that was evaluated based on CLSI (2020) breakpoints.

blsolation date is given as month/year.

TABLE 2 | Metadata of the whole-genome sequenced CRE isolates from Croatia.

	M12	M14	M17	M20	5a	M11	C2	CF1
Species	Escherichia coli	Escherichia coli	Escherichia coli	Escherichia coli	Klebsiella pneumoniae	Klebsiella pneumoniae	Klebsiella pneumoniae	Citrobacter freundii
Genome size (bp)	5,207,851	5,243,436	5,173,484	5,186,809	5,667,286	5,659,465	5,826,947	5,103,429
No. of CDS ^a	5,028	5,082	4,969	5,023	5,339	5,345	5,554	4,865
No. of contings	240	231	207	224	342	289	195	74
No. of contings >1,000 bp	159	164	135	164	204	199	132	45
Average depth (x)	223	185	205	199	224	248	229	264
GC content (%)	50.54	50.55	50.59	50.6	56.92	56.92	57.04	51.84
N ₅₀ (bp)	89,262	87,097	90,500	78,173	70,753	74,345	116,900	372,763
No. of tRNAs	78	81	79	82	77	77	81	78
SRA accession no.	SAMN22028927	SAMN22028930	SAMN22028929	SAMN22028928	SAMN22028932	SAMN22028933	SAMN22028931	SAMN2202893

^aCDS, coding DNA sequences.

cephalosporins, carbapenems, penems, monobactams, fluoroquinolones, aminoglycosides, macrolides, phenicols, quinolones, sulfonamides, trimethoprim, rifampicin, tetracyclines, fosfomycin, nitroimidazoles, peptides, and aminocoumarin (Supplemental Table S1). The isolates were found to harbor between 43 and 90 gene markers associated with resistance phenotypes, with the highest number detected in *E. coli* genomes (Table 3). Most of these genes were associated with intrinsic resistance mechanisms such as regulation and transport by resistance-nodulation-cell division (RND) and major facilitator superfamily (MFS) antibiotic efflux pumps or porin uptake (Supplemental Table S1). Moreover, the isolates harbored three to nine bla genes, out of which four isolates (E. coli M12, M14, and M20 and K. pneumoniae C2) simultaneously possessed two carbapenemase-encoding genes, blaKPC-2 and blaOXA-48. In addition, eight sequenced genomes possessed a total of 20 resistance-associated plasmid replicons and 35 genes involved in bacterial virulence (Table 3; Supplemental Table S1).

Escherichia coli M12, M14, M17, and M20 Isolates

The four $bla_{\rm KPC}$ -carrying E.~coli isolates subjected to WGS were all of serotype O21:H27 and ST2795. Furthermore, the isolates shared a set of 26–30 virulence-related genes (**Table 3**), including those encoding the outer membrane usher protein (FimD), flagellar biosynthesis protein (FlhA), and the locus of enterocyte effacement (LEE) encoding the type III secretion system effector protein (EspX1). A number of other genes involved in pathogenicity were discovered as well, including glutamate decarboxylase (gad), long polar fimbriae (lpfA), tellurium resistance protein (terC), toxin-antitoxin systems (yafQ,~pemK,~pemI), type 1 fimbriae (S - fimbrial adhesion minor subunit; genes sfaH and sfaG), small toxic polypeptide (ldrD), polyamine transport protein D (potD), flagellar fli genes, laminin-binding

fimbriae (*elfG*) and carbon starvation protein A (*cstA*), pointing to the virulence potential of these isolates.

The further similarity between these strains was observed in their plasmid replicon content, with Inc replicons of plasmids FIB(K) and P6 detected in all four strains. Nevertheless, each strain exhibited a unique plasmid replicon pattern, comprising 6-10 replicon types per genome (Table 3). More diversity was observed among the ARGs, of which strains M12, M14, M17, and M20 possessed a total of 90, 79, 82, and 79 genes associated with the regulation or acquisition of antibiotic resistance (Table 3; Supplemental Table S1). Nine bla genes were identified, including the carbapenemase encoding genes blaKPC-2 and blaOXA-48, and ESBL genes bla_{GES-1}, bla_{GES-2}, bla_{OXA-2}, bla_{OXA-10}, and bla_{CTX-M-3}. Three strains (M12, M14 and M20) co-harbored bla_{KPC-2} and bla_{OXA-48}. Among others, ARGs mediating resistance to trimethoprim (dfrA14), quinolones (qnrVC4, qnrS1, qnrB6), aminoglycosides (ant(3")-Ii-aac(6')-IId, ant(3")-Ia, aph(3")-Ib) and sulphonamide (sul1, sul2) were continuously detected.

Moreover, further analysis of the genomes identified the $bla_{\text{KPC-2}}$ gene in IncP6 plasmid contigs of 38,767, 14,644, 25,016, and 14,644 bp in *E. coli* M12, M14, M17, and M20, respectively.

BlastN search against NCBI nr database revealed that a 38,767-bp contig from *E. coli* M12 had query coverage of 91% and nucleotide identity of 99.75%, 99.75%, and 99.68% with IncP6 plasmids deposited in GenBank: p121SC21-KPC2 from Spanish wastewater *C. freundii* (Genbank accession no. LT992437; Yao et al., 2017), pKOX3-P5- KPC from a clinical *Klebsiella oxytoca* in China (GenBank accession no. KY913901; Wang et al., 2017), and pWW14A-KPC2 from wastewater *Klebsiella quasipneumoniae* in Argentina (Ghiglione et al., 2021). Hybrid plasmids pM12-KPC2, pM14-KPC2, pM17-KPC2, and pM20-KPC2 were reconstructed to a size of ~40kb and compared to plasmids of both environmental and clinical origin previously reported in the literature (Dai et al., 2016; Wang et al., 2017;

TABLE 3 | Molecular characteristics of eight whole-genome sequenced CRE isolates.

	M12	M14	M17	M20	5a	M11	C2	CF1
Species	Escherichia coli	Escherichia coli	Escherichia coli	Escherichia coli	Klebsiella pneumoniae	Klebsiella pneumoniae	Klebsiella pneumoniae	Citrobacter freundii
Serotype	O21:H27	O21:H27	O21:H27	O21:H27				
ST	2795	2795	2795	2795	37	37	534	128
Selected	dfrA14, qnrVC4,	dfrA14, qnrS1,	aac(6')-lb-cr,	dfrA14, sul2,	sul1, tet(A),	fosA, oqxA	oqxB, oqxA	bla _{KPC-2} ,
antibiotic	cmIA5, mdf(A),	sul2, mdf(A),	aadA16, bla _{KPC-2} ,	bla _{OXA-10} ,	oqxA, oqxB,	oqxB, bla _{GES-5} ,	aac(6')-lb-cr,	bla _{CMY-157} (43)
resistance	$mph(B)$, bla_{KPC-2} ,	mph(B),	sul1, qnrB6,	bla _{GES-1} ,	aac(6′)-lb-cr	bla_{KPC-2} , bla_{OXA-2} ,	sul1, fosA,	
genes (total no.)a	bla _{GES-2} , bla _{OXA-2} ,	bla _{KPC-2} ,	mdf(A), mph(B),	bla _{KPC-2} ,	bla _{CTX-M-3} , bla _{OXA-2} ,	bla _{CTX-M-3} ,	bla _{KPC-2} , bla _{GES-5} ,	
	bla_{OXA-48} , bla_{OXA-10} ,	bla _{OXA-48} ,	arr-3, dfrA27 (82)	$bla_{OXA-48},$	bla _{GES-5} , bla _{KPC-2} ,	aph(3")-lb,	bla _{SHV-11} , bla _{OXA-48}	
	ant(3")-la,	bla _{GES-1} ,		aph(3")-lb,	fosA, aph(3")-lb,	aph(6)-Id, tet(A),	(58)	
	ant(3")-li-aac(6')-	bla _{OXA-10} ,		mdf(A), qnrS1,	aph(6)-ld, (62)	sul1 (62)		
	IId (90)	aac(6')-lb-cr,		(79)				
		aph(3")-lb, (79)						
Selected genes linked to virulence (total no.) ^a	gad, IpfA, terC, yafQ, sfaH, IdrD, potD, pemK, pemI, filY, filZ, filQ, filA, filG, fill, nagA, fihA, fimD, fdeC, EspX1, epsJ, elfG, cstA (30)	gad, lpfA, terC, sfaH, ldrD, potD, pemK, pemI, nagA, fliY, fliZ, fliQ, fliA, fliG, fill, flhA, fimD, EspX1, epsJ, elfG, cstA (26)	gad, IpfA, terC, IdrD, potD, pemK, pemI, nagA, flhA, fliY, fliZ, fliQ, fliA, fliG, fliI, fimD, EspX1, epsJ, elfG, cstA (26)	gad, IpfA, terC, sfaH, IdrD, potD, pemK, pemI, nagA, flhA, fliY, fliZ, fliQ, fliA, fliG, fliII, fimD, EspX1, epsJ, elfG, cstA (26)	sinR, sfaG, potD, filY, fimD, feoB eutB (9)	sinR, sfaG, potD, filY, fimD, feoB, eutB (9)	sinR, sfaG, potD, pemK, pemI, fliY, fimD, feoB, eutB, cstA (12)	potD, fil/Y, fil/Q, fil/P, fill, fil/G, fil/A fil/A, eutB, cst/ (10)
Plasmids (Inc)	FIB(K), L, P6, C, Col440I, Col(pHAD28)	FIB(K), FII, FII(Yp), L, N, P6, X5, Y, Col440I, Col440II	FIB(K), FII, P6, R, Col440I, Col(pHAD28)	FIB(K), FII, FII(Yp), L, N, P6, Y, Col(IRGK), Col440I,	FIA(HI1), FII(K), FII(Yp), X5, Y, Col440I, Col440II, Col(pHAD28)	FIA(HI1), FII(K), FII(Yp), X5, Y, Col440I, Col440II, Col(pHAD28)	FIB(K), L, R, Q1, FII(pMET), FII(pKP91), Col440I, Col(pHAD28)	FIB(pHCM2)

^aList of total genes associated with the antibiotic resistance and virulence is available in the Supplementary Material.

Yao et al., 2017; Pérez-Vazquez et al., 2019; Ghiglione et al., 2021; **Figure 2**). Analysis of the genetic environment of $bla_{\text{KPC-2}}$ revealed that this gene is located within a $\Delta \text{ISKpn6}/bla_{\text{KPC-2}}$ - $\Delta bla_{\text{TEM-1}}$ -ISKpn27 sequence within a Tn3-based transposon interrupted by an ISApu-flanked element (**Figure 2**), consistent with previous reports (Dai et al., 2016; Ghiglione et al., 2021).

The $bla_{\rm OXA-48}$ was associated with IncL-like plasmids in E.~coli, but the hybrid $bla_{\rm OXA-48}$ -bearing plasmids could not be reconstituted (the contigs containing this carbapenemase gene were 2,231-bp long in all four OXA-48-positive E.~coli and K.~pneumoniae isolates). However, the IS1R element flanking the OXA-48-encoding gene was detected in E.~coli M12 isolate.

Klebsiella pneumoniae 5a, M11 and C2 Isolates

Three KPC-2-producing K. pneumoniae isolates concurrently carried a total of 62, 58, and 62 genes mediating resistance multiple antibiotics, respectively (Table 3; Supplemental Table S1). Isolates 5a and M11 were affiliated to ST37, while C2 belonged to ST534. Moreover, isolates 5a and M11 exhibited the same antibiogram and XDR phenotype (Table 1), and a similar ARGs content (Table 3; Supplemental Table S1). However, ERIC and BOX-PCR typing excluded their clonality, and WGS data analysis revealed that differed by the aac(6')-Ib-cr gene, these conferring fluoroquinolone and aminoglycoside resistance, which was found in the genome of strain 5a and not M11. On the other hand,

K. pneumoniae C2 harbored less ARGs than the former two strains but was resistant to 13 out of 14 tested antimicrobial drugs, remaining susceptible only to trimethoprim-sulfamethoxazole. Nevertheless, identification of sul1 gene in its genome could eventually result in nonsusceptibility even to this antibiotic. Unfortunately, we were not able to reconstruct $bla_{\text{KPC-2}}$ -and $bla_{\text{OXA-48}}$ -bearing plasmids in K. pneumoniae isolates due to the short-read genome sequences.

Moreover, some virulence-related genes have been concurrently detected in all three *K. pneumoniae* genomes (**Table 3**), codifying for type 1 fimbriae (*fimD*, *sfaG*), ferrous ion transport (*feoB*), ethanolamine-ammonia lyase (*eutB*), extracellular matrix production (*sinR*), flagellae (*fliY*), polyamine transport (*potD*), and nickel and cobalt resistance (*cnrA*), respectively. The C2 strain harbored additional three virulence genes encoding the toxin-antitoxin system (*pemK*, *pemI*) and carbon starvation protein A (*cstA*; **Table 3**).

Citrobacter freundii CF1

KPC-2-producing *C. freundii* CF1 belonged to ST128 (**Table 3**). Compared to *E. coli* and *K. pneumoniae* isolates, this strain showed the least diversity of virulence markers, mainly harboring the genes encoding the flagellar apparatus (*fli*), polyamine transport protein D (*potD*), ethanolamine ammonia lyase (*eutB*) and carbon starvation protein A (*cstA*; **Table 3**). Furthermore, this strain contained the fewest ARGs, 43 in total (**Table 3**; **Supplemental Table S1**), and was sensitive to all tested antimicrobial agents (**Table 1**).

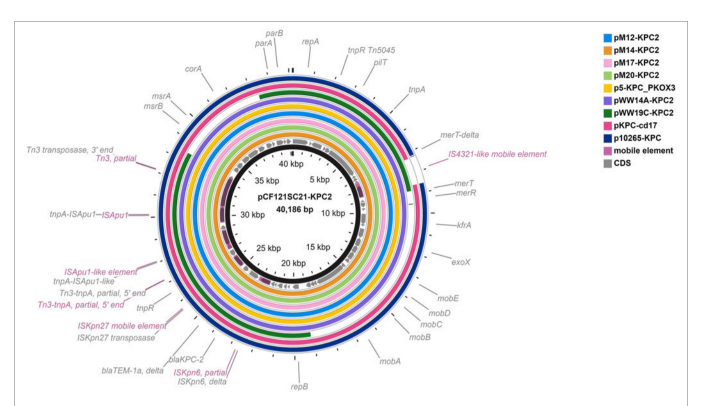


FIGURE 2 | Comparison of plasmids pM12-KPC2, pM14-KPC2, pM17-KPC2, and pM20-KPC2 with IncP6 bla_{KPC-2}-bearing plasmids of environmental and clinical origin described in the literature: pCF121SC21-KPC2 from Spanish wastewater *Citrobacter freundii* (Genbank accession no. LT992437; Yao et al., 2017), pKOX3-P5-KPC from clinical *Klebsiella oxytoca* in China (GenBank accession no. KY913901; Wang et al., 2017), pWW14A-KPC2 from wastewater *Klebsiella quasipneumoniae* in Argentina (GenBank accession no. CP080103; Ghiglione et al., 2021), pWW19C-KPC2 from wastewater *Enterobacter asburiae* in Argentina (GenBank accession no. CP080110; Ghiglione et al., 2021), p10265-KPC from clinical *Pseudomonas aeruginosa* in China (GenBank accession no. KU578314; Dai et al., 2016), and pKPC-cd17 from *Aeromonas* sp. from hospital environment in United States (GenBank accession no. CP026224) which is 100% identical to the *bla*_{KPC-2}-carrying plasmid in *Klebsiella oxytoca* from Spanish hospitals (Pérez-Vazquez et al., 2019). The plasmid pCF121SC21-KPC2 was taken as a reference plasmid (black circle).

DISCUSSION

Carbapenemase-Producing *Escherichia* coli Isolates

All four blaKPC-carrying E. coli isolates that were subjected to WGS belonged to the serotype O21:H27 and ST2795, which was previously identified in the United Kingdom.⁵ The strains carried several genes involved in pathogenicity or other function. For instance, we identified genes encoding the outer membrane usher protein FimD and flagellar biosynthesis protein FlhA associated with the urinary pathogenic E. coli (UPEC), and the effector protein EspX1 of the type III secretion system common to enterohemorrhagic E. coli (EHEC), all of which were previously detected in wastewater from WWTPs (Zhi et al., 2019). Furthermore, gad gene, also detected in all four E. coli genomes, is commonly involved in resistance to gastric acid, allowing E. coli to survive in the acidic host environment (Mates et al., 2007). The lpfA gene, encoding for the long polar fimbriae, was found to be associated with the gut colonization and the attachment to Peyer's patches in mice (Cordonnier et al., 2017), and was identified

adherent-invasive *E. coli* enrolled in the pathogenesis of Crohn's disease (Chassaing et al., 2011). Both genes were also identified in the KPC-producing *E. coli* from the riverine environments (Bleichenbacher et al., 2020). Moreover, *terC* gene, encoding the heavy metal resistance, is found to be significantly correlated with the presence of other virulence factors in the pathogenic strains of *E. coli* isolated from humans, animals, and food (Orth et al., 2006).

The PemK/pemI type II toxin-antitoxin system was also detected in the genomes of $bla_{\rm KPC}$ -positive E.~coli isolates from this study. This module is consisted of a stable toxin and an unstable antitoxin that degrades under stress conditions, enabling the toxin to inhibit the basic cellular processes. Notably, it has been associated with the bacterial persistence in inhospitable conditions, phage inhibition and biofilm formation (Ramage et al., 2009; Hernandez-Ramirez et al., 2017), as well as the IncF plasmid maintenance, conjugation, and spreading (Walling and Butler, 2016; Diaz-Orejas et al., 2017). This system was previously identified in hypermucoviscous carbapenem-resistant E.~coli within and outside the hospital environment (Woodford et al., 2009; Mathers et al., 2015; Zurfluh et al., 2018).

Moreover, total of nine carbapenem-resistant $E.\ coli$ isolates from this study were found to harbor bla_{KPC-2} gene. It is

⁵https://enterobase.warwick.ac.uk/

important to note that the environmental E. coli carrying this carbapenemase gene have rarely been reported, and they have all been recovered from the river water (Poirel et al., 2012; Xu et al., 2015; Yang et al., 2017). To the best of our knowledge, this is the first identification of KPC-producing *E. coli* in coastal marine waters. More importantly, the co-occurrence of bla_{KPC-2} and bla_{OXA-48} in E. coli has not been previously reported in the literature. So far, the bla_{KPC} gene in E. coli has been mainly reported in countries with a high prevalence of KPC-producing K. pneumoniae, indicating the possibility of interspecies gene transfer with K. pneumoniae serving as a blaKPC reservoir (Grundmann et al., 2017). In this regard, we should take into consideration a high prevalence of KPC-producing K. pneumoniae in University Hospital Split (Bedenić et al., 2021) and the fact that the isolates were recovered from the water samples collected near the submarine outfall of the WWTP that treats the hospital wastewater. Nevertheless, this presumption should be carefully addressed in future research, focusing on the genetic environment of the hospital KPC-producing strains. To the best of our knowledge, there are no available data on KPC-producing E. coli or K. pneumoniae from the University Hospital Split analyzed by WGS. The molecular characterization of plasmids harbouring blaKPC-2 gene in Croatia was performed for clinical K. pneumoniae, including those from University Hospital Split that were found to carry this gene on IncFII plasmids (Jelić et al., 2016; Bedenić et al., 2021), or those untyped by PCR-based replicon typing (PBRT; D'Onofrio et al., 2020), as well as in case of river K. pneumoniae that harboured bla_{KPC-2} gene on IncFII plasmids (Jelić et al., 2019). Notably, bla_{KPC-2} _bearing IncP6 plasmids were not previously reported in Croatia.

Moreover, all four bla_{KPC-2} bearing plasmids from E. coli isolates in this study were of ~40 kb and highly similar to the IncP6 bla_{KPC-2}-containing plasmids from wastewater C. freundii in Spain (Yao et al., 2017), clinical K. oxytoca in China (Wang et al., 2017) and wastewater K. quasipneumoniae in Argentina (Ghiglione et al., 2021), pointing to their global circulation. In comparison to plasmids from other incompatibility (Inc) groups, the KPC-2-encoding gene has been rarely detected in IncP6 resistance plasmids (Yao et al., 2017). However, recent studies confirmed that emergence of bla_{KPC-2} gene on mobilizable IncP6 broad-host-range plasmids enhanced its dissemination among different members of Enterobacteriaceae in clinical settings and the environment (Pérez-Vazquez et al., 2019; Ghiglione et al., 2021). This study provides further evidence to this speculation, documenting for the first time the bla_{KPC-2} association with IncP6 plasmids in E. coli in Europe.

Furthermore, 6–10 plasmid replicons were detected in $E.\ coli$ genomes. Among them, IncN and Col-type replicons have been previously associated with the occurrence of $bla_{\rm KPC}$ in human $E.\ coli$ from the global surveillance studies (Stoesser et al., 2017). In addition, three out of four analysed $E.\ coli$ genomes contained the $bla_{\rm OXA-48}$ carbapenemase gene associated with IncL plasmids. Notably, OXA-48-producing Enterobacteriaceae have widely disseminated in Croatian hospitals over the past years (Bedenić et al., 2018), with OXA-48-positive $K.\ pneumoniae$ reported at different wards in University Hospital Split. Since IncL-like plasmids were found to enable the transferability of

 $bla_{\rm OXA-48}$ in *E. coli* strains from northern Croatia (Bedenić et al., 2018; Drenjančević et al., 2019), findings from this study further enhance their relevance as reservoirs of $bla_{\rm OXA-48}$ in Croatia.

Moreover, Ambler class A GES-type ESBLs which were identified in KPC-2-producing *E. coli*, including GES-1 (isolates M14 and M20) and GES-2 (isolate M12) may have additionally enhanced their nonsusceptibility to beta-lactams as these enzymes effectively hydrolyse penicillins and expanded-spectrum cephalosporins (Castanheira et al., 2021). GES-2, in comparison to GES-1, also displayes hydrolytic activity against imipenem (Poirel et al., 2001).

In addition, other resistance determinants identified in the genome of our *E. coli* isolates, including the aminoglycoside resistance gene *aph*(3")-*Ib*, trimethoprim resistance gene *drfA17* and sulfonamide resistance gene *sul2*, were previously identified in clinical *E. coli* from Croatia (Bedenić et al., 2018).

Carbapenemase-Producing Klebsiella pneumoniae Isolates

In this study, we found that two $\mathit{bla}_{\mathrm{KPC-2}}\text{-}\mathrm{positive}$ isolates (5a and M11) belong to ST37. Notably, MDR K. pneumoniae strains of the same lineage, bearing the bla_{KPC-2} (Bedenić et al., 2012) and bla_{OXA-48} (Jelić et al., 2018) carbapenemase genes were previously reported in Croatian hospitals, but not in the natural environment. KPC-producing K. pneumoniae of other STs were previously isolated from aquatic environments (Ekwanzala et al., 2019), including ST258 in river water in Croatia (Jelić et al., 2019). Therefore, this study reports the first identification of KPC-producing K. pneumoniae in the marine environment in Croatia. Both of K. pneumoniae ST37 isolates in this study harbored replicons of plasmids known to enable the spread of ARGs in Enterobacteriaceae. Namely, multiple IncF replicons (FIIK, FIB, FIA, and/or FII) were previously identified in K. pneumoniae and other Enterobacteriaceae (Carattoli, 2009; Huang et al., 2012). On the other hand, K. pneumoniae strain C2 was affiliated to ST534, which was previously detected in the hospital environment in Israel (Adler et al., 2015). Our strain additionally harbored IncL and IncR-type plasmid replicons which have been previously described as vehicles of bla_{KPC-2} (Garbari et al., 2015) as well as bla_{OXA-48} in Croatia (Bedenić et al., 2018; Drenjančević et al., 2019).

K. pneumoniae 51, M11, and C2 isolates produced the Ambler class A GES-5 variant, which confers low carbapenemase activity in addition to penicillins and cephalosporins (Gomi et al., 2018; Castanheira et al., 2021), although GES-5-positive isolates with elevated MICs for imipenem, meropenem, and ertapenem have also been reported (Literacka et al., 2020). In this regard, production of GES-5 may have increased the resistance to carbapenems and other beta-lactams in our isolates. It should be noted that the epidemiology of GES producers is poorly understood, as GES carbapenemase-producing Enterobacteriaceae often stay unreported by resulting falsely negative in the Carba NP test due to the relatively weak activity toward carbapenems (Gomi et al., 2018; Literacka et al., 2020). Nevertheless, hospital outbreaks due to carbapenem-resistant GES-5-positive K. pneumoniae have recently been reported in Portugal (Mendes et al., 2022) and Poland (Literacka et al.,

2020), highlighting their clinical relevance. Apart from the carbapenemases, changes in membrane permeability and activity of membrane efflux pumps may have also contributed to carbapenem resistance in *K. pneumoniae* isolates from this study. Namely, it was observed that a mutant *ompK36* porin gene, like the one detected in these three carbapenemase-producing *K. pneumoniae* isolates, increases nonsusceptibility to this group of antibiotics (Wong et al., 2019). In addition, 7 mutations (P161R, G164A, F172S, R173G, L195V, F197I, and K201M) detected in transcriptional regulator gene *acrR* in C2 isolate were previously shown to highly increase the expression of a major multidrug efflux pump AcrAB-TolC (Sato et al., 2020) that effectively extrude multiple antimicrobials among which carbapenems (Chetri et al., 2019).

Moreover, the environmental K. pneumoniae isolates from our study shared similar determinants of resistance to other classes of antibiotics, which were previously described in K. pneumoniae clinical isolates from Croatia (Bedenić et al., 2018), such as the aminoglycoside and fluoroquinolone resistance gene aac(6')Ib-cr, disinfectant resistance genes oqxA and oqxB, sulfonamide resistance gene sul1, fosA encoding fosfomycin resistance and ESBL gene bla_{CTX-M}. Moreover, an amino acid substitution R256G was detected in PmrB protein sequence in 5a and M11 strains, which has been previously associated with colistin resistance in Enterobacteriaceae (Cheng et al., 2015), as well as in case of CL-resistant and carbapenemase-producing hospital K. pneumoniae in Croatia (D'Onofrio et al., 2020). Namely, variations in the PmrB protein, which is a part of the two-component regulatory system PmrA/PmrB enrolled in modification of lipopolysaccharide (LPS) structure, lead to the neutralization of its negative charge and consequently, the reduced susceptibility to cationic peptide antibiotics such as CL. However, as no increased MIC for CL was observed in these isolates, we speculate that the combined action of multiple mechanisms is likely needed to induce resistance to this antibiotic (Cheng et al., 2015). On the other hand, K. pneumoniae C2 was resistant to CL (MIC 32 µg/ml), yet no variations in PmrB were found. Limited number of studies have pointed to the underestimated role of the energy-driven efflux pump of peptide antibiotics in K. pneumoniae, involving two pumps, AcrAB-TolC and KpnEF (Binsker et al., 2021). It was observed that AcrR deficient mutant strains can successfully extrude polymyxin B, another peptide antibiotic, out of the cell using AcrAB-TolC pump (Padilla et al., 2010). More recent study of Naha et al. (2020) revealed that nonmutated and increasingly expressed RamA, a positive regulator of AcrAB-TolC pump, mediates alterations of LPS which along with the upregulation of the pump have contributed to the CL-resistant phenotype in clinical K. pneumoniae. RamA-mediated changes of lipid A moiety have been previously shown to decrease susceptibility to CL in this pathogen (De Majumdar et al., 2015). Therefore, it is likely that a functional AcrAB-TolC system and RamA could also be involved in nonsuceptibility to CL in K. pneumoniae C2, but this should be addressed more carefully in the future research.

Moreover, several common virulence-related genes were simultaneously detected in all three *K. pneumoniae* genomes, among which those coding the type 1 fimbriae (fimD, sfaG)

and ferrous ion transport (feoB). According to Struve et al. (2008) type 1 fimbriae are significantly enrolled in K. pneumoniae infections of urinary tract. The FeoB is the component of the major prokaryotic ferrous ion transport (Feo) system, and the main protein enabling the iron uptake through the lipid bilayer in almost all bacteria (Cartron et al., 2006; Lau et al., 2007). Nevertheless, in this study it was only detected in *K. pneumoniae*, which could be explained by the fact that a single species can adjust its iron import depending on the type of infection (acute or chronic) and the availability of iron in its environment (Cornelis and Dingemans, 2013). On the other hand, the C2 isolate, like E. coli, additionally harbored two virulence genes (pemK, and pemI) encoding the PemK/PemI type II toxinantitoxin system that has been previously described in hypermucoviscous carbapenem-resistant K. pneumoniae (Fu et al., 2018; Bleriot et al., 2020).

Citrobacter freundii CF1 Isolate

C. freundii CF1 isolate was affiliated to ST128, which was first described by Bonnin et al. (2020) in an isolate from the rectal swab of a French patient. To date, KPC-producing Citrobacter spp. have been isolated from hospital effluents (Zhang et al., 2012), river sediments (Xu et al., 2018), and the recreational areas (Montezzi et al., 2015), but none of them belonged to ST128. To the best of our knowledge, KPC-2-producing C. freundii of the ST128 lineage has not been previously reported. It was unexpected that this carbapenem-sensitive isolate resulted positive by CarbaNP test. However, although rare, there are previous reports of KPC-producing Enterobacterales showing unusual carbapenems susceptibility profile while testing positive by CarbaNP (Shinde et al., 2017; Cury et al., 2020), suggesting low gene expression.

Furthermore, we found that the strain CF1 also carried the $bla_{CMY-159}$, a variant gene identified only recently (Piotrowska et al., 2019) that encodes for the eponymous AmpC betalactamase of the CMY family intrinsic to Citrobacter spp. In a later study, this bla gene was detected in a Citrobacter sp. isolate resistant to cefotaxime, ceftazidime, cefepime and aztreonam, which is similar to the CF1 susceptibility profile. Overall, the majority of the 43 gene markers involved in the antibiotic resistance in this strain was associated with the activity of intrinsic antibiotic efflux including ATP-binding cassette (ABC), RND or MFS pumps. Notably, missense mutations of the AcrAB-TolC efflux pump regulators marR (Y137H) and soxR (T38S) which were detected in this strain were previously found to increase pump expression, leading to the multidrug resistance, among which to beta-lactams and ciprofloxacin (Al-Farsi et al., 2020). Giving the betalactam resistance profile of the CF1 isolate (sensitive to carbapenems and cefepime, but resistant to aztreonam and third generation cephalosporins) we can speculate that the mutation-driven expression of the AcrAB-TolC pump did not influence the activity against carbapenems. This would be a case when coincided with the membrane permeability defects resulted from porin loss or porin structural changes (Pages et al., 2008; Vardakas et al., 2012; Sadeghi, 2019), which have not been detected in CF1 isolate.

Moreover, among the 10 virulence-related genes detected in CF1 genome, the major virulence factors of this pathogen such as Shiga-like and heat-stable toxins, or the cholera toxin B subunit homolog (Bai et al., 2012) were not identified, thus we can speculate about the low virulence potential of this isolate (Pepperell et al., 2002).

CONCLUSION

This study reports the introduction of XDR and carbapenemaseproducing potentially virulent strains of Enterobacteriaceae into the Croatian marine environment through the submarine outfall of the treated wastewater located at a depth of 42 m. Among other antibiotic resistance and virulence determinants previously assigned exclusively to clinical strains, we report for the first time KPC-producing E. coli in coastal waters and the co-occurrence of bla_{KPC-2} and bla_{OXA-48} carbapenemase genes in this species. While *bla*_{OXA-48} was located on an IncL-type plasmids in this species, blaKPC-2 was harbored by recently described broad-host-range IncP6 resistance plasmids, providing first record of their circulation in E. coli and highlighting their importance in the epidemiology of this globally disseminated carbapenemase encoding gene. Leakage of these highly resistant strains into coastal waters through the submarine outlet is of serious concern as it provides a route for their continuous introduction into the marine environment and a reservoir for their further spread.

DATA AVAILABILITY STATEMENT

The genomic sequences are deposited in the NCBI Sequence Read Archive (BioProject number PRJNA768347) under accession numbers listed in **Table 2**. The sequence of plasmid pM12-KPC2 from *E. coli* was deposited in GenBank under accession number CP093216.

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AUTHOR CONTRIBUTIONS

MK and AM: conceptualization and writing—original draft preparation. MDŽ and IŠ: formal analysis. AN, AM, IGB, and MT: validation. MK and AN: investigation. IŠ, JK and AM: resources and funding acquisition. AM: data curation, visualization, and supervision. All authors writing--review and editing and contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022. 858821/full#supplementary-material

Supplementary Table S1 | 480 Molecular characteristics of eight wholegenome sequenced KPC-producing isolates.

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4. ADDITIONAL RESULTS AND DISCUSSION

To gain a deeper insight into our findings, we have carried out an additional analysis. This allowed us to better understand how submarine discharges affect the water quality of the entire water column. For this purpose, we compared our data with information collected at a reference site Split Chanel (SC), which is considered a location free of anthropogenic pressures [100]. These results are discussed in the next subsection 'Comparison of microbial community structure in coastal waters not affected by wastewater'. Moreover, to fortify the validity of our results, to gain new insights and to extend the current knowledge, additional analyses were carried out. This encompasses the simultaneous collection of samples from both influent and effluent of two distinct WWTPs, as well as from the locations of the corresponding submarine outfalls. We have provided detailed explanations of the used methods and the obtained results in the subsection 'Microbiome and resistome of WWTPs and submarine outfalls'. These analyses contribute to a more comprehensive and insightful perspective on our research.

4.1. Comparison of microbial community structure in coastal waters not affected by wastewater

4.1.1. Methodology

As part of a comprehensive investigation, the present chapter compares two geographically distinct locations: one that is affected by wastewater discharges (Katalinića Brig and Stupe submarine outfalls) and one that is unaffected (SC). For this comparative analysis, a dataset from another study was used [100] and compared with the partial datasets of the study presented in Paper II. This analytical endeavour not only enriches our understanding of the dynamics behind the impacts of wastewater discharge but also extends the scientific dissertation concerning the intricate reactions of coastal ecosystems to various factors. Beyond its wastewater-free status, another motivation for selecting the SC as a reference location was its shared physico-chemical characteristics with the nearby submarine outfalls, due to their proximity. We utilized datasets from the winter (March 2021) and summer (August 2021) seasons, collected from both the bottom (SCB) and surface (SCS) locations. To ensure comparability with our analysis, we selected a pair of field samples (from the study in Paper II), opting for those field campaigns that closely aligned with the study conducted by Dželalija

et al. in 2023. Specifically, we included datasets from February 2020 (winter season) and July 2020 (summer season) collected from the bottom (Katalinića Brig (KB) and Stpe (SB)) and surface (Katalinića Brig (KS) and Stpe (SS)) locations.

Consistent with our prior analysis approach (as outlined in Paper II), Dželalija et al., 2023 employed the same metagenomic DNA extraction kit (DNeasy PowerWater Kit) and conducted 16S rRNA amplicon sequencing on the extracted DNA. This strict adherence to methodology ensures the analogy of these datasets. For a comprehensive understanding of the methodology employed and the results obtained in the context of the mentioned study, refer to [100]. Analyses and data visualization in this study were done by Matlab R2021a.

4.1.1. Result and discussion

Differences in microbial community structure between the three selected locations are visible at the phylum level (Figure 4), although they are even more pronounced at the lower taxonomic levels (Figure 5). In winter (Figure 4a), Proteobacteria dominated at the SS and SCS locations (41.66% and 54.77%, respectively), while *Firmicutes* were most abundant at the KS location, with a frequency of 61.66%. Proteobacteria dominated at the SCB location too (39.37%), while at both bottom locations near the submarine outfalls dominated Firmicutes (62.20% at SB and 40.15% at KB). Both phyla, Proteobacteria and Firmicutes, together with Bacteroides are common human-gut members but are also found in coastal waters [100,101]. However, shift in its abundancies are noticed at the locations with different trophic status [100]. In this study, a decrease in *Firmicutes*, and an increase in *Proteobacteria* abundancy at the SCS and SCB locations support such findings. Furthermore, Verrucomicrobia together with Fusobacteria, are commonly found in WWTPs [102,103] and are noticed in our study during the winter only in locations near the submarine outfalls. Actinobacteria was more abundant at SCS (10.47%) compared with SS and KS locations (<6%), and compared with bottom locations, which supported the observed shift in microbial community toward the SCS and SCB locations, similar to the observations in another study [100]. In summer (Figure 4b), locations near Katalinića Brig submarine outfall and locations at Split channel shared the top two phyla. Proteobacteria dominated at KS and KB locations (77.28% and 73.48%, respectively), as at SCS and SCB (57.85% and 66.75%, respectively), followed by Cyanobacteria with abundancy >20% at SCS and SCB locations, and <20% at KS and KB

locations. SS and SB locations showed different microbiome communities. At both locations, Firmicutes were the most abundant phyla (59.58% and 79.87%, respectively), followed by Proteobacteria. Contrarily with previous findings, Actinobacteria shared similar abundancies (>5%) at SS, SB, SCS, and SCB, while at KS and KB were less abundant (<1%). Compared with winter observations, fewer Firmicutes and much more Proteobacteria was observed in KS, KB, SCS, and SCB locations, while at SS and SB location, the opposite finding was observed. The similarity of KS and KB with SCS and SCB locations during the summer could be due to the environmental conditions (higher sea and air temperature) which shaped microbial community. However, SS and SB locations have not been impacted by it, which could be due to the higher wastewater load, as higher FIB concentrations have been founded at this location (Paper II). Moreover, higher CDOM, Turbidity, and Chl a concentration with lower dissolved oxygen values is noticed at this location in this sampling month when compared with KB (Paper I), which support this explanation. Finally, as submarine outfalls do not discharge effluent continuously, but after the pool has been full, variation in microbiome analysis could be thus dependent on the overlapping of sampling time and effluent discharge. However, along with observed results, opposite findings among some phyla and the fact that some phyla contain both, environmental and human-related bacteria, suggest that lower taxonomic levels encompass much more information and are necessary to analyse in order to obtain relevant conclusions.

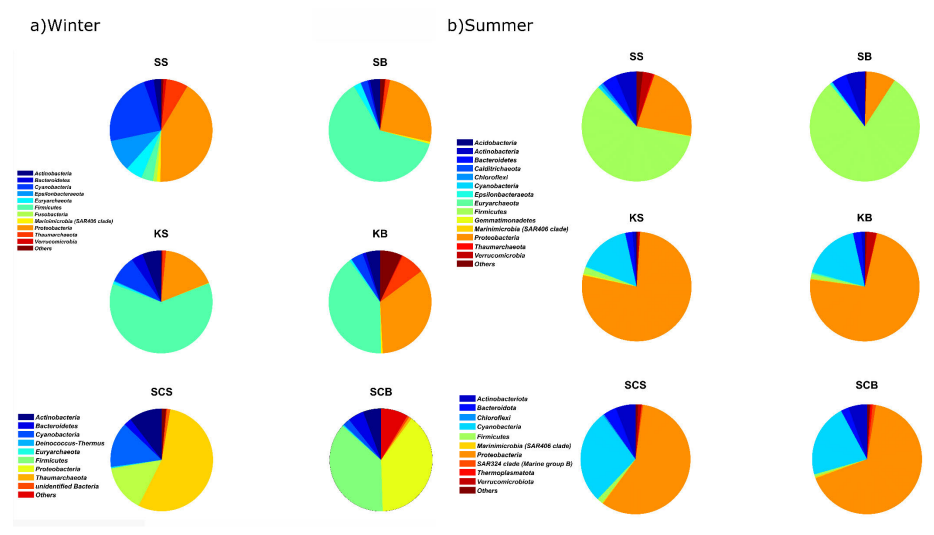


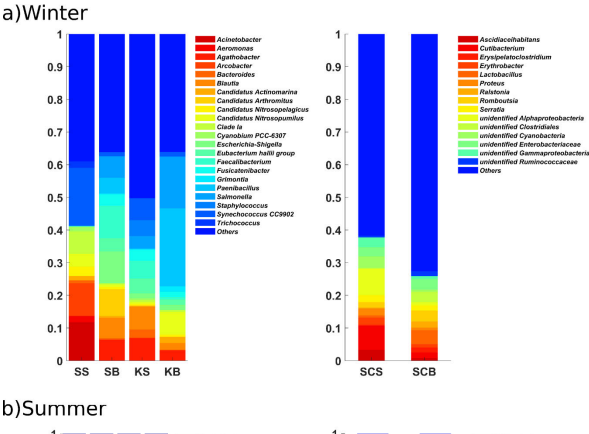
Figure 4. Relative abundance of the ten most abundant taxa at phylum level comprising the bacterial communities from the Stobreč (surface and bottom, SS and SB), Katalinića Brig (surface and bottom, KS and KB) WWTPs effluent sites, and Split channel (surface and bottom, SCS and SCB) reference location.

At the lower taxonomic levels (Figure 5), the differences in microbial community composition are even more visible. During the winter season (Figure 5a), there were no common genera among the selected locations, while in summer only two common genera were found at all selected stations (Synechococcus_CC9902 and Clade_Ia) (Figure 5b). In winter, Faecalibacterium and Escherichia-Shigella were the most abundant on SB (10%), Paenibacillus on KB (23.86%), while Synechococcus_CC9902 and Blautia were predominant on SS (17.67%) and KS (7%), respectively. Acinetobacter was second most abundant in SS (11.78%), with lower abundancies at other locations near submarine outfalls and not among the top 10 in SC locations. Moreover, locations near the submarine outfalls were enriched with human-related genera such as Acinetobacter, Aeromonas, Bacteroides, Blautia, Escherichia-Shigella, Faecalibacterium, Salmonella and Staphylococcus, while at SCS and SCB were not among the top 10 observed. At the SCS and SCB locations, among the top 10 bacterial genera, the most abundant (unidentified_Alphaproteobacteria) comprised abundancy about 8%, which highlights the higher diversity and lower richness of these locations compared with locations near the submarine outfalls. In summer, at SCS and SCB locations, the most abundant genus was Synechococcus_CC9902 (22.55% and 14.38%, respectively). Such increase in richness, along with other autochthonous marine bacteria (AEGEAN-169, Clade and SAR11 lineages) is seasonally driven as previously observed for this region [100,104]. Observation of such native marine bacteria was characteristic for summer at the locations near the submarine outfalls too, although with some differences. Lower abundancies of Synechococcus_CC9902 were observed at all four locations near the submarine outfalls compared with SC locations, while opposite findings were observed for Clade_Ia. Although at the locations near the submarine outfalls, only Clade_Ia was found, its abundance was higher at KS and KB locations (10.74% and 12.62%, respectively), compared with the SC locations (<6%). However, at the SS and SB locations, its abundancy was below 0.2%. In the summer, there were still observed human-related genera, as Bacteroides, Blautia, Dorea, Faecalibacterium, and Salmonella only at the locations near the submarine outfalls. At the SC locations, an increase in abundancy of Acinetobacter and Pseudomonas occurred.

In order to better understand the abundancy of the top 10 genera across sampling locations, a clustergram was obtained (Figure 6). Supporting the above discussed results, in the winter season (Figure 6a) dendrogram clustered submarine locations together, while SC locations clustered separately. However, in summer (Figure 6b), KS and KB locations were clustered together with SCS and SCB, while SS and SB locations clustered separately. This

analysis supported the obtained results at both taxonomic levels addressing the complex dynamics of microbial communities in response to locations and seasons.

This analysis reveals distinct differences in the structure of microbial communities among the selected locations, particularly evident at both the phylum and lower taxonomic levels. Winter and summer seasons exhibit distinct patterns, revealing the influence of submarine outfalls and seasonal changes on microbial populations. Seasonal shifts were observed in both, allochthonous and autochthonous bacteria. However, human-related genera are enriched near the outfalls, underscoring the potential impact of anthropogenic activities on coastal microbial ecosystems. However, future sampling including NGS sequencing together with physico-chemical parameters during the whole year at least are necessary to better understand complex dynamics of microbial communities in response to submarine outfall and seasonal changes. These findings suggest intricate ecological interactions influenced by seasonal and location-specific factors, offering a comprehensive view of the microbial communities' responses to different conditions and impacts.



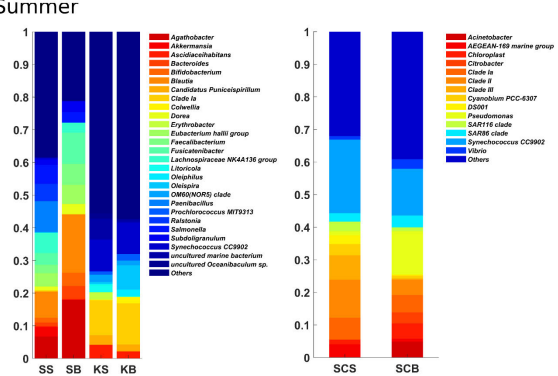
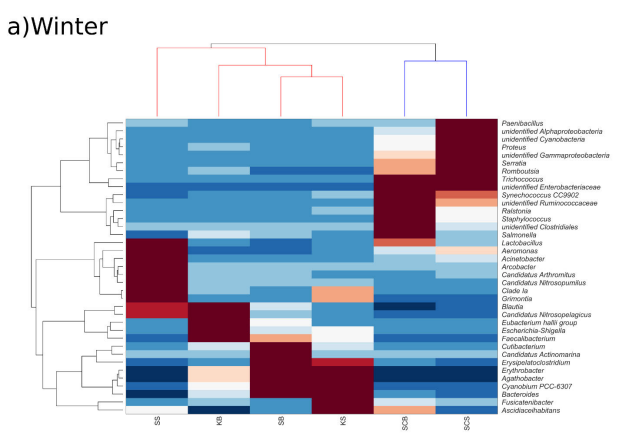


Figure 5. Relative abundance of the ten most abundant taxa at genus level comprising the bacterial communities from the Stobreč (surface and bottom, SS and SB), Katalinića Brig (surface and bottom, KS and KB) WWTPs effluent sites, and Split channel (surface and bottom, SCS and SCB) reference location.



b)Summer

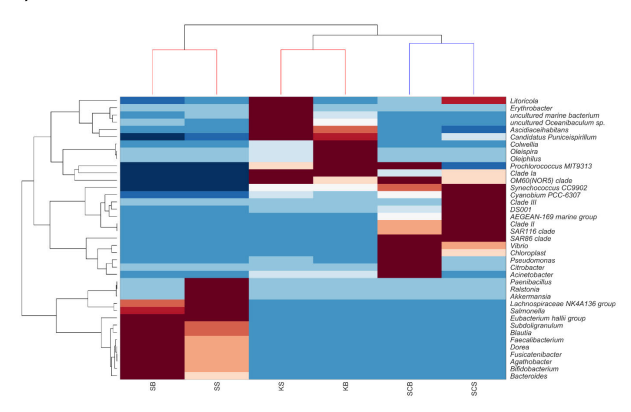


Figure 6. Clustergram (heatmap of relative abundancy at the genus level with dendogram) for winter (a) and summer (b) seasons over the sampling locations. Clustergram standardizes the values along the columns of data. The data has been standardized across all samples, so that the mean is 0 and the standard deviation is 1.

5. CONCLUSION AND FUTURE REMARKS

Within this doctoral dissertation, a comprehensive analysis was undertaken to examine the impact of submarine outfalls on the coastal marine environment's microbiome and resistome. Employing cutting-edge methods, including fast-profiling probes and advanced NGS techniques encompassing 16S rRNA and shotgun sequencing, this study aimed to address knowledge gaps and provide novel insights about the globally disseminated problem of antibiotic resistance. The initial phase (Paper I) focused on elucidating water column stability, a crucial determinant which influence the vertical movement of effluent plume, in tandem with physico-chemical parameters reflecting plume shape. Stratification's essential role in submerging effluents below the pycnocline was confirmed, underscoring the utility of fastprofiling probes. Simultaneously, samples were collected from submarine outfalls' water column bottom and surface locations for metagenomic analysis, as presented in Paper II, which clearly demonstrated the influence of submarine outfalls on the water column's microbial community. This resulted in the proliferation of nonindigenous and pathogenic bacteria, potentially harbouring AR determinants. Further investigations (Paper III) utilized plate-based methods to identify carbapenemase-producing Enterobacteriaceae strains, revealing their presence near the submarine outfall linked to hospital wastewater discharge. Strikingly, environmental carbapenemase-producing E. coli co-occurring bla_{KPC-2} and bla_{OXA-48} carbapenemase genes was detected for the first time, along with genes harbouring on IncP6 resistance plasmids. As this was the first record of IncP6 circulation in E. coli, this study moreover highlighted the importance of environmental monitoring in the epidemiology of this globally disseminated carbapenemase encoding gene. To support and expand our finding, additional results were made and presented in this thesis. Focusing on microbial community structure, we compared locations near the submarine outfalls with one location considered as free of wastewater impact. It reveals differences at both phylum and lower taxonomic levels, revealing the complex ecological dynamics influenced by seasonal and location-specific factors. Finally, an extensive shotgun analysis across both WWTPs (influent and effluent) and corresponding submarine outfalls provided insights into the intricate interplay between bacterial communities and ARGs, emphasizing the persistence and dissemination of AR within WWTPs and its subsequent release into the marine environment. This culmination of knowledge serves as a base for decision-makers, enabling a comprehensive strategies and maintenance of the delicate balance of marine ecosystems and public health. This thesis,

underscores the imperative of holistic approaches in unravelling the complex interactions between human activities and marine ecosystems.

However, it is imperative to acknowledge that the present findings represent a initial step, rather than a conclusive endpoint. This study underscores the potential paths for future research endeavours. Moreover, the nature of this investigation underscores the imperative of interdisciplinary collaboration. Future field campaigns should integrate advanced methodologies, including fast-profiling probes and remotely operated vehicles (ROVs), alongside with established microbiological and molecular techniques. These prospective investigations should encompass both within-WWTP and submarine outfall contexts. The deployment of sophisticated equipment such as probes and ROVs holds the promise of realtime insight into the precise localization, shape, and dynamics of effluent plumes. This, in tandem with assessments of vertical water column stability and seawater currents, promises an enhanced understanding of both vertical and horizontal dispersion patterns of effluent plumes. The integration of diverse seasonal observations could unveil disruptions to water column stability during the delicate human health risk periods of summer. Emerging tools such as NGS coupled in tandem with a spectrum of physiochemical measurements offer a platform for unravelling main stressors and driving forces that shape microbial community dynamics, as well as the distribution and diversity of ARGs. Expanding this investigation with sediment sampling could amplify our understanding and provides a more comprehensive knowledge about the complex interactions between aquatic environments and anthropogenic impacts. The consolidation of findings from diverse chapters seamlessly aligns with the One Health concept, revealing the intricate interdependencies between human, animal, and environmental health. This comprehensive investigation underscores the need for collaborative efforts across disciplines to mitigate the escalating challenges posed by antibiotic resistance and environmental impact.

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7. SUPPLEMENTAL MATERIAL

7.1. Supplemental material to Paper II



Fig. S1. The locations of sampling sites (WWTPs effluents of Stobreč and Katalinića Brig) in the coastal area of Split, central Adriatic Sea

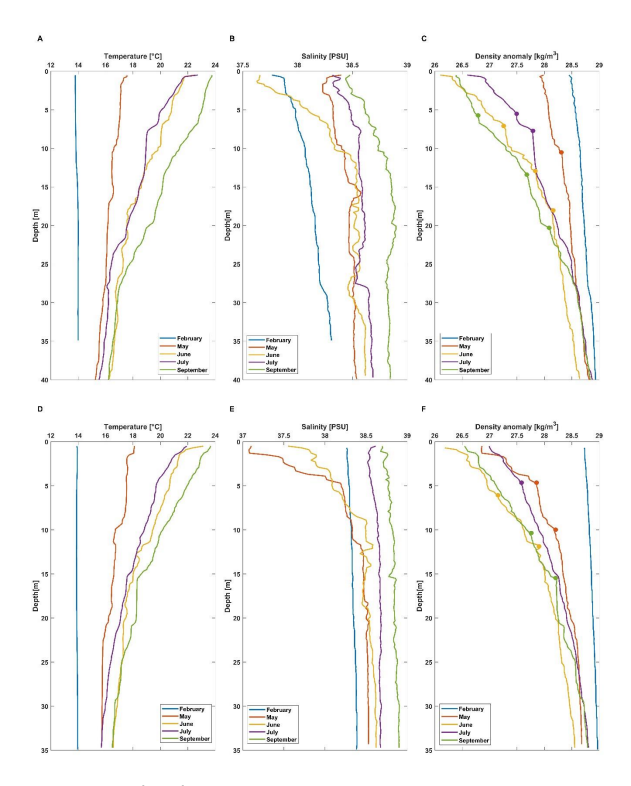


Fig. S2. Vertical profiles of temperature (A), salinity (B) and density anomaly (C) sampled at the Katalinića Brig location and at the Stobreč outfall (temperature (D), salinity (E) and density anomaly (F)) during the sampling period. Pycnocline depth is marked with a colored dot corresponding to the line for each month.

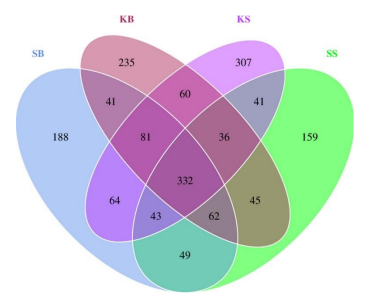


Fig. S3. Venn diagram based on OTUs diversity with values in overlapping parts representing OTUs common to more than one microbiome.

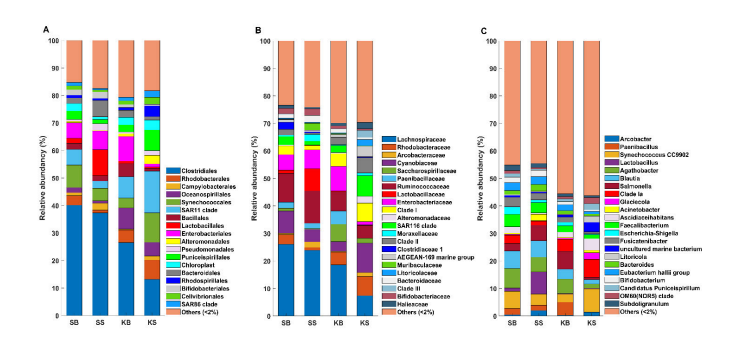


Fig. S4. Relative abundance (%) of bacterial taxa at order (A), family (B) and genus (C) levels in the studied samples. The abundance is expressed as the percentage of the individual taxa in the total number of reads.

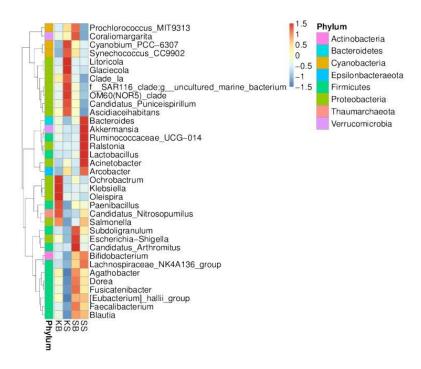


Fig. S5. Taxonomic abundance cluster heatmap plotted by sample name and the 35 most common genera. The absolute 'z' value represents the distance between the raw score and the mean of the standard deviation. 'Z' is negative when the raw score is below the mean, and *vice versa*.

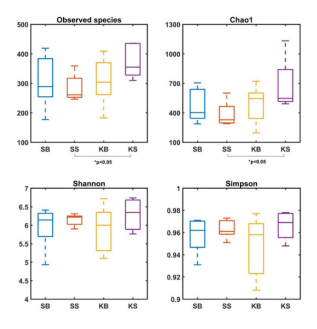


Fig. S6. Boxplots representing alpha diversity by community richness (observed species and Chao1 indices) and diversity (Shannon and Simpson indices) between bacterial communities at two effluent-receiving sites. SB (orange), SS (green), KB (blue), KS (purple). Alpha indices that were significantly different between groups (p<0.05) are marked with an asterisk.

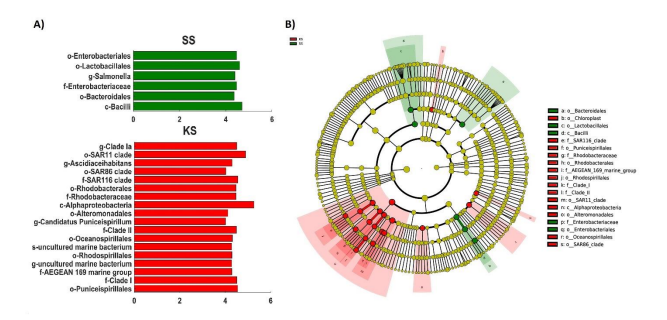


Fig. S7. Histogram of the LDA scores (A) and cladogram (B) showing the phylogenetic distribution of the microbial communities associated with SS and KS with LDA values of 2.0 or higher as determined by LEfSe. Red indicates KS and green indicates SS; yellow represents insignificant difference. The diameter of each circle is proportional to a taxon's abundance. Circles from inner region to outer region represent the phylogenetic levels from class to genus.

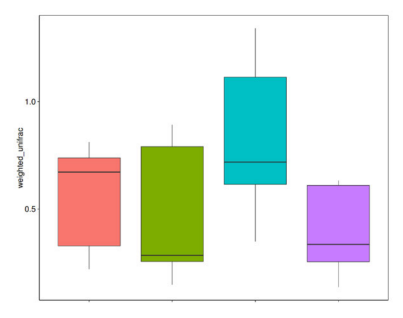


Fig. S8. Weighted UniFrac beta-diversity indices were statistically significant (p<0.05, Wilcox test) in case of KS and KB. SB (orange), SS (green), KB (blue), KS (purple)

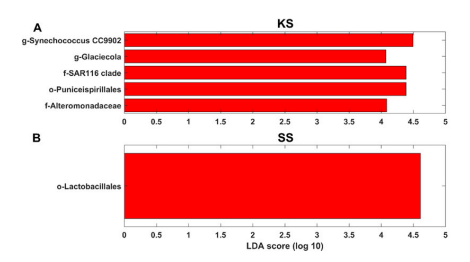


Fig. S9. Indicators of microbial taxa enriched in KS and SS, in comparison to KB and SB, with LDA values higher than 2.0.

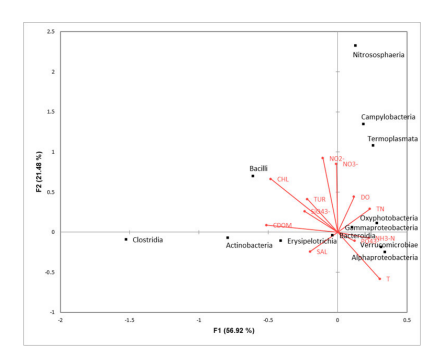


Fig. S10. Canonical correspondence analysis (CCA) plot relates the abundance of bacterial classes to environmental variables. T: temperature; SAL: salinity; CHL: chlorophyll a; TUR: turbidity; CDOM: colored dissolved organic matter; PO₄³⁻: phosphate anions; NH₄⁺: ammonium cation; NO₃⁻: nitrate anions; NO₂⁻: nitrite anions; SiO₄³⁻: silicate anions.

Table S1. Physico-chemical parameters of seawater samples at submarine outfalls: Stobreč(bottom and surface, SB and SS) and Katalinića Brig (bottom and surface, KB and KS)

DATE	SITE	T [°C]	SAL [PSU]	SIGMA [kg/m³]	DO [mg/l]	CHL a [ug/l]	CDOM [ppb]	TUR [NTU]	NO ₃ - [umol/L]	NO2 ⁻ [umol/L]	NH4 ⁺ [umol/L]	TN [umol/L]	PO ₄ ³ - [umol/L]
02/2020	SB	13.96	38.39	28.97	8.46	0.49	0.00	0.60	1.12	0.29	0.04	5.88	0.08
02/2020	SS	13.90	38.01	28.73	8.28	0.45	0.10	0.60	1.08	0.28	0.09	8.40	0.19
02/2020	KB	13.99	38.32	28.93	8.52	0.52	0.00	0.50	0.91	0.24	0.03	6.93	0.11
02/2020	KS	13.79	37.84	28.45	8.70	0.34	0.01	0.50	1.01	0.17	0.04	6.77	0.13
05/2020	SB	15.72	38.53	28.68	8.59	0.22	0.01	0.50	0.17	0.02	0.05	5.58	0.13
05/2020	SS	18.10	37.10	26.86	8.28	0.17	0.00	0.40	0.34	0.01	0.15	5.11	0.08
05/2020	KB	15.27	38.54	28.82	8.00	0.37	0.14	0.70	0.63	0.08	0.53	6.00	0.12
05/2020	KS	17.57	38.30	27.99	8.51	0.14	0.00	0.40	0.48	0.01	0.16	10.65	0.13
06/2020	SB	16.57	38.62	28.56	7.67	0.34	0.04	0.60	0.04	0.01	0.05	6.15	0.17
06/2020	SS	23.12	37.72	26.18	7.53	0.10	0.00	0.30	0.04	0.02	0.04	7.83	0.09
06/2020	KB	16.28	38.62	28.64	7.38	0.46	0.14	0.90	0.15	0.07	0.06	5.11	0.12
06/2020	KS	22.50	37.65	26.10	7.43	0.13	0.00	0.40	0.03	0.02	0.04	5.20	0.38
07/2020	SB	15.69	38.68	28.81	7.85	0.37	0.34	0.50	0.13	0.05	0.05	5.78	0.19
07/2020	SS	21.92	38.62	26.99	8.25	0.11	0.00	0.20	0.29	0.02	0.01	5.62	0.11
07/2020	KB	15.57	38.69	28.87	8.07	0.28	0.19	0.50	0.12	0.02	0.02	5.24	0.12
07/2020	KS	22.72	38.34	26.84	7.79	0.06	0.00	0.40	0.09	0.01	0.17	7.82	0.22
09/2020	SB	16.50	38.90	28.79	7.95	0.17	0.30	0.48	0.03	0.04	0.03	6.95	0.08
09/2020	SS	23.67	38.70	26.54	7.98	0.16	0.00	0.27	0.10	0.04	0.01	10.72	0.07
09/2020	KB	16.23	38.84	28.84	7.88	0.28	0.27	0.46	0.08	0.06	0.01	7.38	0.10
09/2020	KS	23.76	38.46	26.39	7.85	0.09	0.00	0.29	0.09	0.03	0.09	9.63	0.10

Abbreviations: T: temperature; SAL: salinity; Sigma: density anomaly; CHL: chlorophyll a; CDOM: color dissolved organic matter; TUR: turbidity; PO43-: phosphate anions; NH4+: ammonium cation; NO3-: nitrate anions; NO2-: nitrite anions; SiO43-: silicate anions

Table S2. Results of analysis of similarities (ANOSIM) and multi-response permutation procedure (MRPP) analysis evaluating the variation among and within bacterial communities from WWTP effluents Stupe (bottom and surface, SB and SS) and Katalinića Brig (bottom and surface, KB and KS)

	ANOSIM		MRPP						
Group	R-value	P-value	A	observed- delta	expected- delta	Significance			
SB-SS	-0.052	0.566	-0.01314	0.6933	0.6843	0.422			
KB-SS	-0.034	0.455	-0.004446	0.7154	0.7123	0.378			
KB-SB	-0.148	0.897	-0.04571	0.7583	0.7251	0.91			
KS-SS	0.428 ^a	0.03 ^b	0.1342 ^c	0.617	0.7127	0.013^{d}			
KS-SB	0.136 ^a	0.124	0.03692 ^c	0.6599	0.6852	0.139			
KS-KB	0.172 ^a	0.087	0.03587 ^c	0.6821	0.7074	0.149			

^a Inter-group variation is considered significant.

b, d Statistically significant (p -value <0.05)

^c Variations among groups are larger than variation within groups.

Table S3. PICRUSt

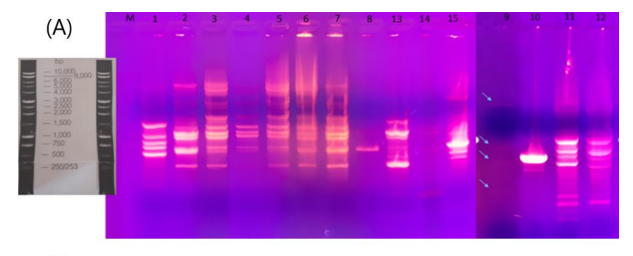
				SB.6 SS.6 KB.6 KS 6 SB.5 SS 5 KB.5 KS.5 SB.9 SS.9 KB 9 KS.9 SB.7 SS.7 KI	B.7 KS 7 SB.2 SS.2	2 KB.2 KS.2	stat SBvsSS	p SBvsSS	stat KBvsKS	p KBvsKS	stat SBvsKB	p SBvsKB	stat SSvsKS	p SSvsKS	stat SBvsKS	p SBvsKS	stat KBvsSS p	p KBvsSS
0	Bene verlan s	Aminoglycoside res stance genes	16S rRNA methylt ans e ases	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	.00 0.00 0.00 0.00	0.00 0.00	0.77	0.8	-1.01	0.35	1.01	0.37	0.92	0.39	0.91	0. 1	-1.7	0.1
1 1	Bene verlan s	Phenicol resistance genes	23S rRNA methylt ans e ase	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	.00 0.00 0.00 0.00	0.00 0.00	-05	0.60	0.60	0.57	0.27	0.80	1.6	0.1	0.93	0.38	-0.81	0.
2	Bene verlan s	Macrol de res s ance genes	23S rRNA methylt ans e ases	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	.00 0.00 0.00 0.00	0 0.00 0.01	-0 63	0.55	-0.19	0.85	0.55	0.60	0.60	0.57	0.18	0.86	-1.2	0.25
3	Bene varian s	Quino one resistance genes	ADP-ribosyl transferases	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0.00 0.00	-0 57	0.60	-0.09	0.93	0.29	0.79	0.25	0.81	-0.58	0.59	-0.29	0.78
	Bene varian s	Phenicol resistance genes	Acetyltrans erases	0.05 0.06 0.05 0.02 0.02 0.0 0 03 0.02 0.02 0.0 0.0 0 01 0.06 0.0 0	1.01 0.01 0.0 0.01	1 0.03 0.05	-0 07	0.95	1.02	0.3	0.58	0.58	1.67	0.1	1.58	0.15	-0.65	0 53
5	Bene sets	CAMP resistance modules	Cat onic an imicrob al peptide (CAMP) resistance, VraFG transporter MD:M00730]	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0 0.00 0.01	-0.9	0.6	-1.0	0.36	0.70	0.51	-0.99	0.38	-1.01	0.37	-0.98	0.36
6	Bene sets	CAMP resistance modules	Cat onic an imicrob al peptide (CAMP) resistance, dtABCD operon [MD:M00725]	0.0 0.0 0.03 0.01 0.00 0.08 0.01 0.0 0.03 0.07 0.02 0.01 0.0 0.03 0	1.01 0.01 0.02 0.06	6 0.01 0.05	-2. 6	0.0	-0.77	0. 7	1.27	0.25	2. 5	0.0	0.26	0.81	-3.86	0 01
7	Bene sets	CAMP resistance modu es	Cat onic an imicrob al peptide (CAMP) resistance, lysyl-phosphat dylglycerol L-PG) synthase MprF [MD:M00726]	0.01 0.00 0.00 0.01 0.00 0.02 0.01 0.01	1.01 0.01 0.00 0.00	0 0.00 0.02	-1 51	0.19	-2.1	0.08	0.01	0.99	0.52	0.62	-1.67	0.13	-1.62	0.17
8		CAMP resistance modu es	Cat onic an imicrob al peptide (CAMP) resistance, pro ease PgtE [MD:M007]	0.02 0.02 0.02 0.0 0.03 0.03 0.03 0.03 0		1 0.0 0.02	0.52	0.62	1.1	0.29	1.29	0.2	-0.9	0.37	-0.28	0.79	2 27	0.05
9		beta-Lactamase genes	Class A MTJ	0.00 0.00 0.00 0.00 0.00 0.01 0.01 0.00 0.00 0.01 0.01 0.01 0.00 0.00 0.00			-1 86	0.12	2.67	0.05	2.	0.07	2.3	0.07	0.73	0. 9	1 29	0.2
10	Bene varian s	beta-Lactamase genes	Class B MTj	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0.00 0.00	0.99	0.38	-0.07	0.95	0.31	0.77	-2.27	0.08	0.30	0.78	1 36	0.25
11	Bene verlan s	beta-Lactamase genes	Class C MTj	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0.00 0.00	-0 90	0.0	1.15	0.31	1.13	0.31	0.92	0. 0	-0.08	0.9	0 31	0.77
12	Bene varian s	beta-Lactamase genes	Class D MTJ	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0 0.01 0.00	-2.5	0.0	1.79	0.1	1.32	0.25	.03	0.01	1.18	0.28	0 21	0.8
13	Bene varian s	Trimethoprim resistance genes	Dihydrofolate reductases	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0 0.01 0.00	-1 55	0.17	1. 7	0.19	1.57	0.18	1.3	0.22	0.12	0.91	0. 2	0.69
1 1	Bene varian s	Sul onam de res s ance genes	Dihydropteroa e synthase	0.01 0.01 0.01 0.00 0.00 0.01 0.00 0.00	1.00 0.01 0.01 0.00	0 0.02 0.01	-1 56	0.17	1.62	0.18	1.30	0.25	2.1	0.09	0.8	0.	0 27	0.80
15	Bene sets	beta-Lactam res stance modules	Imipenem esistance, repress on of po in Op D [MD:M007 5]	0.02 0.01 0.02 0.0 0.05 0.02 0.05 0.0 0.0 0.02 0.03 0.0 0.01 0.0 0	0.0 0.0 0.03 0.0	0.03 0.01	0.36	0.73	0.33	0.75	0.8	0.3	-1.15	0.28	-0.66	0.53	1 22	0.26
16	Bene sets	beta-Lactam res stance modules	Methicil in res s ance [MD M00625]	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0 0.00 0.01	-0.10	0.92	-1.06	0.35	1.1	0.29	-0.99	0.38	-1.00	0.38	-0.91	0.0
17	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, et lux pump AbcA [MD M00700]	0.01 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00 0.00	0 0.01 0.01	-0 03	0.97	0.23	0.83	0.35	0.7	-0.06	0.96	-0.08	0.9	0 32	0.76
18	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, ef lux pump AcrEF-To C [MD M00696]	0.0 0.0 0.05 0.12 0.11 0.06 0.11 0.09 0.08 0.06 0.07 0.12 0.03 0.05 0	1.13 0.1 0.08 0.06	6 0.09 0.0	0.88	0. 2	-0.57	0.58	1.07	0.32	-2.67	0.05	-1. 9	0.18	2. 6	0.06
19	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, et lux pump AdeABC [PATH:map01501] [MD:M006_9]	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.01 0.01 0.00 0.01 0.00 0.00 0.00 0	1.00 0.00 0.00 0.00	2 0.00 0.00	-0.51	0.6	0.05	0.96	0.53	0.61	0.28	0.79	-0.5	0.61	-0.26	0.81
20	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, of lux pump BpeEF OprC [MD:M00698]	0.00 0.00 0.00 0.02 0.02 0.00 0.01 0.01	1.01 0.02 0.00 0.00	2 0.00 0.00	-00	0.97	4.1	0.29	0.18	0.86	-1.03	0.33	-1.11	0.30	0.12	0.90
21	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, ef lux pump MdtEF-TolC [MD:M00697]	0.03 0.02 0.03 0.13 0.12 0.0 0.12 0.10 0.09 0.03 0.0 0.13 0.02 0.03 0	1.13 0.1 0.06 0.06	6 0.0 0.03	1.30	0.25	-1.07	0.32	0.37	0.72	-3.1	0.03	-1.52	0.17	1 61	0.17
22	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, et lux pump MepA [MD:M00705]	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	1.00 0.00 0.00	0.00 0.00	0.09	0.93	0.69	0.52	0. 6	0.66	0.31	0.77	0. 9	0.6	0. 9	0.6
23	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, ef lux pump MexAB-OprM [MD:M00718]	0.06 0.05 0.06 0.23 0.25 0.08 0.22 0.28 0.21 0.08 0.09 0.23 0.0 0.08 0	1.29 0.26 0.08 0.32	2 0.10 0.05	0.10	0.93	-0.9	0.37	0.3	0.68	-1.39	0.20	-1. 1	0.19	0. 9	0.6
2	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, et lux pump MexCD-OprJ [MD:M00639]	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	0.00 0.00 0.00	0.00 0.00	0.	0.68	-1.39	0.20	0.13	0.90	-2.09	0.08	-1.66	0.1	0.51	0.63
25	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, of lux pump MexEF-OprN MD:M006 1]	0.00 0.00 0.00 0.01 0.01 0.00 0.01 0.01	.01 0.01 0.00 0.00	0.00 0.00	1.33	0.25	-0.99	0.35	0.59	0.57	-3.62	0.02	-1.86	0.10	1 67	0.17
26	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, ef lux pump MexJK OprM MD:M006 2]	0.01 0.01 0.01 0.0 0.0 0.0 0.02 0 0 0.06 0.05 0.02 0.02 0 03 0.01 0.02 0	1.03 0.0 0 01 0.09	9 0.02 0.01	-0 60	0.57	-1. 3	0.20	0.19	0.85	-0.28	0.79	-1.33	0.22	-0.5	0.62
27	Sene sets	Mu tipdrug resistance modules	Mult d ug res s ance, ef lux pump MexPQ-OpmE [MD:M00769]	0.01 0.01 0.01 0.01 0.00 0.02 0.01 0.01	.01 0.01 0.01 0.00	0 0.03 0.01	-16	0.15	1.93	0.12	1.65	0.16	2.18	0.09	0.51	0.63	0 52	0.62
28	Sene sets	Mu tipdrug resistance modules	Mult d ug res s ance, ef lux pump MexXY-OprM [MD M006 3]	0.01 0.01 0.01 0.02 0.02 0.02 0.02 0.05 0.03 0.02 0.02 0.02 0.01 0.02 0	1.02 0.02 0.01 0.07	7 0.02 0.01	-1 08	0.32	-1.07	0.3	0.25	0.81	0.35	0.7	-1.0	0.33	-1.05	0.35
29	Bene sets	Mu tipdrug resistance modules	Mult d ug res s ance, et lux pump NorB [MD:M00702]	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	.00 0.00 0.00 0.00	0 0.00 0.01	-0 07	0.95	-0.29	0.78	1.02	0.35	-0.80	0. 6	-0.83	0. 5	0.95	0.38
30	Sene sets	Mu tipdrug resistance modules	Mult d ug res s ance, ef lux pump QacA [MD:M0071]	0.01 0.01 0.01 0.01 0.01 0.01 0.01 0.01	.01 0.01 0.01 0.01	1 0.03 0.01	-188	0.10	1.79	0.1	1.72	0.15	2.2	0.07	0.09	0.93	0.8	0.
31	Sene sets	Mu tipdrug resistance modules	Mult d ug res s ance, epression of porin OmpF [MD:M007-6]	0.01 0.01 0.02 0.01 0.01 0.02 0.00 0.00	.01 0.00 0.01 0.00	0 0.0 0.01	-1. 0	0.21	2.09	0.10	1. 0	0.22	2.63	0.0	1.6	0.15	0. 9	0.6
32	Bene verlan s	Aminoglycoside res stance genes	N-Acetyltransferases	0.03 0.02 0.02 0.03 0.02 0.02 0.02 0.02	1.02 0.02 0.02 0.01	1 0.0 0.02	0.32	0.76	1.17	0.29	1.03	0.35	-0.0	0.97	0.29	0.78	1.18	0.28
33	Sene verlan s	Aminoglycoside res stance genes	O Nucleotidy transferases	0.0 0.0 0.0 0.01 0.01 0.0 0 02 0.01 0.01	.01 0.01 0.0 0.01	1 0.0 0.0	-0.36	0.73	1.51	0.17	0.15	0.89	2.09	0.07	1.75	0.12	-0. 9	0.6
3	Sene verlan s	Aminoglycoside res stance genes	O Phosphotransferases	0.03 0.02 0.0 0.02 0.02 0.02 0.02 0.01 0.01	1.01 0.01 0.02 0.01	1 0.03 0.02	0.51	0.62	2.13	0.08	0. 1	0.69	1.19	0.28	1.82	0.12	0.89	0.0
35	Bene varian s	Quino one resistance genes	Other	0.00 0.00 0.00 0.01 0.01 0.00 0.01 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0	1.01 0.00 0.00 0.00	0.00 0.00	0.9	0.39	-0.0	0.70	0.08	0.9	-2.18	0.07	-0.37	0.72	1 57	0.17
36	Sene verlan s	Aminoglycoside res stance genes	Other acetylt ans e ases	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00	.00 0.00 0.00 0.00	0.00 0.00	0.62	0.56	0.71	0.51	0.17	0.87	0.11	0.92	0.65	0.5	0.68	0.53
37	Bene varian s	Phenical resistance genes	Others	0.06 0.07 0.06 0.02 0.02 0.05 0.03 0.03 0.03 0.05 0.05 0.02 0.08 0.05 0	.01 0.01 0.05 0.02	2 0.0 0.07	-0.08	0.9	0.68	0.52	0.78	0. 6	1. 5	0.19	1.29	0.23	-0.93	0.38
38	Sene verlan s	Phenicol resistance genes	Phosphot ans e ase	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.01 0	0.00 0.00 0.00	0 0.01 0.00	-0.26	0.81	0.83	0. 5	0.55	0.61	0.7	0.50	0.93	0.39	0 35	0.7
39	Sene sets	Vancomycin res stance modules	Tetracyc ine resistance, efflux pump Tet38 [MD:M0070]	0.01 0.00 0.00 0.00 0.00 0.00 0.00 0.00	.00 0.00 0.00 0.00	0 0.01 0.01	0.05	0.96	0.38	0.72	0.12	0.91	0.31	0.76	0.35	0.7	0.16	0.88
0	Bene varian s	Quino one resistance genes	Transpor er	0.01 0.01 0.01 0.01 0.01 0.01 0.01 0.01	.01 0.01 0.01 0.01	1 0.03 0.01	-188	0.10	1.79	0.1	1.72	0.15	2.2	0.06	0.09	0.93	0.85	0.
1 1		Quino one resistance genes	Transpor ers	0.10 0.09 0.09 0.05 0.05 0.09 0.06 0.0 0.0 0.08 0.09 0.05 0.09 0.09 0			-0.26	0.80	1.18	0.28	0.06	0.96	1.50	0.18	1.30	0.23	-0.31	0.77
2	Bene sets	Vancomycin res stance modules	Vancomycin resistance, D-A a-D-Lac type IMD:M006511	0.18 0.19 0.17 0.07 0.09 0.13 0.09 0.06 0.16 0.1 0.12 0.13 0.19 0.1 0	.08 0.08 0.18 0.08	8 0.08 0.18	1.03	0.33	0.20	0.8	2.11	0.07	1.09	0.31	2.00	0.08	-1.0	0.33
	Bene sets		Vancomycin resistance, D-A a-D-Ser type IMD M006521	0.12 0.1 0.12 0.01 0.00 0.09 0.02 0.02 0.03 0.08 0.08 0.01 0.16 0.10 0			0.11	0.91	0.61	0.56	0.8	0. 3	1.50	0.17	1.38	0.21	-0.87	0. 1
			beta Lac am resistance. Bla system (MD:M00627)	0.08 0.08 0.07 0.0 0.0 0.06 0.03 0.03 0.06 0.06 0.06 0			0.35	0.7	-0.37	0.72	1.52	0.17	1.18	0.27	1.2	0.26	-1.55	0.16

h6	SB	SS	KB	KS	avg	avg*100 (%)
ADP-ribosyl transferases	0.001342229	0.001623039	0.001446958	0.001486454	0.00147467	0.147467001
Acetyltransferases	0.036832739	0.037536826	0.030911291	0.021517864	0.03169968	3.169967984
VraFG transporter	0.000251372	0.000306535	0.000171207	0.002939252	0.000917092	0.091709152
dltABCD operon	0.027018209	0.055880038	0.016331228	0.024057014	0.030821622	3.082162228
lysyl-phosphatidylglycerol (L-PG) synthase MprF	0.00541115	0.01158994	0.005387239	0.009437595	0.007956481	0.795648092
protease PgtE	0.022481244	0.019387717	0.030375753	0.024247784	0.024123125	2.412312455
Class A	0.002252081	0.004136278	0.006730991	0.001891764	0.003752779	0.375277867
Class B	0.000788467	0.000123782	0.000556001	0.000581753	0.000512501	0.051250083
Class C	0.0001799	0.000297554	0.000353645	0.000185848	0.000254237	0.025423662
Class D	0.001319883	0.002479593	0.002705788	0.000889434	0.001848675	0.184867463
Dihydrofolate reductases	0.001457877	0.002452775	0.002893114	0.001382149	0.002046479	0.204647883
Dihydropteroate synthase	0.006125511	0.010195775	0.011424358	0.005064495	0.008202535	0.820253488
Imipenem resistance, repression of porin OprD	0.027474768	0.024135991	0.036509284	0.03329835	0.030354598	3.035459827
Methicillin resistance	0.000206904	0.000220837	0.000107258	0.001832217	0.000591804	0.059180384
AbcA	0.002988691	0.00304596	0.003894927	0.003184482	0.003278515	0.327851523
AcrEF-ToIC	0.068704083	0.054457257	0.091101018	0.104207054	0.079617353	7.961735296
AdeABC	0.00244	0.004666247	0.003514679	0.003423486	0.003511103	0.351110299
BpeEF-OprC	0.006874798	0.007087075	0.007636009	0.01226429	0.008465543	0.8465543
MdtEF-ToIC	0.061207689	0.03476614	0.071874535	0.104273032	0.068030349	6.803034888
MepA	0.000334035	0.00030423	0.000605177	0.000205293	0.000362184	0.036218397
MexAB-OprM	0.127009013	0.120835282	0.153058435	0.208901989	0.15245118	15.24511799
MexCD-OprJ	0.000408128	0.00025167	0.000463581	0.001150779	0.00056854	0.056853965
MexEF-OprN	0.002903268	0.000780739	0.00449564	0.007271262	0.003862727	0.38627273
MexJK-OprM	0.022033512	0.032431813	0.023760085	0.037393704	0.028904779	2.89047786
MexPQ-OpmE	0.008220534	0.014245997	0.017434764	0.007202127	0.011775855	1.177585534
MexXY-OprM	0.016546445	0.029761695	0.017942367	0.025252785	0.022375823	2.237582305
NorB	0.001547211	0.001576409	0.002263197	0.002697599	0.002021104	0.202110424
QacA	0.008078168	0.011180356	0.014270761	0.007970031	0.010374829	1.037482904
repression of porin OmpF	0.008811087	0.0140996	0.017326298	0.004966068	0.011300763	1.13007632
N-Acetyltransferases	0.020964709	0.020004378	0.025756344	0.020135602	0.021715258	2.171525827
O-Nucleotidyltransferases	0.0299335	0.033340925	0.02847499	0.014508119	0.026564384	2.656438355
O-Phosphotransferases	0.024552221	0.021909598	0.026836898	0.016975348	0.022568516	2.256851643
Other	0.002533946	0.000825704	0.002688746	0.003282774	0.002332792	0.233279241
Other acetyltransferases	0.000274431	0.000201947	0.000306526	0.000194228	0.000244283	0.024428318
Others	0.047271576	0.048307619	0.037541667	0.02922992	0.040587696	4.058769572
Phosphotransferase	0.001859922	0.002195908	0.003030235	0.001305046	0.002097778	0.209777771
Tetracycline resistance, efflux pump Tet38	0.003489989	0.003393747	0.003813737	0.002683234	0.003345177	0.334517684
Transporter	0.008078073	0.011164182	0.014270393	0.007967346	0.010369998	1.036999818
Transporters	0.072000073	0.07647995	0.071094601	0.053980284	0.068388727	6.838872704
Vancomycin resistance, D-Ala-D-Lac type	0.160461379	0.133907208	0.108643111	0.103063231	0.126518732	12.65187322
Vancomycin resistance, D-Ala-D-Ser type	0.08343058	0.07933171	0.052300592	0.03327916	0.06208551	6.208551032
beta-Lactam resistance, Bla system	0.070650547	0.065374999	0.047391736	0.051761329	0.058794653	5.879465269

Table S4_KEGG BRITE ARGs

	Aminoglycoside resistance genes		Macrolide resistance genes	Multipdrug resistance modules	Phenicol resistance genes	Quinolone resistance genes	Sulfonamide resistance genes	Tetracycline resistance genes	Vancomycin resistance modules	beta-Lactam resistance modules	beta- Lactamase genes
g_Acinetobacter-SS	0.09	0.00	0.09	1.54	0.09	0 09	0.00	0.00	0 00	0.18	0.00
g Acinetobacter-KS	0.72	0.00	0.90	15.81	0.09	0.72	0.00	0.00	0 00	1 80	0.00
g Acinetobacter-KS			2.17	40.15		1 94		0.00		4 54	0.00
	1.92	0.00			2.25		0.00		0 00		
gAcinetobacter-KB	0.09	0.00	0.09	1.80	0.09	0 09	0.00	0.00	0 00	0.19	0.00
g_Arcobacter-SS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Arcobacter-KS	0.00	2.73	0.00	16.50	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Arcobacter-SB	0.00	9.50	0.00	57.22	0.00	0 00	0.00	0.00	0 00	0 00	0.00
gArcobacter-KB	0.00	0.19	0.00	1.14	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Bacteroides-SS	0.00	0.00	1.84	25.41	1.90	0 00	0.00	0.55	1.49	1 67	0.03
g_Bacteroides-KS	0.00	0.00	0.33	2.68	0.37	0 00	0.00	0.20	0 24	0 25	0.01
g_Bacteroides-SB	0.00	0.00	0.63	6.29	0.70	0 00	0.00	0.41	0 57	0 68	0.01
gBacteroides-KB	0.00	0.00	0.98	13.84	1.17	0 00	0.00	0.55	0 95	1 00	0.01
gBifidobacterium-SS	0.37	0.00	1.13	1.36	0.00	0 00	0.00	2.19	1.43	0 00	0.37
gBifidobacterium-KS	0.00	0.00	0.02	0.05	0.00	0 00	0.00	0.05	0 04	0 00	0.00
gBifidobacterium-SB	0.00	0.00	0.08	0.25	0.00	0 00	0.00	0.20	0.16	0 00	0.00
gBifidobacterium-KB	0.05	0.00	0.62	1.42	0.00	0 00	0.00	1.38	1 04	0 00	0.05
gClostridium_sensu_stricto_1-SS	0.18	0.05	0.02	0.00	0.00	0 00	0.00	0.03	0.76	0 20	0.00
gClostridium_sensu_stricto_1-KS	0.04	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0.14	0 04	0.00
gClostridium_sensu_stricto_1-SB	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
gClostridium_sensu_stricto_1-KB	0.00	0.01	0.00	0.00	0.00	0 00	0.00	0.01	0 01	0 00	0.00
gEscherichia-Shigella-SS	0.00	0.97	0.48	10.18	0.00	0 00	0.00	0.00	0 00	0 00	0.00
gEscherichia-Shigella-KS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
gEscherichia-Shigella-SB	0.00	0.05	0.03	0.53	0.00	0 00	0.00	0.00	0 00	0 00	0.00
gEscherichia-Shigella-KB	0.00	3.26	1.63	34.22	0.00	0 00	0.00	0.00	0 00	0 00	0.00
gHelicobacter-SS	0.09	0.00	0.00	0.73	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Helicobacter-KS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Helicobacter-SB	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Helicobacter-KB	0.09	0.00	0.00	0.76	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Klebsiella-SS	0.14	0.09	0.09	1.00	0.05	0 09	0.05	0.00	0 00	0.18	0.14
g_Klebsiella-KS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Klebsiella-SB	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Klebsiella-KB	3.37	2.25	2.25	24.72	1.12	2 25	1.12	0.00	0 00	4.49	3.37
g_Lactobacillus-SS	0.77	111.46	19.31	0.73	0.00	0 00	0.00	0.00	21.44	3 86	0.00
g_Lactobacillus-KS	0.07	0.47	0.14	0.00	0.00	0 00	0.00	0.00	0 04	0.18	0.00
g_Lactobacillus-SB	0.23	0.91	0.00	0.45	0.00	0 00	0.00	0.00	0 00	0 23	0.00
g_Lactobacillus-KB	0.05	1.32	0.09	0.09	0.00	0 00	0.00	0.00	0.19	0.14	0.00
g_Ochrobactrum-SS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0.00	0 00	0.00
g_Ochrobactrum-KS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Ochrobactrum-SB	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Ochrobactrum-KB	0.00	4.30	0.00	30.13	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g Pseudomonas-SS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
gPseudomonas-KS	2.39	2.53	0.70	34.89	0.70	0.70	0.00	0.70	1.40	12.11	0.00
gPseudomonas-SB	0.57	0.91	0.15	9.50	0.15	0.15	0.00	0.15	0.30	3.13	0.00
gPseudomonas-KB	0.57	0.91	0.15	2.75	0.15	0.15	0.00	0.15	0 09	0 95	0.00
gr seddomonas-RB g_Salmonella-SS	3.41	5.12		22.19	0.00	1.71	0.00	0.00		0 00	0.00
gsalmonella-ss g_Salmonella-KS	0.01	0.02	1.71 0.01	0.07	0.00	0 01	0.00	0.00	1.71 0 01	0 00	0.00
						0 01					
g_Salmonella-SB	0.04	0.06	0.02	0.28	0.00		0.00	0.00	0 02	0 00	0.00
g_Salmonella-KB	4.23	6.35	2.12	27.51	0.00	2.12	0.00	0.00	2.12	0 00	0.00
g_Staphylococcus-SS	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Staphylococcus-KS	0.00	0.47	0.00	0.18	0.00	0 00	0.00	0.00	0 04	0.11	0.00
g_Staphylococcus-SB	0.00	0.49	0.00	0.19	0.00	0 00	0.00	0.00	0 04	0.11	0.00
g_Staphylococcus-KB	0.00	0.00	0.00	0.00	0.00	0 00	0.00	0.00	0 00	0 00	0.00
g_Streptococcus-SS	0.15	0.83	0.00	0.00	0.00	0 00	0.00	0.05	0.45	0 00	0.00
g_Streptococcus-KS	0.01	0.11	0.00	0.00	0.00	0 00	0.00	0.04	0 04	0 00	0.00
g_Streptococcus-SB	0.11	1.13	0.00	0.00	0.00	0 00	0.00	0.30	0.47	0 00	0.00
gStreptococcus-KB	0.04	0.28	0.00	0.00	0.00	0 00	0.00	0.05	0.14	0 00	0.00

7.1. Supplemental material to Paper III



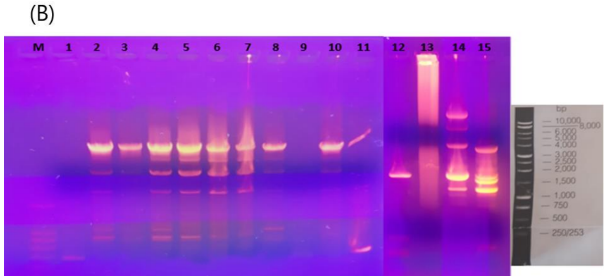


Figure S1. (A) BOX-PCR and (B) ERIC-PCR profiles of KPC-producing *Enterobacteriaceae*: *E. coli* isolates M20 (line 2), M19 (line 3), M16 (line 4), M13 (line 5), M14 (line 6), M15 (line 7), M17 (line 8), M12 (line 10), M18 (line 13); *K. pneumoniae* isolates 5a (line 1), M11 (line 9), C2 (line 11) and C1 (line 12); *C. freundii* isolates CF1 (line 14) and CF2 (line 15). 1 kb DNA ladder (Promega, USA) was used.

M12 E. coli (Table S1)

ResFinder 4.1

Resistance gene	Identity (%)	Alignment Length/Gene Length	Coverage (%)	Position in reference	Phenotype	Accession no.
dfrA14	100	474/474	100	1475	Trimethoprim resistance	KF921535
qnrVC4	100	657/657	100	1658	Quinolone resistance	GQ891757
cmlA1	99.68	1260/1260	100	11261	Phenicol resistance	M64556
mdf(A)	99.92	1233/1233	100	11234	Macrolide resistance	Y08743
blaKPC-2	100	882/882	100	1883	Beta-lactam resistance	AY034847
blaGES-2	100	864/864	100	1865	Beta-lactam resistance	AF326355
blaOXA-2	100	828/828	100	1829	Beta-lactam resistance	DQ112222
blaOXA-48	100	798/798	100	1799	Beta-lactam resistance	AY236073
blaOXA-10	100	801/801	100	1802	Beta-lactam resistance	J03427
					Alternate name; PSE-2	
ant(3")-Ii-aac(6')-IId	98.71	1393/1392	99.78	11394	Aminoglycoside resistance	AF453998
ant(3")-Ia	98.97	972/972	99.28	1973	Aminoglycoside resistance	X02340

PlasmidFinder 2.1

Database	Plasmid	Identity	Query / Template length	Accession number
enterobacteriaceae	Col440I	96.49	114 / 114	CP023920
enterobacteriaceae	IncC	100	417 / 417	JN157804
enterobacteriaceae	IncFIB(K)	99.82	560 / 560	JN233704
enterobacteriaceae	IncL	100	661 / 661	JN626286
enterobacteriaceae	IncP6	99.88	806 / 806	JF785550

MLST 2.0 (Multi-Locus Sequence Typing)

Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
adk	100	100	536	536	0	adk_64	2795
fumC	100	100	469	469	0	fumC_4	
gyrB	100	100	460	460	0	gyrB_5	
icd	100	100	518	518	0	icd_1	
mdh	100	100	452	452	0	mdh_8	
purA	100	100	478	478	0	purA_8	
recA	100	100	510	510	0	recA_6	

SerotypeFinder 2.0

Database	Gene	Serotype	Identity	Template / HSP length	Accession number
O_type	wzy	O21	100	1083 / 1083	EU694098
O_type	WZX	O21	100	1236 / 1236	EU694098
H_type	fliC	H27	91.17	1338 / 1461	AM231154

VirulenceFinder 2.0

Database	Virulence factor	Identity	Query / Template length	Protein function	Accession number
virulence_ecoli	gad	100	1401 / 1401	Glutamate decarboxylase	AP009240
virulence_ecoli	gad	99.93	1401 / 1401	Glutamate decarboxylase	AP010953
virulence_ecoli	lpfA	99.83	573 / 573	Long polar fimbriae	AP010953
virulence_ecoli	lpfA	91.45	524 / 573	Long polar fimbriae	CP002185
virulence_ecoli	lpfA	91.62	559 / 573	Long polar fimbriae	HE616528
virulence_ecoli	terC	99.02	714 / 714	Tellurium ion resistance protein	CP007491
virulence_ecoli	terC	98.96	966 / 966	Tellurium ion resistance protein	MG591698
virulence_ecoli	terC	100	1041 / 1041	Tellurium ion resistance protein	UGAE01000003

Plasmids detected by ARESdb cloud platform

Plasmid	Identity	Query / Template length	Accession number
Col(pHAD28)	91.60%	131 / 131	KU674895
Col(pHAD28)	92.50%	120 / 131	KU674895
Col440I	96.49%	114 / 114	CP023920
IncC	100.00%	417 / 417	JN157804
IncFIB(K)	99.82%	560 / 560	JN233704
IncL	100.00%	661 / 661	JN626286
IncP6	99.88%	806 / 806	JF785550

Virulence Factors detected by ARESdb cloud platform

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
virulence factors	ydeH	-	442 aa	98.6%	100%
virulence factors	ydaM	-	410 aa	98%	100%
virulence factors	yafQ	-	82 aa	98.8%	100%
virulence factors	soj	-	261 aa	99.6%	100%
virulence factors	sfaH	-	304 aa	97.4%	100%
virulence factors	potD	-	348 aa	91.1%	100%
virulence factors	pemK	-	110 aa	96.4%	100%
virulence factors	pemI	-	85 aa	96.5%	100%
virulence factors	nagA	-	139 aa	98.6%	94.6%
virulence factors	mqsA	-	131 aa	100%	100%
virulence factors	mqo	-	548 aa	99.3%	100%
virulence factors	manZ	-	269 aa	98.5%	100%
virulence factors	manX	-	144 aa	97.9%	100%
fimbria	LpfA	-	190 aa	100%	100%
virulence factors	ldrD	-	91 aa	94.5%	100%
virulence factors	fliZ	-	108 aa	98.1%	99.1%
virulence factors	fliZ	-	62 aa	96.8%	100%
virulence factors	fliY	-	266 aa	92.1%	100%
virulence factors	fliQ	-	89 aa	93.3%	100%
virulence factors	fliI	-	456 aa	95.6%	100%
virulence factors	fliG	-	332 aa	96.7%	100%
virulence factors	fliA	-	239 aa	99.6%	100%
virulence factors	flhA	-	692 aa	94.4%	100%
virulence factors	fimD	-	517 aa	98.6%	100%
virulence factors	fdeC	-	1416 aa	93.6%	100%
virulence factors	EspX1	-	473 aa	91.5%	100%
virulence factors	epsJ	-	300 aa	98%	100%
virulence factors	$\it elfG$	-	357 aa	100%	100%
virulence factors	cstA	-	701 aa	92.4%	100%

Antibiotic resistance genes detected by ARESdb cloud platform

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
ABC efflux pump	<i>YojI</i>	-	547 aa	100%	100%
putative efflux transport	yhe I	-	590 aa	98.1%	100%
Vga	Vga(C)	-	74 aa	94.6%	97.4%
efflux	TolC	-	495 aa	100%	100%
beta-lactamase TEM	TEM-160	-	190 aa	98.9%	66.4%
efflux regulation	SoxR	R4K	152 aa	93.4%	100%
RND efflux transporter	SilC	-	461 aa	99.3%	100%
RND efflux transporter	SilB	-	430 aa	99.3%	100%
RND efflux transporter	SilA	-	1048 aa	98.8%	100%
target protection	Rpsl	-	124 aa	93.5%	100%
regulation of resistance	rpoS	-	172 aa	100%	100%
MFS efflux transporter	RosB	-	558 aa	91.9%	100%
Qnr	QnrVC4	-	218 aa	100%	100%
multidrug efflux SMR transporter	QacF	-	110 aa	99.1%	100%
Pmr	PmrL	-	660 aa	100%	100%
Pmr	PmrF	-	322 aa	100%	100%
Pmr	PmrE	-	388 aa	99%	100%
Pmr	PmrC	-	480 aa	99.2%	87.8%
efflux regulation	PhoQ	-	486 aa	99.4%	100%
ABC efflux pump	PatA	-	459 aa	99.8%	100%
DNA topoisomerase subunit	parC	T57S	752 aa	94.9%	100%
beta-lactamase OXA	OXA-48	-	265 aa	100%	100%
beta-lactamase OXA	OXA-2	-	275 aa	100%	100%
beta-lactamase OXA	OXA-10	-	266 aa	100%	100%
regulation of resistance	ompR	-	239 aa	100%	100%
porin	OmpC	-	368 aa	96.5%	100%
ABC efflux pump	MsrB	-	137 aa	100%	100%
ABC efflux pump	MsbA	-	582 aa	100%	100%
macrolide phosphotransferase	Mph(B)	-	157 aa	98.1%	99.4%
compensation mechanism	MgrB	-	46 aa	100%	97.9%
regulation of resistance	MerR	-	144 aa	100%	100%
MFS efflux transporter	MdtP	-	488 aa	97.7%	100%

MFS efflux transporter	MdtO	-	683 aa	99.6%	100%
MFS efflux transporter	MdtN	-	343 aa	100%	100%
MFS efflux transporter	MdtM	-	410 aa	98%	100%
MFS efflux transporter	MdtH	-	402 aa	100%	100%
MFS efflux transporter	MdtG	-	408 aa	99.8%	100%
RND efflux transporter	MdtF	-	1037 aa	99.7%	100%
RND efflux transporter	MdtE	-	385 aa	99.7%	100%
RND efflux transporter	MdtC	-	1024 aa	99.7%	99.9%
RND efflux transporter	MdtB	-	1040 aa	99.6%	100%
RND efflux transporter	MdtA	-	415 aa	99.3%	100%
ABC efflux pump	mdlB	-	593 aa	99.5%	100%
MFS efflux transporter	MdfA	-	410 aa	99.8%	100%
efflux regulation	MarR	G103S, Y137H	144 aa	97.9%	100%
efflux regulation	MarA	_	129 aa	99.2%	100%
ABC efflux pump	MacA	-	371 aa	99.7%	100%
fimbria	LpfA	-	190 aa	100%	100%
beta-lactamase KPC	KPC-2	-	293 aa	100%	100%
efflux regulation	KdpE	-	225 aa	99.1%	100%
efflux regulation	H-NS	-	137 aa	100%	100%
DNA topoisomerase subunit	gyrA	S83L	875 aa	99.9%	100%
putative efflux transport	gspA	-	270 aa	98.1%	100%
influx	GlpT	G206A	423 aa	93.1%	93.8%
influx	GlpT	E448K	452 aa	98.2%	100%
beta-lactamase GES	GES-2	-	287 aa	100%	100%
efflux regulation	GadX	-	274 aa	98.5%	100%
efflux regulation	GadW	-	242 aa	93.4%	100%
efflux regulation	EvgS	-	1065 aa	99.2%	89%
efflux regulation	EvgA	-	204 aa	100%	100%
beta-lactamase ESC	ESC-148	-	420 aa	98.3%	100%
MFS efflux transporter	EmrY	-	512 aa	99.6%	100%
MFS efflux transporter	EmrK	-	372 aa	98.1%	96.1%
multidrug efflux SMR transporter	EmrE	-	110 aa	98.2%	100%
MFS efflux transporter	EmrD	-	394 aa	99.7%	100%
MFS efflux transporter	<i>EmrB</i>	-	512 aa	100%	100%
MFS efflux transporter	EmrA	-	390 aa	99.7%	100%
Dfr	DfrA14	-	160 aa	99.4%	100%

influx	cycA	S263A	467 aa	98.3%	100%
putative efflux transport	cusR	-	227 aa	95.2%	100%
putative efflux transport	cusF	-	110 aa	97.3%	100%
efflux regulation	CRP	-	210 aa	99.5%	100%
efflux regulation	CpxA	-	457 aa	99.1%	100%
MFS efflux transporter	CmlA5	-	437 aa	99.8%	100%
BaeSR	BaeS	-	467 aa	100%	100%
BaeSR	BaeR	-	240 aa	100%	100%
undecaprenyl-diphosphatase	BacA	-	273 aa	100%	100%
beta-lactamase regulation	argR	-	156 aa	94.2%	100%
beta-lactamase regulation	ampR	-	105 aa	94.3%	85.4%
beta-lactamase ampC	ampH	-	385 aa	99.5%	100%
beta-lactamase ampC	ampC1	-	434 aa	99.8%	100%
efflux regulation	AcrS	-	220 aa	99.5%	100%
RND efflux transporter	AcrF	-	1034 aa	99.3%	100%
RND efflux transporter	AcrE	-	385 aa	100%	100%
RND efflux transporter	AcrD	-	1037 aa	99.9%	100%
RND efflux transporter	AcrB	-	1049 aa	100%	100%
RND efflux transporter	AcrA	-	397 aa	100%	100%
ANT(3")-Ia	aadA11	-	269 aa	100%	95.7%
ANT(3")-Ia	aadA	-	275 aa	100%	99.6%
AAC(6')-I	<i>AAC</i> (6')- <i>Ib</i>	-	192 aa	99%	94.6%

M14 E. coli (Table S2)

ResFinder 4.1 **Resistance Identity** Alignment **Coverage Position** Phenotype Accession Length/Gene gene (%)(%)in no. Length reference dfrA14 100 474/474 100 1..475 Trimethoprim resistance KF921535 1..658 100 657/657 100 Quinolone resistance qnrS1 AB187515 97.67 99.83 Fluoroquinolone and aminoglycoside resistance *aac*(6')-*Ib*-599/600 5..604 DQ303918 99.88 *aph*(3")-*Ib* 804/804 100 1..805 Aminoglycoside resistance AF321551 99.92 Macrolide resistance Y08743 mdf(A)1233/1233 100 1..1234 blaKPC-2 100 882/882 100 1..883 Beta-lactam resistance AY034847 blaOXA-100 798/798 100 1..799 Beta-lactam resistance AY236073 48 100 blaGES-1 864/864 100 1..865 Beta-lactam resistance HQ170511 Beta-lactam resistance Alternate name; PSE-2 100 801/801 100 1..802 J03427 blaOXA-10 sul2 100 816/816 100 1..817 Sulphonamide resistance AY034138

PlasmidFinder 2.1

Database	Plasmid	Identity (%)	Query / Template length	Accession number
enterobacteriaceae	IncFIB(K)	99.82	560 / 560	JN233704
enterobacteriaceae	IncFII	99.62	261 / 261	AY458016
enterobacteriaceae	IncL	100	661 / 661	JN626286
enterobacteriaceae	IncN	99.81	514 / 514	AY046276
enterobacteriaceae	IncP6	99.88	806 / 806	JF785550
enterobacteriaceae	IncX5	100	576 / 576	MF062700
enterobacteriaceae	IncY	98.95	765 / 765	K02380

VirulenceFinder 2.0

Database	Virulence factor	Identity	Query / Template length	Protein function	Accession number
virulence_ecoli	gad	100	1401 / 1401	Glutamate decarboxylase	AP009239
virulence_ecoli	gad	99.93	1401 / 1401	Glutamate decarboxylase	AP009240
virulence_ecoli	<i>lpfA</i>	99.83	573 / 573	Long polar fimbriae	AP010953
virulence_ecoli	<i>lpfA</i>	91.45	524 / 573	Long polar fimbriae	AP010953
virulence_ecoli	<i>lpfA</i>	91.62	559 / 573	Long polar fimbriae	CP002185
virulence_ecoli	terC	99.02	714 / 714	Tellurium ion resistance protein	HE616528
virulence_ecoli	terC	98.96	966 / 966	Tellurium ion resistance protein	CP007491
virulence_ecoli	terC	100	1041 / 1041	Tellurium ion resistance protein	MG591698

MLST 2.0 (Multi-Locus Sequence Typing)

Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
adk	100	100	536	536	0	adk_64	2795
fumC	100	100	469	469	0	fumC_4	
gyrB	100	100	460	460	0	gyrB_5	
icd	100	100	518	518	0	icd_1	
mdh	100	100	452	452	0	mdh_8	
purA	100	100	478	478	0	purA_8	
recA	100	100	510	510	0	recA_6	

SerotypeFinder 2.0

Database	Gene	Serotype	Identity	Template / HSP length	Accession number
O_type	wzy	O21	100	1083 / 1083	EU694098
O_type	WZX	O21	100	1236 / 1236	EU694098
H_type	fliC	H27	91.17	1338 / 1461	AM231154

Plasmids detected by ARESdb cloud platform

Plasmid	Identity	Query / Template length	Accession number
Col440I	95.61%	114 / 114	CP023920
Col440I	94.74%	114 / 114	CP023920
Col440II	91.10%	281 / 282	CP023921
IncFIB(K)	99.82%	560 / 560	JN233704
IncFII	99.62%	261 / 261	AY458016
IncFII(Yp)	92.64%	231 / 230	CP000670
IncL	100.00%	661 / 661	JN626286
IncN	99.81%	514 / 514	AY046276
IncP6	99.88%	806 / 806	JF785550
IncX5	100.00%	576 / 576	MF062700
IncY	98.95%	765 / 765	K02380

Virulence Factors detected by ARESdb cloud platform

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
virulence factors	ydeH	-	442 aa	98.6%	100%
virulence factors	ydaM	-	410 aa	98%	100%
virulence factors	sfaH	-	304 aa	97.4%	100%
virulence factors	potD	-	348 aa	91.1%	100%
virulence factors	pemK	-	110 aa	100%	100%
virulence factors	pemI	-	85 aa	100%	100%
virulence factors	nagA	-	139 aa	98.6%	94.6%
virulence factors	mqsA	-	131 aa	100%	100%
virulence factors	mqo	-	548 aa	99.3%	100%
virulence factors	manZ	-	269 aa	98.5%	100%
virulence factors	manX	-	144 aa	97.9%	100%
virulence factors	ldrD	-	91 aa	94.5%	100%
virulence factors	fliZ	-	108 aa	98.1%	99.1%
virulence factors	fliZ	-	62 aa	96.8%	100%
virulence factors	fliY	-	266 aa	92.1%	100%
virulence factors	fliQ	-	89 aa	93.3%	100%
virulence factors	fliI	-	456 aa	95.6%	100%
virulence factors	fliG	-	332 aa	96.7%	100%
virulence factors	fliA	-	239 aa	99.6%	100%
virulence factors	flhA	-	692 aa	94.4%	100%
virulence factors	fimD	-	517 aa	98.6%	100%
virulence factors	EspX1	-	473 aa	91.5%	100%
virulence factors	epsJ	-	300 aa	98%	100%
virulence factors	$\it elfG$	-	357 aa	100%	100%
virulence factors	cstA	-	701 aa	92.4%	100%
fimbria	LpfA	-	190 aa	100%	100%

Antibiotic resistance genes detected by ARESdb cloud platform

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
Vga	Vga(C)	-	74 aa	94.6%	97.4%
undecaprenyl-diphosphatase	BacA	-	273 aa	100%	100%
target protection	Rpsl	-	124 aa	93.5%	100%
Sul	Sul2	-	282 aa	98.9%	98.3%
RND efflux transporter	MdtF	-	1037 aa	99.7%	100%
RND efflux transporter	MdtE	-	385 aa	99.7%	100%
RND efflux transporter	MdtC	-	1024 aa	99.7%	99.9%
RND efflux transporter	MdtB	-	1040 aa	99.6%	100%
RND efflux transporter	MdtA	-	415 aa	99.3%	100%
RND efflux transporter	AcrF	-	1034 aa	99.3%	100%
RND efflux transporter	AcrE	-	385 aa	100%	100%
RND efflux transporter	AcrD	-	1037 aa	99.9%	100%
RND efflux transporter	AcrB	-	1049 aa	100%	100%
RND efflux transporter	AcrA	-	397 aa	100%	100%
regulation of resistance	rpoS	-	172 aa	100%	100%
regulation of resistance	ompR	-	239 aa	100%	100%
Qnr	QnrS7	-	218 aa	100%	100%
putative efflux transport	yheI	-	590 aa	98.1%	100%
putative efflux transport	gspA	-	270 aa	98.1%	100%
putative efflux transport	cusR	-	227 aa	95.2%	100%
putative efflux transport	cusF	-	110 aa	97.3%	100%
porin	OmpC	-	368 aa	96.5%	100%
Pmr	PmrL	-	660 aa	99.8%	100%
Pmr	PmrF	-	322 aa	100%	100%
Pmr	PmrE	-	388 aa	99%	100%
multidrug efflux SMR transporter	EmrE	-	110 aa	98.2%	100%
MFS efflux transporter	RosB	-	558 aa	91.9%	100%
MFS efflux transporter	MdtM	-	410 aa	98%	100%
MFS efflux transporter	MdtH	-	402 aa	100%	100%
MFS efflux transporter	MdtG	-	408 aa	99.8%	100%
MFS efflux transporter	MdfA	-	410 aa	99.8%	100%
MFS efflux transporter	EmrY	-	512 aa	99.6%	100%
MFS efflux transporter	EmrK	-	372 aa	98.1%	96.1%

MFS efflux transporter	EmrD	-	394 aa	99.7%	100%
MFS efflux transporter	EmrB	-	512 aa	100%	100%
MFS efflux transporter	EmrA	-	390 aa	99.7%	100%
macrolide phosphotransferase	Mph(B)	-	157 aa	98.1%	99.4%
influx	GlpT	G206A	423 aa	93.1%	93.8%
influx	GlpT	E448K	452 aa	98.2%	100%
influx	cycA	S263A	467 aa	98.3%	100%
fimbria	LpfA	-	190 aa	100%	100%
efflux regulation	SoxR	R4K	152 aa	93.4%	100%
efflux regulation	PhoQ	-	486 aa	99.4%	100%
efflux regulation	MarR	G103S, Y137H	144 aa	97.9%	100%
efflux regulation	MarA	-	129 aa	99.2%	100%
efflux regulation	KdpE	-	225 aa	99.1%	100%
efflux regulation	H-NS	-	137 aa	100%	100%
efflux regulation	GadX	-	274 aa	98.5%	100%
efflux regulation	GadW	-	242 aa	93.4%	100%
efflux regulation	EvgS	-	1197 aa	99.3%	100%
efflux regulation	EvgA	-	204 aa	100%	100%
efflux regulation	CRP	-	210 aa	99.5%	100%
efflux regulation	CpxA	-	457 aa	99.1%	100%
efflux regulation	AcrS	-	220 aa	99.5%	100%
efflux	TolC	-	495 aa	100%	100%
DNA topoisomerase subunit	parC	T57S	752 aa	95.1%	100%
DNA topoisomerase subunit	gyrA	S83L	875 aa	99.9%	100%
Dfr	DfrA14	-	160 aa	99.4%	100%
compensation mechanism	MgrB	-	46 aa	100%	97.9%
beta-lactamase TEM	TEM-160	-	190 aa	98.9%	66.4%
beta-lactamase regulation	argR	-	156 aa	94.2%	100%
beta-lactamase regulation	ampR	-	105 aa	94.3%	85.4%
beta-lactamase OXA	OXA-48	-	265 aa	100%	100%
beta-lactamase OXA	OXA-10	-	266 aa	100%	100%
beta-lactamase KPC	KPC-2	-	293 aa	100%	100%
beta-lactamase GES	GES-1	-	287 aa	100%	100%
beta-lactamase ESC	ESC-148	-	420 aa	98.3%	100%
beta-lactamase ampC	ampH	-	385 aa	99.7%	100%
beta-lactamase ampC	ampC1	-	434 aa	99.8%	100%
BaeSR	BaeS	-	467 aa	100%	100%

BaeSR	BaeR	-	240 aa	100%	100%
APH(3")	<i>APH</i> (3")-	-	277 aa	99.6%	91.7%
	<i>Ib</i>				
ABC efflux pump	YojI	-	547 aa	100%	100%
ABC efflux pump	PatA	-	459 aa	99.8%	100%
ABC efflux pump	MsrB	-	137 aa	100%	100%
ABC efflux pump	MsbA	-	582 aa	100%	100%
ABC efflux pump	mdlB	-	593 aa	99.5%	100%
ABC efflux pump	MacA	-	371 aa	99.7%	100%
AAC(6')-I	<i>AAC</i> (6')- <i>Ib</i>	-	192 aa	100%	94.6%

M17 E. coli (Table S3)

ResFinder 4.1								
Resistance	Identity	Alignment	Coverage	Position	Phenotype	Accession no.		
gene	(%)	Length/Gene	(%)	in				
		Length		reference				
aac(6')-Ib-	100	600/600	100	1601	Fluoroquinolone and	DQ303918		
cr					aminoglycoside resistance			
aadA16	99.65	846/846	100	1847	Aminoglycoside resistance	EU675686		
blaKPC-2	100	882/882	100	1883	Beta-lactam resistance	AY034847		
aac(6')-Ib-	100	600/600	100	1601	Fluoroquinolone and	DQ303918		
cr					aminoglycoside resistance			
qnrB6	100	645/645	100	1646	Quinolone resistance	EF523819		
mdf(A)	99.92	1233/1233	100	11234	Macrolide resistance	Y08743		
sul1	100	840/840	100	1841	Sulphonamide resistance	U12338		
ARR-3	99.45	543/543	100	1544	Rifampicin resistance	FM207631		
ARR-3	100	453/453	100	1454	Rifampicin resistance	JF806499		
dfrA27	100	474/474	100	1475	Trimethoprim resistance	FJ459817		

PlasmidFinder 2.1

Database	Plasmid	Identity (%)	Query / Template length	Accession number
enterobacteriaceae	IncFIB(K)	99.82	560 / 560	JN233704
enterobacteriaceae	IncFII	99.62	261 / 261	AY458016
enterobacteriaceae	IncP6	99.88	806 / 806	JF785550
enterobacteriaceae	IncR	99.2	251 / 251	DQ449578

VirulenceFinder 2.0

Database	Virulence factor	Identity	Query / Template length	Protein function	Accession number
virulence_ecoli	gad	100	1401 /	Glutamate	AP009240
			1401	decarboxylase	
virulence_ecoli	gad	99.93	1401 /	Glutamate	AP010953
			1401	decarboxylase	
virulence_ecoli	<i>lpfA</i>	99.83	573 / 573	Long polar fimbriae	AP010953
virulence_ecoli	<i>lpfA</i>	96.68	573 / 573	Long polar fimbriae	CP002185
virulence_ecoli	terC	95.24	686 / 714	Tellurium ion	CP007491
				resistance protein	
virulence_ecoli	terC	98.96	966 / 966	Tellurium ion	MG591698
				resistance protein	
virulence_ecoli	terC	100	1041 /	Tellurium ion	UGAE01000003
			1041	resistance protein	

MLST 2.0 (Multi-Locus Sequence Typing)

Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
adk	100	100	536	536	0	adk_64	2795
fumC	100	100	469	469	0	fumC_4	
gyrB	100	100	460	460	0	gyrB_5	
icd	100	100	518	518	0	icd_1	
mdh	100	100	452	452	0	mdh_8	
purA	100	100	478	478	0	purA_8	
recA	100	100	510	510	0	recA_6	

SerotypeFinder 2.0

Database	Gene	Serotype	Identity	Template / HSP length	Accession number
H_type	fliC	H27	91.17	1337 / 1461	AM231154
O_type	wzy	O21	98.8	1083 / 1083	EU694098
O_type	WZX	O21	100	1236 / 1236	EU694098

Plasmids detected by ARESdb cloud platform

Plasmid	Identity	Query / Template length	Accession number
Col(pHAD28)	92.37%	131 / 131	KU674895
Col440I	90.99%	111 / 114	CP023920
IncFIB(K)	99.82%	560 / 560	JN233704
IncFII	99.62%	261 / 261	AY458016
IncFII(Yp)	92.64%	231 / 230	CP000670
IncP6	99.88%	806 / 806	JF785550
IncR	99.20%	251 / 251	DQ449578

Virulence Factors detected by ARESdb cloud platform

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
virulence factors	ydeH	-	442 aa	98.6%	100%
virulence factors	ydaM	-	410 aa	98%	100%
virulence factors	sfaH	-	304 aa	97.4%	100%
virulence factors	potD	-	348 aa	91.1%	100%
virulence factors	pemK	-	110 aa	100%	100%
virulence factors	pemI	-	85 aa	100%	100%
virulence factors	nagA	-	139 aa	98.6%	94.6%
virulence factors	mqsA	-	131 aa	100%	100%
virulence factors	mqo	-	548 aa	99.3%	100%
virulence factors	manZ	-	269 aa	98.5%	100%
virulence factors	manX	-	144 aa	97.9%	100%
virulence factors	ldrD	-	91 aa	94.5%	100%
virulence factors	fliZ	-	108 aa	98.1%	99.1%
virulence factors	fliZ	-	62 aa	96.8%	100%
virulence factors	fliY	-	266 aa	92.1%	100%
virulence factors	fliQ	-	89 aa	93.3%	100%
virulence factors	fliI	-	456 aa	95.6%	100%
virulence factors	fliG	-	332 aa	96.7%	100%
virulence factors	fliA	-	239 aa	99.6%	100%
virulence factors	flhA	-	692 aa	94.4%	100%
virulence factors	fimD	-	517 aa	98.6%	100%
virulence factors	EspX1	-	473 aa	91.5%	100%
virulence factors	epsJ	-	300 aa	98%	100%
virulence factors	$\it elfG$	-	357 aa	100%	100%
virulence factors	cstA	-	701 aa	92.4%	100%
fimbria	LpfA	-	190 aa	100%	100%

Antibiotic resistance genes detected by ARESdb cloud platform

Marker Class1	Marker2	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
Vga	Vga(C)	-	74 aa	94.6%	97.4%
undecaprenyl- diphosphatase	BacA	-	273 aa	100%	100%
target protection	Rpsl	-	124 aa	93.5%	100%
Sul	Sul1	-	308 aa	99.4%	100%
RND efflux	MdtF	-	1037 aa	99.7%	100%
transporter RND efflux	MdtE	-	385 aa	99.7%	100%
transporter RND efflux	MdtC	-	1024 aa	99.7%	99.9%
transporter RND efflux	MdtB	-	1040 aa	99.6%	100%
transporter RND efflux	MdtA	-	415 aa	99.3%	100%
transporter RND efflux	AcrF	-	1034 aa	99.3%	100%
transporter RND efflux	AcrE	-	385 aa	100%	100%
transporter RND efflux	AcrD	-	1037 aa	99.9%	100%
transporter RND efflux	AcrB	-	1049 aa	100%	100%
transporter RND efflux	AcrA	-	397 aa	100%	100%
transporter regulation of	virB	-	324 aa	91%	100%
resistance regulation of	rpoS	-	172 aa	100%	100%
resistance regulation of	ompR	-	239 aa	100%	100%
resistance Qnr	QnrB6	-	226 aa	98.2%	100%
putative efflux transport	yheI	-	590 aa	98.1%	100%
putative efflux transport	gspA	-	270 aa	98.1%	100%
putative efflux	cusR	-	227 aa	95.2%	100%
transport putative efflux	cusF	-	110 aa	97.3%	100%
transport porin	OmpC	-	368 aa	96.7%	100%
Pmr	PmrL	-	660 aa	99.8%	100%

Pmr	PmrF	-	322 aa	100%	100%
Pmr	PmrE	-	388 aa	98.7%	100%
Pmr	PmrC	-	547 aa	99.6%	100%
multidrug efflux SMR transporter	EmrE	-	110 aa	98.2%	100%
MFS efflux transporter	RosB	-	558 aa	91.9%	100%
MFS efflux transporter	MdtP	-	488 aa	97.7%	100%
MFS efflux transporter	MdtO	-	683 aa	99.6%	100%
MFS efflux transporter	MdtN	-	343 aa	100%	100%
MFS efflux	MdtM	-	410 aa	98%	100%
transporter MFS efflux	MdtH	-	402 aa	100%	100%
transporter MFS efflux	MdtG	-	408 aa	99.8%	100%
transporter MFS efflux	MdfA	-	410 aa	99.8%	100%
transporter MFS efflux	EmrY	-	512 aa	99.6%	100%
transporter MFS efflux	EmrK	-	372 aa	98.1%	96.1%
transporter MFS efflux	EmrD	-	394 aa	99.7%	100%
transporter MFS efflux	EmrB	-	512 aa	100%	100%
transporter MFS efflux	EmrA	-	390 aa	99.7%	100%
transporter macrolide	Mph(B)	-	157 aa	98.1%	99.4%
phosphotransferase influx	GlpT	G206A	423 aa	93.1%	93.8%
influx	GlpT	E448K	452 aa	98.2%	100%
influx	cycA	S263A	467 aa	98.3%	100%
fimbria	LpfA	-	190 aa	100%	100%
efflux regulation	SoxR	R4K	152 aa	93.4%	100%
efflux regulation	PhoQ	-	486 aa	99.4%	100%
efflux regulation	MarR	G103S, Y137H	144 aa	97.9%	100%
efflux regulation	MarA	-	129 aa	99.2%	100%
efflux regulation	<i>KdpE</i>	-	225 aa	99.1%	100%
efflux regulation	H-NS	-	137 aa	100%	100%

efflux regulation	GadX	-	274 aa	98.5%	100%
efflux regulation	GadW	-	242 aa	93.4%	100%
efflux regulation	EvgS	-	1197 aa	99.3%	100%
efflux regulation	EvgA	-	204 aa	100%	100%
efflux regulation	CRP	-	210 aa	99.5%	100%
efflux regulation	CpxA	-	457 aa	99.1%	100%
efflux regulation	AcrS	-	220 aa	99.5%	100%
efflux	TolC	-	495 aa	100%	100%
DNA topoisomerase	parC	T57S	752 aa	95.1%	100%
subunit DNA topoisomerase subunit	gyrA	S83L	875 aa	99.9%	100%
Dfr	DfrA27	-	157 aa	99.4%	100%
compensation	MgrB	-	46 aa	100%	97.9%
mechanism beta-lactamase TEM	TEM- 160	-	190 aa	98.9%	66.4%
beta-lactamase	argR	-	156 aa	94.2%	100%
regulation beta-lactamase regulation	ampR	-	105 aa	94.3%	85.4%
beta-lactamase KPC	KPC-2	-	293 aa	100%	100%
beta-lactamase ESC	ESC-148	-	420 aa	98.3%	100%
beta-lactamase	атрН	-	385 aa	99.7%	100%
ampC beta-lactamase ampC	ampC1	-	434 aa	99.8%	100%
BaeSR	BaeS	-	467 aa	100%	100%
BaeSR	BaeR	-	240 aa	100%	100%
Arr	Arr-3	-	162 aa	99.4%	82.2%
ANT(3")-Ia	aadA16	-	281 aa	100%	100%
ABC efflux pump	YojI	-	547 aa	100%	100%
ABC efflux pump	PatA	-	459 aa	99.8%	100%
ABC efflux pump	MsrB	-	137 aa	100%	100%
ABC efflux pump	MsbA	-	582 aa	100%	100%
ABC efflux pump	mdlB	-	593 aa	99.5%	100%
ABC efflux pump	MacA	-	371 aa	99.7%	100%
AAC(6')-I	AAC(6') -Ib-cr	-	199 aa	100%	100%

M20 E. coli (Table S4)

ResFinder 4.1								
Resistance gene	Identity (%)	Alignment Length/Gene Length	Coverage (%)	Position in reference	Phenotype	Accession no.		
dfrA14	100	474/474	100	1475	Trimethoprim resistance	KF921535		
sul2	100	816/816	100	1817	Sulphonamide resistance	AY034138		
blaOXA-10	100	801/801	100	1802	Beta-lactam resistance Alternate name; PSE-2	J03427		
blaGES-1	100	864/864	100	1865	Beta-lactam resistance	HQ170511		
blaKPC-2	100	882/882	100	1883	Beta-lactam resistance	AY034847		
blaOXA-48	100	798/798	100	1799	Beta-lactam resistance	AY236073		
aph(3")-Ib	99.88	804/804	100	1805	Aminoglycoside resistance	AF321551		
mdf(A)	99.92	1233/1233	100	11234	Macrolide resistance	Y08743		
qnrS1	100	657/657	100	1658	Quinolone resistance	AB187515		

PlasmidFinder 2.1

Database	Plasmid	Identity	Query / Template length	Accession number
enterobacteriaceae	Col(IRGK)	98.37	185 / 184	AY543071
enterobacteriaceae	IncFIB(K)	99.82	560 / 560	JN233704
enterobacteriaceae	IncFII	99.62	261 / 261	AY458016
enterobacteriaceae	IncL	100	661 / 661	JN626286
enterobacteriaceae	IncN	99.81	514 / 514	AY046276
enterobacteriaceae	IncP6	99.88	806 / 806	JF785550
enterobacteriaceae	IncY	98.95	765 / 765	K02380

Database	Virulence factor	Identity	Query / Template length	Protein function	Accession number
virulence_ecoli	gad	100	1401 / 1401	Glutamate decarboxylase	AP009240
virulence_ecoli	gad	99.93	1401 / 1401	Glutamate decarboxylase	AP010953
virulence_ecoli	lpfA	100	573 / 573	Long polar fimbriae	AP010953
virulence_ecoli	lpfA	98.78	573 / 573	Long polar fimbriae	CP002185
virulence_ecoli	terC	99.02	714 / 714	Tellurium ion resistance protein	CP007491
virulence_ecoli	terC	98.96	966 / 966	Tellurium ion resistance protein	MG591698
virulence_ecoli	terC	100	1041 / 1041	Tellurium ion resistance protein	UGAE01000003

MLST 2.0 (Multi-Locus Sequence Typing)

Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
adk	100	100	536	536	0	adk_64	2795
fumC	100	100	469	469	0	fumC_4	
gyr B	100	100	460	460	0	gyrB_5	
icd	100	100	518	518	0	icd_1	
mdh	100	100	452	452	0	mdh_8	
purA	100	100	478	478	0	purA_8	
recA	100	100	510	510	0	recA_6	

Plasmids detected by ARESdb cloud platform

Plasmid	Identity	Query / Template length	Accession number
Col(IRGK)	98.37%	184 / 184	AY543071
Col440I	95.61%	114 / 114	CP023920
Col440II	91.10%	281 / 282	CP023921
IncFIB(K)	99.82%	560 / 560	JN233704
IncFII	99.62%	261 / 261	AY458016
IncFII(Yp)	92.64%	231 / 230	CP000670
IncL	100.00%	661 / 661	JN626286
IncN	99.81%	514 / 514	AY046276
IncP6	99.88%	806 / 806	JF785550
IncY	98.95%	765 / 765	K02380

Virulence Factors detected by ARESdb cloud platform

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
virulence factors	ydeH	-	442 aa	98.6%	100%
virulence factors	ydaM	-	410 aa	98%	100%
virulence factors	sfaH	-	304 aa	97.4%	100%
virulence factors	potD	-	348 aa	91.1%	100%
virulence factors	pemK	-	110 aa	100%	100%
virulence factors	pemI	-	85 aa	100%	100%
virulence factors	nagA	-	139 aa	98.6%	94.6%
virulence factors	mqsA	-	131 aa	100%	100%
virulence factors	mqo	-	548 aa	99.3%	100%
virulence factors	manZ	-	269 aa	98.5%	100%
virulence factors	manX	-	144 aa	97.9%	100%
virulence factors	ldrD	-	91 aa	94.5%	100%
virulence factors	fliZ	-	108 aa	98.1%	99.1%
virulence factors	fliZ	-	62 aa	96.8%	100%
virulence factors	fliY	-	266 aa	92.1%	100%
virulence factors	fliQ	-	89 aa	93.3%	100%
virulence factors	fliI	-	456 aa	95.6%	100%
virulence factors	fliG	-	332 aa	96.7%	100%
virulence factors	fliA	-	239 aa	99.6%	100%
virulence factors	flhA	-	692 aa	94.4%	100%
virulence factors	fimD	-	517 aa	98.6%	100%
virulence factors	EspX1	-	473 aa	91.5%	100%
virulence factors	epsJ	-	300 aa	98%	100%
virulence factors	$\it elfG$	-	357 aa	100%	100%
virulence factors	cstA	-	701 aa	92.4%	100%
fimbria	<i>LpfA</i>	-	190 aa	100%	100%

Antibiotic resistance genes detected by ARESdb cloud platform

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
Vga	Vga(C)	-	74 aa	94.6%	97.4%
undecaprenyl- diphosphatase	BacA	-	273 aa	100%	100%
target protection	Rpsl	-	124 aa	93.5%	100%
Sul	Sul2	-	282 aa	98.9%	98.3%
RND efflux	MdtF	-	1037 aa	99.7%	100%
transporter RND efflux	MdtE	_	385 aa	99.7%	100%
transporter					
RND efflux	MdtC	-	1024 aa	99.7%	99.9%
transporter RND efflux	MdtB	-	1040 aa	99.6%	100%
transporter RND efflux	MdtA	-	415 aa	99.3%	100%
transporter RND efflux	AcrF	-	1034 aa	99.3%	100%
transporter RND efflux	AcrE	-	385 aa	100%	100%
transporter RND efflux	AcrD	-	1037 aa	99.9%	100%
transporter RND efflux	AcrB	_	1049 aa	100%	100%
transporter RND efflux	AcrA	-	397 aa	100%	100%
transporter regulation of	rpoS	_	172 aa	100%	100%
resistance	rpos		172 dd	10070	10070
regulation of resistance	ompR	-	239 aa	100%	100%
Qnr	QnrS1	-	218 aa	100%	100%
putative efflux transport	yheI	-	590 aa	98.1%	100%
putative efflux	gspA	-	270 aa	98.1%	100%
transport putative efflux	cusR	-	227 aa	95.2%	100%
transport putative efflux	cusF	-	110 aa	97.3%	100%
transport porin	OmpC	_	368 aa	96.5%	100%
Pmr	PmrL	_	660 aa	99.8%	100%
Pmr	PmrF	_	322 aa	100%	100%
		-			
Pmr	PmrE	-	388 aa	99%	100%

multidrug efflux SMR transporter	EmrE	-	110 aa	98.2%	100%
MFS efflux	RosB	_	558 aa	91.9%	100%
transporter					
MFS efflux	MdtM	-	410 aa	98%	100%
transporter MFS efflux	M 4411		402 aa	100%	100%
transporter	MdtH	-	402 aa	100%	100%
MFS efflux	MdtG	-	408 aa	99.8%	100%
transporter					
MFS efflux	MdfA	-	410 aa	99.8%	100%
transporter MFS efflux	EmrY	_	512 aa	99.6%	100%
transporter	Liiii I	_	312 dd	<i>77.</i> 070	10070
MFS efflux	EmrK	-	372 aa	98.1%	96.1%
transporter					
MFS efflux	EmrD	-	394 aa	99.7%	100%
transporter MFS efflux	EmrB	_	512 aa	100%	100%
transporter	Lini		312 dd	10070	10070
MFS efflux	EmrA	-	390 aa	99.7%	100%
transporter	16.1(D)		1.57	00.10/	00.40/
macrolide phosphotransferase	Mph(B)	-	157 aa	98.1%	99.4%
influx	GlpT	G206A	423 aa	93.1%	93.8%
influx	GlpT	E448K	452 aa	98.2%	100%
influx	cycA	S263A	467 aa	98.3%	100%
fimbria	LpfA	-	190 aa	100%	100%
efflux regulation	SoxR	R4K	152 aa	93.4%	100%
efflux regulation	PhoQ	_	486 aa	99.4%	100%
efflux regulation	MarR	G103S,	144 aa	97.9%	100%
citiax regulation	Man	Y137H	11144	J1.J/0	10070
efflux regulation	MarA	-	129 aa	99.2%	100%
efflux regulation	KdpE	-	225 aa	99.1%	100%
efflux regulation	H-NS	-	137 aa	100%	100%
efflux regulation	GadX	-	274 aa	98.5%	100%
efflux regulation	GadW	-	242 aa	93.4%	100%
efflux regulation	EvgS	_	1197 aa	99.3%	100%
efflux regulation	EvgA	_	204 aa	100%	100%
_					
efflux regulation	CRP	-	210 aa	99.5%	100%
efflux regulation	CpxA	-	457 aa	99.1%	100%
efflux regulation	AcrS	-	220 aa	99.5%	100%
efflux	TolC	-	495 aa	100%	100%

DNA	parC	T57S	752 aa	95.1%	100%
topoisomerase subunit					
DNA	gyrA	S83L	875 aa	99.9%	100%
topoisomerase	8,771	50312	075 dd	<i>JJ</i> . <i>J</i> /0	10070
subunit					
Dfr	DfrA14	-	160 aa	99.4%	100%
compensation	MgrB	-	46 aa	100%	97.9%
mechanism					
beta-lactamase	<i>TEM-</i>	-	190 aa	98.9%	66.4%
TEM beta-lactamase	160 argR	_	156 aa	94.2%	100%
regulation	ui gre		130 aa	J 1.270	10070
beta-lactamase	ampR	-	105 aa	94.3%	85.4%
regulation					
beta-lactamase	<i>OXA-48</i>	-	265 aa	100%	100%
OXA beta-lactamase	OXA-10	_	266 aa	100%	100%
OXA	OM1 10		200 44	10070	10070
beta-lactamase	KPC-2	-	293 aa	100%	100%
KPC				100.	
beta-lactamase	GES-1	-	287 aa	100%	100%
GES beta-lactamase ESC	ESC-148	_	420 aa	98.3%	100%
beta-lactamase ampC	атрН	-	385 aa	99.7%	100%
beta-lactamase	ampC1	_	434 aa	99.8%	100%
ampC	1				
BaeSR	BaeS	-	467 aa	100%	100%
BaeSR	BaeR	-	240 aa	100%	100%
APH(3")	<i>APH</i> (3")	-	277 aa	99.6%	91.7%
	-Ib				
ABC efflux pump	YojI	-	547 aa	100%	100%
ABC efflux pump	PatA	-	459 aa	99.8%	100%
ABC efflux pump	MsrB	-	137 aa	100%	100%
ABC efflux pump	MsbA	-	582 aa	100%	100%
ABC efflux pump	mdlB	-	593 aa	99.5%	100%
ABC efflux pump	MacA	_	371 aa	99.7%	100%
AAC(6')-I	AAC(6')		192 aa	100%	94.6%
AAC(0)-1	-Ib	-	1 <i>94</i> aa	100 70	J +. U%
	10				

5a K. pneumoniae

(Table S5)

ResFinder 4.1

Resistance gene	Identity	Alignment Length/Gene Length	Coverage	Position in reference	Phenotype	Accession no.
sul1	100	840/840	100	1841	Sulphonamide resistance	U12338
tet(A)	100	1200/1200	100	11201	Tetracycline resistance	AJ517790
OqxA	100	1176/1176	100	11177	Disinfectant resistance	EU370913
OqxB	99.97	3153/3153	100	13154	Disinfectant resistance	EU370913
oqxB	99.97	3153/3153	100	13154	Quinolone resistance	EU370913
aac(6')-Ib-cr	94.61	519/519	95.95	1520	Fluoroquinolone and aminoglycoside resistance	EF636461
oqxA	100	1176/1176	100	11177	Quinolone resistance	EU370913
blaCTX-M-3	100	876/876	100	1877	Beta-lactam resistance	Y10278
blaOXA-2	100	828/828	100	1829	Beta-lactam resistance	DQ112222
blaGES-5	100	864/864	100	1865	Beta-lactam resistance	DQ236171
aph(3")-Ib	100	804/804	100	1805	Aminoglycoside resistance	AF024602
aph(6)- Id	100	837/837	100	1838	Aminoglycoside resistance	M28829
fosA	99.29	420/420	100	1421	Fosfomycin resistance	ACWO01000079
blaKPC-2	100	882/882	100	1883	Beta-lactam resistance	AY034847

PlasmidFinder 2.1

Database	Plasmid	Identity	Query / Template length	Accession number
enterobacteriaceae	Col(pHAD28)	95.42	131 / 131	KU674895
enterobacteriaceae	Col440II	97.16	282 / 282	CP023921
enterobacteriaceae	IncFIA(HI1)	96.91	388 / 388	AF250878
enterobacteriaceae	IncFII(K)	97.97	148 / 148	CP000648
enterobacteriaceae	IncX5	100	576 / 576	MF062700
enterobacteriaceae	IncY	99.48	765 / 765	K02380

MLST 2.0 (Multi-Locus Sequence Typing)

Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
gapA	100	100	450	450	0	gapA_2	37
infB	100	100	318	318	0	infB_9	
mdh	100	100	477	477	0	mdh_2	
pgi	100	100	432	432	0	pgi_1	
phoE	100	100	420	420	0	phoE_13	
rpoB	100	100	501	501	0	rpoB_1	
tonB	100	100	414	414	0	tonB_16	

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
Vga	Vga(C)	-	74 aa	94.6%	97.4%
target protection	Rpsl	-	124 aa	94.4%	100%
Sul	Sul 1	-	308 aa	99.4%	100%
RND efflux transporter	SilC	-	461 aa	100%	100%
RND efflux transporter	SilB	-	430 aa	97.9%	100%
RND efflux transporter	SilA	-	1048 aa	99%	100%
RND efflux transporter	OqxB	-	1050 aa	99.9%	100%
RND efflux transporter	OqxA	-	391 aa	100%	100%
RND efflux transporter	MdtC	-	1024 aa	91.6%	99.9%
RND efflux transporter	MdtB	-	1040 aa	90.1%	100%
RND efflux transporter	AcrD	-	1037 aa	91%	100%
RND efflux transporter	AcrB	-	1049 aa	91.5%	100%
RND efflux transporter	AcrA	-	397 aa	100%	100%
regulation of resistance	virB	-	324 aa	91.7%	100%
regulation of resistance	rpoS	-	172 aa	99.4%	100%
regulation of resistance	rnfC	-	562 aa	91.6%	75.9%
regulation of resistance	ompR	-	239 aa	100%	100%
regulation of resistance	mtfA	-	265 aa	100%	100%
regulation of resistance	MerR	-	121 aa	91.7%	100%
regulation of resistance	MerR	-	144 aa	100%	100%
regulation of resistance	MerR	-	116 aa	93.1%	80.6%
regulation of resistance	leuO	-	308 aa	99.4%	100%
Qnr	QnrB12	-	215 aa	92.1%	100%
putative efflux transport	cusR	-	227 aa	99.1%	100%
porin	OmpK37	D275T, E244D, I128M, I70M, M233Q, N230G, N274S, R239K, V277I	384 aa	94.3%	100%
porin	OmpK36	-	369 aa	99.2%	100%
porin	OmpK36	A217S, D223G, E232R, F198Y, F207Y, G189T, L229A, L59V, N304E, T222L	369 aa	93.5%	100%
porin	OmpK35	-	195 aa	100%	85.9%
Pmr	PmrE	-	388 aa	99.5%	100%

phospholipid biosynthesis pathway	tesA	-	184 aa	92.4%	100%	
MFS efflux transporter	Tet(A)	_	424 aa	100%	100%	
MFS efflux transporter	RosB	_	558 aa	100%	100%	
MFS efflux transporter	RosA	-	406 aa	99.3%	100%	
MFS efflux transporter	KpnH	-	512 aa	93.4%	100%	
MFS efflux transporter	KpnG	_	390 aa	99.7%	100%	
MFS efflux transporter	KpnF	_	109 aa	100%	100%	
MFS efflux transporter	KpnE	-	120 aa	99.2%	100%	
MFS efflux transporter	KdeA	-	410 aa	100%	100%	
MFS efflux transporter	EmrD	-	394 aa	99.5%	100%	
influx regulation	bamB	A176T	360 aa	94.4%	100%	
influx	UhpT	E350Q	456 aa	96.1%	98.5%	
influx	GlpT	G206A	423 aa	93.4%	93.8%	
influx	cycA	S263A	467 aa	92.7%	100%	
glutathione transferase	FosA	-	139 aa	99.3%	100%	
efflux regulation	MarA	-	123 aa	93.5%	96.9%	
efflux regulation	H-NS	-	135 aa	93.3%	98.5%	
efflux regulation	CRP	-	210 aa	99%	100%	
efflux regulation	CpxA	-	461 aa	98.7%	100%	
DNA topoisomerase subunit	parC	T57S	752 aa	94.3%	100%	
DNA topoisomerase subunit	gyrA	S83Y	877 aa	99.8%	100%	
beta-lactamase regulation	argR	-	156 aa	100%	100%	
beta-lactamase OXA	OXA-2	-	275 aa	100%	100%	
beta-lactamase GES	GES-5	-	287 aa	100%	100%	
beta-lactamase CTX-M	CTX-M-3	-	291 aa	100%	100%	
BasRS	BasS synonym PmrB	R256G	365 aa	99.5%	100%	
BaeSR	BaeS	-	492 aa	99.4%	100%	
BaeSR	BaeR	-	240 aa	99.6%	100%	
APH(6)	APH(6)-Id	-	278 aa	100%	100%	
APH(3")	<i>APH(3'')-Ib</i>	-	290 aa	99.7%	96%	
ABC efflux pump	PatA	-	459 aa	94.6%	100%	
ABC efflux pump	MsbA	-	582 aa	91.9%	100%	
AAC(6')-I	AAC(6')- Ib	-	179 aa	100%	88.2%	

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
virulence factors	sinR	-	180 aa	90%	99.4%
virulence factors	sfaG	-	182 aa	100%	100%
virulence factors	potD	-	348 aa	94.8%	100%
virulence factors	gmr	-	551 aa	100%	100%
virulence factors	fliY	-	310 aa	100%	100%
virulence factors	fliY	-	266 aa	92.1%	100%
virulence factors	fimD	-	859 aa	99.5%	100%
virulence factors	feoB	T755A	772 aa	94.2%	100%
virulence factors	eutB	-	453 aa	94.9%	100%
virulence factors	cnrA	-	1030 aa	99.8%	100%

Plasmids detected by ARESdb cloud platform

Plasmid	Identity	Query / Template length	Accession number
Col(pHAD28)	93.89%	131 / 131	KU674895
Col440I	95.61%	114 / 114	CP023920
Col440II	97.16%	282 / 282	CP023921
IncFIA(HI1)	96.91%	388 / 388	AF250878
IncFII(K)	97.97%	148 / 148	CP000648
IncFII(Yp)	92.21%	231 / 230	CP000670
IncX5	100.00%	576 / 576	MF062700
IncY	99.48%	765 / 765	K02380

M11 K. pneumoniae (Table S6)

ResFinder 4.1

Resistance gene	Identity (%)	Alignment Length/Gene Length	Coverage (%)	Position in reference	Phenotype	Accession no.
fosA	99.29	420/420	100	1421	Fosfomycin resistance	ACWO01000079
·					•	
OqxA	100	1176/1176	100	11177	Disinfectant resistance	EU370913
OqxB	99.97	3153/3153	100	13154	Disinfectant resistance	EU370913
blaGES-5	100	864/864	100	1865	Beta-lactam resistance	DQ236171
blaKPC-2	100	882/882	100	1883	Beta-lactam resistance	AY034847
blaOXA-2	100	828/828	100	1829	Beta-lactam resistance	DQ112222
blaCTX-M-3	100	876/876	100	1877	Beta-lactam resistance	Y10278
aph(3")-Ib	99.88	803/804	99.88	2805	Aminoglycoside resistance	AF024602
aph(6)- Id	100	837/837	100	1838	Aminoglycoside resistance	M28829
oqxA	100	1176/1176	100	11177	Quinolone resistance	EU370913
oqxB	99.97	3153/3153	100	13154	Quinolone resistance	EU370913
tet(A)	97.57	1244/1275	97.57	11245	Tetracycline resistance	AF534183
sul1	100	840/840	100	1841	Sulphonamide resistance	U12338

PlasmidFinder 2.1

Database	Plasmid	Identity (%)	Query / Template length	Accession number
enterobacteriaceae	Col(pHAD28)	95.42	131 / 131	KU674895
enterobacteriaceae	Col440II	97.16	282 / 282	CP023921
enterobacteriaceae	IncFIA(HI1)	96.91	388 / 388	AF250878
enterobacteriaceae	IncFII(K)	97.97	148 / 148	CP000648
enterobacteriaceae	IncX5	100	576 / 576	MF062700
enterobacteriaceae	IncY	99.48	765 / 765	K02380

MLST 2.0 (Multi-Locus Sequence Typing)

Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
gapA	100	100	450	450	0	gapA_2	37
infB	100	100	318	318	0	infB_9	
mdh	100	100	477	477	0	mdh_2	
pgi	100	100	432	432	0	pgi_1	
phoE	100	100	420	420	0	phoE_13	
rpoB	100	100	501	501	0	rpoB_1	
tonB	100	100	414	414	0	tonB_16	

Marker Class1	Marker2	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
virulence factors	sinR	-	180 aa	90%	99.4%
virulence factors	sfaG	-	182 aa	100%	100%
virulence factors	potD	-	348 aa	94.8%	100%
virulence factors	gmr	-	551 aa	100%	100%
virulence factors	fliY	-	310 aa	100%	100%
virulence factors	fliY	-	266 aa	92.1%	100%
virulence factors	fimD	-	859 aa	99.5%	100%
virulence factors	feoB	T755A	772 aa	94.2%	100%
virulence factors	eutB	-	453 aa	94.9%	100%
virulence factors	cnrA	-	1030 aa	99.8%	100%

Plasmids detected by ARESdb cloud platform

Plasmid	Identity	Query / Template length	Accession number
Col(pHAD28)	94.66%	131 / 131	KU674895
Col440I	95.61%	114 / 114	CP023920
Col440II	97.16%	282 / 282	CP023921
IncFIA(HI1)	96.91%	388 / 388	AF250878
IncFII(K)	97.97%	148 / 148	CP000648
IncFII(Yp)	92.21%	231 / 230	CP000670
IncX5	100.00%	576 / 576	MF062700
IncY	99.48%	765 / 765	K02380

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
Vga	Vga(C)	-	74 aa	94.6%	97.4%
target protection	Rpsl	-	124 aa	94.4%	100%
Sul	Sul1	-	308 aa	99.4%	100%
RND efflux transporter	SilC	-	461 aa	100%	100%
RND efflux transporter	SilB	-	430 aa	97.9%	100%
RND efflux transporter	SilA	-	1048 aa	99%	100%
RND efflux transporter	OqxB	-	1050 aa	99.9%	100%
RND efflux transporter	OqxA	-	391 aa	100%	100%
RND efflux transporter	MdtC	-	1024 aa	91.6%	99.9%
RND efflux transporter	MdtB	-	1040 aa	90.1%	100%
RND efflux transporter	AcrD	-	1037 aa	91%	100%
RND efflux transporter	AcrB	-	1049 aa	91.5%	100%
RND efflux transporter	AcrA	-	397 aa	100%	100%
regulation of resistance	virB	-	324 aa	91.7%	100%

regulation of resistance	rpoS	-	172 aa	99.4%	100%	
regulation of resistance	rnfC	-	562 aa	91.6%	75.9%	
regulation of resistance	ompR	-	239 aa	100%	100%	
regulation of resistance	mtfA	-	265 aa	100%	100%	
regulation of resistance	MerR	-	121 aa	91.7%	100%	
regulation of resistance	MerR	-	144 aa	100%	100%	
regulation of resistance	MerR	-	116 aa	93.1%	80.6%	
regulation of resistance	leuO	-	308 aa	99.4%	100%	
Qnr	QnrB12	-	215 aa	92.1%	100%	
putative efflux transport	cusR	-	227 aa	99.1%	100%	
porin	OmpK37	D275T, E244D, I128M, I70M, M233Q, N230G, N274S, R239K, V277I	384 aa	94.3%	100%	
porin	OmpK36	-	368 aa	99.2%	100%	
porin	OmpK36	A217S, D223G, E232R, F198Y, F207Y, G189T, L229A, L59V, N304E, T222L	368 aa	93.5%	100%	
porin	OmpK35	-	195 aa	100%	85.9%	
Pmr	PmrE	-	388 aa	99.5%	100%	
phospholipid biosynthesis pathway	tesA	-	184 aa	92.4%	100%	
MFS efflux transporter	Tet(A)	-	424 aa	100%	100%	
MFS efflux transporter	RosB	-	558 aa	100%	100%	
MFS efflux transporter	RosA	-	406 aa	99.3%	100%	
-						
MFS efflux transporter	КрпН	-	512 aa	93.4%	100%	
	KpnH KpnG	- -	512 aa 390 aa	93.4% 99.7%	100% 100%	

MFS efflux transporter	KpnE	-	120 aa	99.2%	100%
MFS efflux transporter	KdeA	-	410 aa	100%	100%
MFS efflux transporter	EmrD	-	394 aa	99.5%	100%
influx regulation	bamB	A176T	360 aa	94.4%	100%
influx	UhpT	E350Q	456 aa	96.1%	98.5%
influx	GlpT	G206A	423 aa	93.4%	93.8%
influx	cycA	S263A	467 aa	92.7%	100%
glutathione transferase	FosA	-	139 aa	99.3%	100%
efflux regulation	MarA	-	123 aa	93.5%	96.9%
efflux regulation	H-NS	-	135 aa	93.3%	98.5%
efflux regulation	CRP	-	210 aa	99%	100%
efflux regulation DNA	CpxA	-	461 aa	98.7%	100%
topoisomerase subunit DNA	parC	T57S	752 aa	94.3%	100%
topoisomerase subunit	gyrA	S83Y	877 aa	99.8%	100%
beta-lactamase regulation	argR	-	156 aa	100%	100%
beta-lactamase OXA	OXA-2	-	275 aa	100%	100%
beta-lactamase GES	GES-5	-	287 aa	100%	100%
beta-lactamase CTX-M	CTX-M-3	-	291 aa	100%	100%
BasRS	BasS synonym PmrB	R256G	365 aa	99.5%	100%
BaeSR	BaeS	-	492 aa	99.4%	100%
BaeSR	BaeR	-	240 aa	99.6%	100%
APH(6)	APH(6)-Id	_	278 aa	100%	100%
APH(3")	APH(3'')- Ib	-	290 aa	99.7%	96%
ABC efflux pump	PatA	-	459 aa	94.6%	100%
ABC efflux pump	MsbA	-	582 aa	91.9%	100%
AAC(6')-I	AAC(6')- Ib	-	179 aa	100%	88.2%

C2 K. pneumoniae (Table S7)

ResFinder 4.1

Resistance gene	Identity (%)	Alignment Length/Gene Length	Coverage	Position in reference	Phenotype	Accession no.
oqxB	98.92	3153/3153	100	13154	Quinolone resistance	EU370913
oqxA	99.32	1176/1176	100	11177	Quinolone resistance	EU370913
aac(6')-Ib-cr	98	600/600	100	1601	Fluoroquinolone and aminoglycoside resistance	DQ303918
sul1	100	840/840	100	1841	Sulphonamide resistance	U12338
blaKPC-2	100	882/882	100	1883	Beta-lactam resistance	AY034847
blaGES-5	100	864/864	100	1865	Beta-lactam resistance	DQ236171
blaSHV-11	92.8	861/861	92.8	1862	Beta-lactam resistance	AF535128
blaOXA-48	100	798/798	100	1799	Beta-lactam resistance	AY236073
fosA	99.05	420/420	100	1421	Fosfomycin resistance	ACWO01000079
OqxB	98.92	3153/3153	100	13154	Disinfectant resistance	EU370913
OqxA	99.32	1176/1176	100	11177	Disinfectant resistance	EU370913

PlasmidFinder 2.1

Database	Plasmid	Identity (%)	Query / Template length	Accession number
enterobacteriaceae	Col(pHAD28)	100	131 / 131	KU674895
enterobacteriaceae	IncFIB(K)	98.93	560 / 560	JN233704
enterobacteriaceae	IncFII(pKP91)	95.22	230 / 230	CP000966
enterobacteriaceae	IncL	100	661 / 661	JN626286
enterobacteriaceae	IncR	99.6	251 / 251	DQ449578

MLST 2.0 (Multi-Locus Sequence Typing)

Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
gapA	100	100	450	450	0	gapA_2	534
infB	100	100	318	318	0	infB_1	
mdh	100	100	477	477	0	mdh_1	
pgi	100	100	432	432	0	pgi_2	
phoE	100	100	420	420	0	phoE_61	
rpoB	100	100	501	501	0	rpoB_4	
tonB	100	100	408	408	0	tonB_132	

Marker Class	Marker	Variant(s)	Alignment	Sequence	Sequence
			Length	Identity	Coverage
virulence factors	sinR	-	180 aa	90%	99.4%
virulence factors	sfaG	-	183 aa	97.8%	100%
virulence factors	potD	-	348 aa	95.1%	100%
virulence factors	pemK	-	110 aa	96.4%	100%
virulence factors	pemI	-	85 aa	96.5%	100%
virulence factors	gmr	-	551 aa	100%	100%
virulence factors	fliY	-	310 aa	99.7%	100%
virulence factors	fimD	-	859 aa	98.7%	100%
virulence factors	feoB	T755A	772 aa	94.2%	100%
virulence factors	eutB	-	453 aa	94.9%	100%
virulence factors	cstA	-	701 aa	92.6%	100%
virulence factors	cnrA	-	1030 aa	99.2%	100%

Marker Class	Marker	Varian t(s)	Alignment Length	Sequence Identity	Sequence Coverage
Vga	Vga(C)	-	76 aa	100%	100%
target protection	Rpsl	-	124 aa	94.4%	100%
Sul	Sul1	-	308 aa	99.4%	100%
RND efflux transporter	SilC	-	461 aa	100%	100%
RND efflux transporter	SilB	-	430 aa	97.9%	100%
RND efflux transporter	SilA	-	1048 aa	98.9%	100%
RND efflux transporter	MdtC	-	1024 aa	91.2%	99.9%
RND efflux transporter	MdtB	-	1040 aa	90.7%	100%
RND efflux transporter	AcrD	-	1037 aa	91%	100%
RND efflux transporter	AcrB	-	1049 aa	91.4%	100%
RND efflux transporter	AcrA	-	397 aa	99.7%	100%
regulation of resistance	virB	-	324 aa	92.3%	100%
regulation of resistance	rpoS	-	172 aa	99.4%	100%
regulation of resistance	rnfC	-	562 aa	91.8%	75.9%
regulation of resistance	ompR	-	239 aa	100%	100%
regulation of resistance	mtfA	-	265 aa	100%	100%
regulation of resistance	leuO	-	308 aa	99.4%	100%
Qnr	QnrB12	-	215 aa	92.1%	100%
putative efflux transport	setB	A200E	393 aa	99.5%	100%
putative efflux transport	cusR	-	227 aa	98.7%	100%
porin	OmpK37	I128M, I70M	374 aa	99.5%	100%
porin	OmpK36	-	375 aa	95.5%	100%
porin	OmpK36	A217S, D223G, E232R, F198Y, F207Y, G189T, L229A, L59V, T222L	375 aa	90.9%	100%
porin	OmpK35	-	226 aa	100%	99.6%
Pmr	PmrE	-	388 aa	99%	100%

phospholipid biosynthesis pathway	tesA	-	184 aa	92.4%	100%
OqxB	OqxB19	-	1050 aa	100%	100%
OqxA	OqxA10	-	391 aa	100%	100%
MFS efflux transporter	RosB	-	558 aa	100%	100%
MFS efflux transporter	RosA	-	406 aa	100%	100%
MFS efflux transporter	KpnH	-	512 aa	93.4%	100%
MFS efflux transporter	KpnG	-	390 aa	99.5%	100%
MFS efflux transporter	KpnF	-	109 aa	100%	100%
MFS efflux transporter	KpnE	-	120 aa	100%	100%
MFS efflux transporter	KdeA	-	410 aa	99.8%	100%
MFS efflux transporter	EmrD	-	394 aa	99.5%	100%
influx regulation	bamB	A176T	360 aa	94.4%	100%
influx	UhpT	E350Q	456 aa	96.1%	98.5%
influx	GlpT	G206A	423 aa	93.4%	93.8%
influx	cycA	S263A	467 aa	92.5%	100%
glutathione transferase	FosA	-	139 aa	99.3%	100%
efflux regulation	RamA	-	113 aa	92%	91.1%
efflux regulation	MarA	-	123 aa	93.5%	96.9%
efflux regulation	H-NS	-	135 aa	93.3%	98.5%
efflux regulation	CRP	-	210 aa	99%	100%
efflux regulation	CpxA	-	461 aa	98.5%	100%
efflux regulation	AcrR	F172S, F197I, G164A, K201M , L195V, P161R, R173G	216 aa	96.8%	100%
DNA topoisomerase subunit	parC	T57S	752 aa	94.1%	100%
DNA topoisomerase subunit	gyrA	S83Y	877 aa	99.8%	100%
beta-lactamase SHV	SHV-11	-	286 aa	100%	100%
beta-lactamase regulation	argR	-	156 aa	100%	100%
beta-lactamase OXA	OXA-48	-	265 aa	100%	100%
beta-lactamase GES	GES-5	-	287 aa	100%	100%
BaeSR	BaeS	-	492 aa	99.4%	100%

BaeSR	BaeR	-	240 aa	99.2%	100%
ABC efflux pump	PatA	-	459 aa	94.6%	100%
ABC efflux pump	MsbA	-	582 aa	91.9%	100%
AAC(6')-I	<i>AAC</i> (6')- <i>Ib7</i>	-	210 aa	99%	100%

Plasmids detected by ARESdb cloud platform

Plasmid	Identity	Query / Template length	Accession number
Col(pHAD28)	100.00%	131 / 131	KU674895
Col440I	95.61%	114 / 114	CP023920
IncFIB(K)	98.93%	560 / 560	JN233704
IncFII(pKP91)	96.07%	229 / 230	CP000966
IncFII(pMET)	90.92%	584 / 577	EU383016
IncL	100.00%	661 / 661	JN626286
IncQ1	93.58%	779 / 796	M28829
IncR	99.60%	251 / 251	DQ449578

CF1 C. freundii (Table S8)

ResFinder 4.1							
Resistance	Identity	Alignment	Coverage	Position	Phenotype	Accession	
gene	(%)	Length/Gene		in		no.	
		Length		reference			
blaKPC-2	100	882/882	100	1883	Beta-lactam resistance	AY034847	

PlasmidFinder 2.1

Database	Plasmid	Identity	Query / Template length	Accession number
enterobacteriaceae	IncFIB(pHCM2)	96.69	875 / 875	AL513384

T.	ΛT	CT	2	Λ	$M_{\rm D}$	ılti_	Locus	Sequence	Tyning)	١
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Locus	Identity	Coverage	Alignment Length	Allele Length	Gaps	Allele	ST
arcA	100	100	435	435	0	arcA_14	128
aspC	100	100	513	513	0	aspC_23	
clpX	100	100	567	567	0	clpX_20	
dnaG	100	100	444	444	0	dnaG_58	
fadD	100	100	483	483	0	fadD_36	
lysP	100	100	477	477	0	lysP_62	
mdh	100	100	549	549	0	mdh_60	

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverag e
undecaprenyl- diphosphatase	BacA	-	273 aa	92.7%	100%
target protection	Rpsl	-	124 aa	93.5%	100%
RND efflux transporter	SilC	-	461 aa	97.6%	100%
RND efflux transporter	SilB	-	430 aa	96.5%	100%
RND efflux transporter	SilA	-	1048 aa	98.9%	100%
RND efflux transporter	MdtC	-	1025 aa	92.7%	100%
RND efflux transporter	MdtB	-	1040 aa	92.8%	100%
RND efflux transporter	AcrD	-	1037 aa	94.7%	100%
RND efflux transporter	AcrB	-	1049 aa	94.5%	100%
RND efflux transporter	AcrA	-	397 aa	91.7%	100%
regulation of resistance	rpoS	-	172 aa	100%	100%
regulation of resistance	rnfC	-	560 aa	91.1%	83.2%
regulation of resistance	ompR	-	239 aa	100%	100%
putative efflux transport	sugE	-	105 aa	99%	100%
putative efflux transport	cusR	-	226 aa	97.3%	99.6%
phospholipid biosynthesis pathway	tesA	-	183 aa	90.2%	99.5%
MFS efflux transporter	RosB	-	557 aa	92.5%	99.8%
MFS efflux transporter	MdtH	-	402 aa	91.3%	100%
MFS efflux transporter	MdtG	-	406 aa	91.4%	99.5%
MFS efflux transporter	MdfA	-	410 aa	91%	100%
MFS efflux transporter	EmrB	-	512 aa	95.3%	100%
influx	UhpT	E350Q	463 aa	95.7%	100%
influx	GlpT	G206A	423 aa	92.9%	93.8%

cycA	S263A	466 aa	96.1%	99.8%	
SoxR	R4K	152 aa	93.4%	100%	
SoxR	T38S	152 aa	94.1%	98.7%	
MarR	Y137H	144 aa	90.3%	100%	
MarA	-	128 aa	93%	99.2%	
KdpE	-	224 aa	90.6%	99.6%	
H-NS	-	137 aa	94.2%	100%	
CRP	-	210 aa	99%	100%	
CpxA	-	457 aa	98%	100%	
parC	T57S	752 aa	96%	100%	
gyrA	S83T	875 aa	92.5%	99.8%	
argR	-	156 aa	94.2%	100%	
cmy-157	-	381 aa	97.4%	100%	
ampH	-	385 aa	91.7%	100%	
BaeR	-	239 aa	96.2%	99.6%	
YojI	-	547 aa	91.2%	100%	
PatA	-	458 aa	95%	99.8%	
MsbA	-	582 aa	94.7%	100%	
mdlB	-	592 aa	91%	99.8%	
MacB	-	376 aa	94.9%	100%	
	SoxR SoxR MarR MarA KdpE H-NS CRP CpxA parC gyrA argR cmy-157 ampH BaeR YojI PatA MsbA mdlB	SoxR R4K SoxR T38S MarR Y137H MarA - KdpE - H-NS - CRP - CpxA - parC T57S gyrA S83T argR - cmy-157 - ampH - BaeR - YojI - PatA - MsbA - mdlB -	SoxR R4K 152 aa SoxR T38S 152 aa MarR Y137H 144 aa MarA - 128 aa KdpE - 224 aa H-NS - 137 aa CRP - 210 aa CpxA - 457 aa parC T57S 752 aa gyrA S83T 875 aa argR - 156 aa cmy-157 - 381 aa ampH - 385 aa BaeR - 239 aa YojI - 547 aa PatA - 458 aa MsbA - 582 aa mdlB - 592 aa	SoxR R4K 152 aa 93.4% SoxR T38S 152 aa 94.1% MarR Y137H 144 aa 90.3% MarA - 128 aa 93% KdpE - 224 aa 90.6% H-NS - 137 aa 94.2% CRP - 210 aa 99% CpxA - 457 aa 98% parC T57S 752 aa 96% gyrA S83T 875 aa 92.5% argR - 156 aa 94.2% cmy-157 - 381 aa 97.4% ampH - 385 aa 91.7% BaeR - 239 aa 96.2% YojI - 547 aa 91.2% PatA - 458 aa 95% MsbA - 582 aa 94.7% mdlB - 592 aa 91%	SoxR R4K 152 aa 93.4% 100% SoxR T38S 152 aa 94.1% 98.7% MarR Y137H 144 aa 90.3% 100% MarA - 128 aa 93% 99.2% KdpE - 224 aa 90.6% 99.6% H-NS - 137 aa 94.2% 100% CRP - 210 aa 99% 100% CpxA - 457 aa 98% 100% parC T57S 752 aa 96% 100% gyrA S83T 875 aa 92.5% 99.8% argR - 156 aa 94.2% 100% cmy-157 - 381 aa 97.4% 100% ampH - 385 aa 91.7% 100% BaeR - 239 aa 96.2% 99.6% YojI - 547 aa 91.2% 100% PatA - 458 aa 95%

Marker Class	Marker	Variant(s)	Alignment Length	Sequence Identity	Sequence Coverage
virulence factors	potD	-	348 aa	93.4%	100%
virulence factors	fliY	-	266 aa	94%	100%
virulence factors	fliQ	-	89 aa	91%	100%
virulence factors	fliP	-	245 aa	91.4%	100%
virulence factors	fliI	-	456 aa	95.8%	100%
virulence factors	fliG	-	332 aa	96.4%	100%
virulence factors	fliA	-	239 aa	96.2%	100%
virulence factors	flhA	-	692 aa	94.5%	100%
virulence factors	eutB	-	453 aa	97.6%	100%
virulence factors	cstA	-	701 aa	91.4%	100%

8. CURRICULUM VITAE

MARIJA KVESIĆ IVANKOVIĆ

Date of birth: 8th September 1994

e-mail: mkvesic@unist.hr

EDUCATION

• 09/2020.-today

Postgraduate university study in Biophysics, Faculty of Science, University of Split

• 09/2016.-10/2018.

Graduate university study in Chemistry, Faculty of Chemistry and Technology, University of Split

• 09/2013.-09/2016.

Undergraduate university study in Biology and Chemistry, Faculty of Science, University of Split

EMPLOYMENT

• 03/2019.-today

Research Asistent at Center of Excellence for Science and Technology - Integration of Mediterranean region (STIM), University of Split, Croatia

PROJECTS

• 2022.-today:

Project Leader 'Impact of Submarine Discharges on Antibiotic Resistance in Marine Environment', Croatian Academy of Sciences and Arts

 2021/2022: Project Leader 'Impact of Submarine Discharges on the Development and Spread of Antibiotic-Resistant Human Pathogens', Croatian Academy of Sciences and Arts

AWARDS

• Dean award, Faculty of Science, University of Split, 2016

TEACHING

- Faculty of Science, University of Split, Part-time adjunct assistant for the course *Molecular biology*
- Faculty of Science, University of Split, Part-time adjunct assistant for the course *Algae* and fungi

9. LIST OF PUBLICATIONS

Dželalija, M.; **Kvesić Ivanković**, **M.**; Jozić, S.; Ordulj, M.; Kalinić, H.; Pavlinović, A.; Šamanić, I.; Maravic, A. Marine Resistome of a Temperate Zone: Distribution, Diversity, and Driving Factors across the Trophic Gradient. Water Research 2023. (Under Revision)

Galešić Divić, M., **Kvesić Ivanković, M**., Kišević, M., Panić, M., Lugonja, P., Crnojević, V., and Andričević, R. Prediction of water quality parameters in oligotrophic coastal waters using UAV-obtained hyperspectral data. Ecological Indicators 2023. (Under Revision)

Dželalija, M., **Kvesić, M**., Novak, A., Fredotović, Ž., Kalinić, H., Šamanić, I., Ordulj, M., Jozić, S., Barišić, I.G., Tonkić, M. and Maravić, A., 2022. Microbiome profiling and characterization of virulent and vancomycin-resistant Enterococcus faecium from treated and untreated wastewater, beach water and clinical sources. Science of The Total Environment, 159720.

Kvesić, M., Šamanić, I., Novak, A., Fredotović, Ž., Dželalija, M., Kamenjarin, J., Goić Barišić, I., Tonkić, M. and Maravić, A., 2022. Submarine Outfalls of Treated Wastewater Effluents are Sources of Extensively-and Multidrug-Resistant KPC-and OXA-48-Producing Enterobacteriaceae in Coastal Marine Environment. Frontiers in Microbiology, 1486.

Kvesić, **M.**, Kalinić, H., Dželalija, M., Šamanić, I., Andričević, R. and Maravić, A., 2022. Microbiome and antibiotic resistance profiling in submarine effluent-receiving coastal waters in Croatia. Environmental Pollution, 292, 118282.

Kvesić, **M.**, Vojković, M., Kekez, T., Maravić, A. and Andričević, R., 2021. Spatial and Temporal Vertical Distribution of Chlorophyll in Relation to Submarine Wastewater Effluent Discharges. Water, 13(15), p.2016.