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ABSTRACTS

Content

Торіс	Abbreviation	Page
HUMAN MICROBIOME	HM	4
MEDICAL MICROBIOLOGY	MM	15
NOSOCOMIAL INFECTIONS	Nos	90
NEW ANTIMICROBIAL AGENTS	AntM	122
ANTIBIOTIC RESISTANCE	AntRes	168
VIRAL PATHOGENS	V	229
INFECTIOUS IMMUNOLOGY AND BRM	ll II	270
ZOONOSES	Zoo	281
VETERINARY MICROBIOLOGY	VM	291
FOOD MICROBIOLOGY AND FOOD SAFETY	FM	298
PANEL: MICROBIOLOGY OF ANIMAL	Panel	319
PRODUCTS AND CONSUMER HEALTH		
MICROBIAL BIOTECHNOLOGIES	MB	324
EXTREMOPHILIC MICROORGANISMS	ExM	343
PROBIOTICS & PREBIOTICS	PP	350
ENVIRONMENTAL MICROBIOLOGY	EnvM	366
PARASITOLOGY	Р	390
VARIA	Var	409
WORKSHOP: FROM RESULTS TO A PAPER	WSh	415

EnvM-14

DETERMINATION OF THE LEVEL AND SOURCE OF MICROBIAL POLLUTION IN THE SAVA RIVER BASIN

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Introduction. The contamination of water bodies by sewage or manure is generally determined by fecal indicator bacteria. Specific differentiation between sources of contamination is of particular importance, since the health risk to humans is usually considered higher from human, than from animal fecal contamination. Quantitative PCR (qPCR)-based assays for analysis of human- or animal-associated genetic Bacteroidetes fecal markers have gained increased popularity in the field of microbial source tracking (MST).

Aim. The aim of the present study was to assess the level and source of microbial pollution in the Sava River Basin (SRB).

Materials and methods. The sampling at the SRB was performed in September 2015, at 15 locations. Additional samples were collected from 4 wastewater outlets. Quantification of total coliforms, *E. coli* and enterococci was performed with Colilert Quanti-Tray 2000 and MPN approach. The human-associated BacHum and HF183II, the ruminant-associated BacR and the pig-associated Pig2Bac fecal markers were enumerated by quantitative PCR (qPCR).

Results. According to the concentrations of fecal indicator bacteria, the river samples were scattered within the classes I and II (slight to moderate pollution). According to all monitored indicators, samples collected at 3 wastewater outlets were excessively contaminated. The results of MST revealed the presence of human-associated fecal markers BacHum and HF183II at 13 locations. The ruminant-associated BacR and the pig-associated Pig-2-Bac markers were not detected. High correlation was observed between the standard fecal indicators and human associated fecal markers.

Conclusions. The results reveal human origin of fecal pollution in the SRB and indicates the urgent need for effective wastewater treatment plants in water management.

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