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## Water and microbial monitoring technologies towards the near future space exploration

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#### 23 Abstract

24 Space exploration is demanding longer lasting human missions and water resupply from 25 Earth will become increasingly unrealistic. In a near future, the spacecraft water monitoring 26 systems will require technological advances to promptly identify and counteract contingent 27 events of waterborne microbial contamination, posing health risks to astronauts with lowered 28 immune responsiveness. The search for bio-analytical approaches, alternative to those applied 29 on Earth by cultivation-dependent methods, is pushed by the compelling need to limit waste 30 disposal and avoid microbial regrowth from analytical carryovers. Prospective technologies 31 will be selected only if first validated in a flight-like environment, by following basic 32 principles, advantages, and limitations beyond their current applications on Earth. Starting 33 from the water monitoring activities applied on the International Space Station, we provide a 34 critical overview of the nucleic acid amplification-based approaches (i.e., loop-mediated 35 isothermal amplification, quantitative PCR, and high-throughput sequencing) and early-36 warning methods for total microbial load assessments (i.e., ATP-metry, flow cytometry), 37 already used at a high readiness level aboard crewed space vehicles. Our findings suggest that 38 the forthcoming space applications of mature technologies will be necessarily bounded by a 39 compromise between analytical performances (e.g., speed to results, identification depth, 40 reproducibility, multiparametricity) and detrimental technical requirements (e.g., reagent 41 usage, waste production, operator skills, crew time). As space exploration progresses toward 42 extended missions to Moon and Mars, miniaturized systems that also minimize crew 43 involvement in their end-to-end operation are likely applicable on the long-term and suitable 44 for the in-flight water and microbiological research.

45

## 46 Keywords:

47 International Space Station; space missions; biomonitoring; water biological contamination

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## 66 1 Introduction

Liquid water is essential for all known Earth-derived life forms living in space conditions, 67 68 including microbes (McKay, 2014). The on-going space exploration has not yet demonstrated 69 whether the presence of extra-terrestrial water could indicate itself the occurrence of 70 microbial life, but the prevailing paradigm is that living microorganisms necessitate temporal 71 and spatial proximity with aqueous solutions for their metabolism (Martín-Torres et al., 2015; 72 Merino et al., 2019; Stevenson et al., 2015). In turn, the microbial contamination cannot be 73 thoroughly eliminated from Earth's waters but only controlled and attenuated on the long-74 term time scale (Lopez et al., 2019; Rettberg et al., 2019). Since space exploration has been 75 demanding longer lasting missions, the analysis of waterborne microorganisms turned out of 76 utmost importance for future human spaceflights, planetary outposts, and life-support systems 77 (Horneck et al., 2010).

78 The prevention of crew infectious waterborne diseases is retained among the highest 79 priorities particularly for long duration missions (Ott et al., 2014), since emergency resupply 80 is unrealistic and recycled water could represent the only suitable source for the on-board 81 activities. Most of the aquatic microorganisms found aboard the International Space Station 82 (ISS) do not generally constitute a severe hazard for human health (Blaustein et al., 2019; 83 Checinska Sielaff et al., 2019; Sobisch et al., 2019). However, they may threat astronauts 84 with reduced immune response, mostly following the microgravity stress conditions (Garrett-85 Bakelman et al., 2019; Ott et al., 2016). Other concerning issues arise from microbial 86 influences on spacecraft integrity and function, owing to the potential corrosion and 87 degradation of stainless steel and other materials associated with the electronic equipment 88 and life support systems (Horneck et al., 2010; Yang et al., 2018; Zea et al., 2018). Therefore, 89 there is an increasing interest to improve the spacecraft water monitoring systems to identify 90 and possibly counteract contingent events of microbial contamination (Van Houdt et al.,

91 2012; Yamaguchi et al., 2014).

92 The definition, identification, and test of the microbial monitoring approaches suitable for the 93 on-board water quality control are challenged by several technical constraints (e.g., material 94 safety compatibility, resistance to launch vibration) and a minimal availability for managing 95 excess power, storage, volume, mass, and crew time (Allen et al., 2018). Moreover, selected 96 devices and their supporting reagents must remain viable for years, while operating safely 97 and reliably in extreme conditions (e.g., in the absence of gravity). Technology flexibility is 98 also critical, since monitoring systems should be able to detect different microbial targets 99 (e.g., fungi, protists, prokaryotes, viruses) and to accept samples of various origin, spanning 100 from biomedical (e.g., blood, urine, saliva samples, routine chemistry, cell cultures) to water 101 and environmental samples (Nelson, 2011).

102	In the next years, the Chinese modular space station, built on the experience gained from its
103	precursors Tiangong-1 and Tiangong-2, will be placed and operating in the Low Earth orbit
104	(Gibney, 2019). The sustainable human exploration of the Moon is programmed in the
105	meanwhile (El-Jaby et al., 2019; Pittman et al., 2016), whereas the human missions to the
106	surface of Mars are envisioned before 2040 (ISECG, 2018). Therefore, it is hypothesized that
107	only prospective methodological applications at a high technology readiness level (i.e., at
108	least validated in a flight-like environment; Straub, 2015) will be selected by following their
109	basic principles and current uses in Earth and space-analogue settings.
110	A number of review papers has recently emphasized the need for high-throughput
111	technologies to timely monitor and achieve the stringent microbial quality requirements of
112	future crewed space habitats (De Middeleer et al., 2019; Karouia et al., 2017; Liu, 2017;
113	Moissl-Eichinger et al., 2016; Mora et al., 2016; Yamaguchi et al., 2014).
114	In this article, we narrow the focus on promising bio-analytic technologies for quality
115	assessments of waters in space, with the aim to explore advantages and limitations beyond
116	their current applications on Earth. Starting from the rigorous housekeeping program and the
117	consolidated results of water monitoring activity on the ISS (Duncan et al., 2008; Limero and
118	Wallace, 2017), we provide a critical overview of the microbial monitoring approaches,
119	based on flexible technologies for the identification of microbial components and a total
120	contamination assessment (total microbial burden) in waters circulating on crewed space
121	vehicles.
100	

# 123 2 Water recycle and microbial monitoring aboard the International Space Station 124 2.1 The ISS water cycle

The water recycling system innovations required to support ISS activities have been listed
among the major benefits for humanity (Detsis and Detsis, 2013; NASA et al., 2019). The

127 ISS is provided with potable water from different suppliers, coordinated by the space 128 agencies of United States (National Aeronautics and Space Administration - NASA), Russia 129 (Russian Federal Space Agency - Roscosmos), Europe (European Space Agency - ESA), and 130 Japan (Japanese Aerospace Exploration Agency - JAXA) (Bruce et al., 2005; Van Houdt et 131 al., 2012). All the possible necessary precautions to prevent external contamination are 132 applied throughout water transferring and loading steps over the entire treatment period 133 before the liftoff of supply modules. For instance, the American and Russian waters are 134 produced in conditioned and limited-access areas and preparation facilities, with no risk of 135 accidental water quality modifications during the production process. At the research center 136 of the Italian Società Metropolitana Acque Torino (SMAT), purified waters for space travels 137 are also processed upon selecting well and spring waters that most closely meet the physical, 138 chemical, and bacteriological quality standards for astronauts (Lobascio et al., 2004). 139 Currently on ISS, waters for direct human consumption are regularly delivered and recovered 140 in order to guarantee approximately 4 L per person per day (Figure 1). A reserve of potable 141 water (up to approx. 2000 L) is stored in contingency containers to maintain ISS operations in 142 response to emergency scenarios (Carter et al., 2018). Although routinely monitored and kept 143 constant, the overall water mass balance represents a recurrent major challenge owing to the 144 various ISS water needs (Pickett et al., 2020).

145 Beside the on-demand crew consumption, on-board waters are distributed for different

146 purposes, comprising hygiene and cleaning practices, urinal flushing, oxygen generation via

147 electrolysis, life-support systems, and flexible water-based experimental activities (e.g.,

148 vegetable and food production systems, animal physiology and behavioral adaptation tests)

149 (Baiocco et al., 2018; Chatani et al., 2015; Massa et al., 2016; Niederwieser et al., 2018;

150 Ronca et al., 2019; Wolff et al., 2018). Wastewaters are continuously collected and recycled

151 at high efficiency level (Pickett et al., 2020). In the US segment, the Water Recovery and

152 Management System was reported to recuperate up to 85% from crew urine and flush water,

along with the water content from liquid wastes and humidity condensate from the cabin.

154 Various containers, reservoirs, tanks and bellows are also necessary to maintain water

155 pressure and circulation through the distribution network (Carter et al., 2018).

156 Since microbial growth is unavoidable in persistent stagnation zones and at varying residence 157 times along the water distribution network (Lautenschlager et al., 2010; Ling et al., 2018), the 158 pre- and in-flight addition of biocides is used for residual microbial control. Molecular iodine 159 is applied in the U.S. segment, while the ionic silver level is amended in Russian waters, both 160 at low concentrations (i.e., not detrimental for human health) (Artemyeva, 2016; Lobascio et 161 al., 2004). Moreover, high temperature in the catalytic reactor, multifiltration beds within the 162 Water Processor Assembly, UV-C LEDs within the CO<sub>2</sub> Concentration Assembly of the Advanced Closed Loop System, and novel antimicrobial coatings on various ISS surfaces 163 164 were proven effective against potential microbial biomass growth (Bockstahler et al., 2017; 165 Carter et al., 2018; Perrin et al., 2018; Petala et al., 2020; Roman et al., 2006; Sobisch et al., 166 2019). Finally, the ISS is maintained at pressure and oxygen levels very close to those at sea 167 level on Earth, with a cabin temperature of about 22°C and a relative humidity of about 60%, 168 in order to minimize detrimental growth of microbial biofilms on cabin surfaces (Pierson et al., 2013). 169

170

## 171 2.2 On-board water monitoring and microbial contamination

The achievement and maintenance of water quality standards are evaluated by systematic
monitoring procedures (Limero and Wallace, 2017). Major water physical-chemical
parameters including conductivity, pH, total organic, total inorganic and total carbon, nitrate,
potassium, chloride and ammonium are monitored in-flight, together with iodine and silver
levels (Artemyeva, 2016). Moreover, a robust monitoring program was implemented to verify

that risks of microbial contamination were within acceptable limits in samples collected from different sites of the Russian and US segments, respectively once every three months and each month (Pierson et al., 2013; Van Houdt and Leys, 2012). Crewmembers use handheld equipment to monitor ISS waters and humidity condensate from surfaces. Chemical and biological samples are taken concurrently and at a frequency that may change due to realtime flight necessities (Pierson et al., 2013).

183 Water samples can be processed on-board by cultivation-based methods using the US-184 supplied Water Microbiology Kit for the quantification of total heterotrophic bacteria and 185 coliforms (Bruce et al., 2005). The maximum total number of aerobic heterotrophic viable 186 bacterial cells, counted as colony forming units on a rich agar medium, was internationally 187 defined according to the concentration levels that are achievable with the current prevention 188 and monitoring technologies available and applicable for space (i.e., HPC  $\leq$  50 CFU/ml). 189 Microbial quality standards are also set for ISS internal surfaces, from which humidity 190 condensate is collected (maximum bacterial load =  $10000 \text{ CFU}/100 \text{ cm}^2$ ; maximum fungal  $load = 100 \text{ CFU}/100 \text{ cm}^2$ ) (Van Houdt and Levs, 2012). Along with the on-board monitoring 191 192 activities, archival water samples are regularly collected in teflon bags (Figure 1), preserved, 193 and returned to Earth approximately every three months for post-flight analyses (Limero and 194 Wallace, 2017).

The microbial contamination level was above the former acceptability limits several times during previous in-flight monitoring surveys. Events of microbial biofilm growth within space vehicles were mainly associated with the water layer covering internal surfaces and life support systems (La Duc et al., 2004; Novikova et al., 2006; Roman et al., 2006). Thus, waters collected from humidity condensate were retained among the major sources of microbiological hazard for potable water quality deterioration (Horneck et al., 2010).

201



202

Figure 1. A water drop floating on the ISS (upper left). The Italian astronaut Samantha
Cristoforetti and the Russian cosmonaut Salizhan Sharipov showing the Teflon bag (upper
right), the water tank (lower left) and removal system (lower right), used for drinking
purposes and water storage on the ISS (Credits: ESA and NASA).

208 Several cultivable microbial isolates obtained from spaceflights were mainly affiliated to 209 Bacteria and Fungi (Bruce et al., 2005; Coil et al., 2016; Novikova et al., 2006). For the 210 purposes of this review, it is worth noting that the analysis of the microbial cultivable fraction 211 were likely to provide only a limited snapshot of the highly diverse community found on the 212 ISS by cultivation-independent methods (Checinska Sielaff et al., 2019; Coil et al., 2016; De 213 Middeleer et al., 2019; Ichijo et al., 2016; Lang et al., 2017; Mora et al., 2019; Morris et al., 214 2012). Despite major advantages arise from target-specific isolation and characterization of 215 different types of waterborne microorganisms and pathogens in pure culture, a number of technical and logistic disadvantages characterizes the space application of cultivation-based 216 217 methods. The microgravity conditions provide conflicting results on microbial growth and

218	virulence of opportunistic human pathogens (Morrison et al., 2017; Zhang et al., 2019), while
219	the analysis of archival samples generates a detrimental time gap for a flight-supportive result
220	interpretation (Huang et al., 2018; Novikova et al., 2006). HPC was proven to be also
221	affected by carbon sources in cultivation media, the incubation time, and the initial microbial
222	load level (Amalfitano et al., 2018b). Finally, the prompt development and requirement
223	update of alternate bio-analytical technologies for microbial monitoring during space
224	exploration is pushed fundamentally by the need to avoid microbial regrowth from analytical
225	wastes (Wong et al., 2017).
226	
227	3 Nucleic acid amplification-based methods for microbial identification in space
228	waters
229	Methods based on nucleic acid amplification offer the advantage of specific and fast
230	detection, easy automatization and standardization. Many of the limitations of cultivation-
231	based techniques adopted for bacterial detection and water quality monitoring are
232	overwhelmed by largely reducing the time required for the microbial identification including
233	also viable but not cultivable bacteria and un-cultivable pathogens.
234	
235	3.1 Sample pre-treatments and nucleic acid extraction in space
236	Sample pre-treatment and nucleic acid extraction are key starting points for most of the
237	molecular approaches and product developments dedicated to microbial water quality
238	monitoring. Nucleic acids extracted from water samples are amplified to specific markers.
239	The extraction and purification of nucleic acids rely on cell lysis and the selective binding of
240	cellular DNA and RNA to solid surfaces through filtration and column-based protocols.
241	A fast biomolecular analysis can be conducted in space settings with the in-line
242	implementation of optimized protocols for fast DNA and RNA extraction and sample

243 preparation for sequencing. Published studies reported new promising field-deployable 244 amplification devices and approaches for on board applications (Boguraev et al., 2017; 245 Montague et al., 2018) (Figure 2). Simplified, sample processing and DNA purification, 246 strategies have currently been tested aboard the International Space Station (ISS). The 247 Wetlab-2 project has already developed and tested on the ISS a Sample Preparation Module 248 (SPM) to lyse cells, and to provide high quality extraction of nucleic acids extract, by 249 circumventing operational issues related to microgravity, surface tension alteration, reduced 250 operational space and handling expertise (Parra et al., 2017). 251 Numerous commercialized kits for solid-phase extraction allow handy and rapid nucleic acid 252 purification procedures, in which potential contaminants are removed through sequential 253 washing steps based on centrifugation or DNA separation by paramagnetic beads (Tan and 254 Yiap, 2009). Methods for the direct PCR amplification without DNA extraction were also 255 developed (Williams et al., 2017). Recent papers reported simplified methods for nucleic acid 256 purification using filtration membranes and DNA amplification form the nucleic acid directly 257 on filters (Kaliyappan et al., 2012; Rodriguez et al., 2016). Notably, a cellulose-paper-based 258 dipstick was used to efficiently bind, wash, and elute purified nucleic acids from different 259 matrices without any pipetting or electrical equipment (Zou et al., 2017). 260 Nevertheless, a major bottleneck for the in-flight application of amplification based 261 approaches still lies on the need for time-consuming and waste-producing sample 262 concentration and DNA extraction (Girones et al., 2010). In particular, the method sensibility 263 is lowered when amplifying targets present at low levels over the total extracted DNA 264 (Brandt and Albertsen, 2018). Suboptimal DNA extraction and purification are known to 265 affect results of biomolecular analysis, mainly owing to uncomplete lysis of more resistant 266 bacterial populations and poor removal of DNA polymerase inhibitors (Albertsen et al., 2015; 267 Girones et al., 2010). DNA purification requires effective cell disruption, inactivation of

268 nucleases, and purification of DNA from contaminants that might interfere with the 269 amplification efficiency. Residual extracellular DNA could not be easily discriminated from 270 that of viable dangerous microorganisms, thus leading to an overestimation of risk for human 271 health (Girones et al., 2010). Moreover, waterborne substances can concentrate together with 272 DNA during sample processing and inhibit polymerase enzymes, thus influencing the 273 sensitivity and reliability of the PCR-based microbial detection. Overall, the relative accuracy 274 and precision (i.e., repeatability and reproducibility) of nucleic acid amplification based 275 methods are affected by sample type and pre-treatment methods (concentration and DNA 276 extraction), thus requiring to be evaluated across the whole sample processing (Hospodsky et 277 al., 2010; Kralik and Ricchi, 2017).

278

#### 279 **3.2** Target-based techniques for detection and quantification of nucleic acids

280 The polymerase chain reaction (PCR) is widely used for detecting genes and microorganisms 281 of health concern in water (Ramírez-Castillo et al., 2015). Among these target-based 282 detection approaches, the loop-mediated isothermal amplification (LAMP) of nucleic acids 283 was retained as a rapid and sensible option (Zhao et al., 2015), with minimal requirements for 284 the in-flight water quality monitoring and pathogen detection (Ott et al., 2014). The 285 amplification takes place at isothermal temperature (60-65°C) and positive reactions can be 286 visualised by naked eye (i.e., without post-amplification steps) following the increase of 287 sample turbidity or colour owing to the addition of fluorescent dyes (Notomi et al., 2015). 288 Being less sensible than PCR to inhibition and not significantly influenced by non-target 289 DNA, direct LAMP assays are currently used with good analytical performances (Etchebarne 290 et al., 2017; Samhan et al., 2017). Low amount of DNA can be amplified up to generating  $10^9$ 291 copies within 1 h and producing as final amplification product a complex stem-loop DNA, 292 with several inverted repeats of the target and cauliflower-like structures. Commercial kits for

the rapid on-site detection of water pathogens are already available for terrestrial

applications, also comprising quantitative real-time LAMP, reverse transcription RT-LAMP,

295 in situ LAMP, and viable LAMP (Notomi et al., 2015). For the scopes of this review, it is

worth noting that a rapid method for detecting approximately 1 CFU/100 ml of *Legionella* 

297 *pneumophila* in tap water was developed and efficiently applied on field in less than 2

298 working hours through a direct on-filter LAMP amplification with live/dead propidium

299 monoazide (PMA) differentiation (Samhan et al., 2017).

300 Among the target-based quantification approaches, the quantitative PCR (qPCR) represents a 301 popular technique for the in-flight water quality monitoring (Oubre et al., 2013). The

302 quantification of target sequences is based on the development of a fluorescent signal

303 proportional to the amount of amplified product obtained during the PCR thermal cycles.

304 Multiplex qPCR can be used for the simultaneous detection and quantification of multiple

305 pathogens, consistently reducing analytical time and costs (Ibekwe et al., 2002; LaGier et al.,

306 2004). With low detection limits (<400 cells per sample) and volume requirements (<100 µl

307 of sample), qPCR assays are routinely applied in monitoring plans for the detection and

308 quantification of waterborne pathogens (Girones et al., 2010; Ramírez-Castillo et al., 2015).

309 Good repeatability and reproducibility of qPCR outcomes were reported for gene targets of

310 fecal origin and a standardized workflow achieved consistent results, with low intra- and

inter-laboratory coefficients of variation (median CV = 0.1-3.3% and 1.9.7.1%, respectively)

312 (Ebentier et al., 2013; Orin et al., 2012). A large body of the scientific literature has

313 documented the development and use of qPCR for pathogenic viruses, bacteria, protozoa, and

314 fungi (Kralik and Ricchi, 2017; Ramírez-Castillo et al., 2015). Through a pre-treatment with

315 cell membrane impermeant DNA intercalating dyes, the so-called Viable qPCR was applied

316 for discriminating between viable (with intact membrane) and dead (with damaged

317 membrane) bacteria, allowing the quantification of water- and food-borne pathogens such as

318 Campylobacter, E. coli O157:H7, Legionella pneumophila, Salmonella, Cryptosporidium

319 (Banihashemi et al., 2012; Brescia et al., 2009; Delgado-Viscogliosi et al., 2009).

320 Given the versatile applications of the numerous available assays, both LAMP and qPCR

321 have been retained as suitable water monitoring methods for long-term exploration missions.

322 Owing to the high sensitivity, specificity, and simple post-amplification steps to detect the

323 amplified targets, LAMP was proposed by the Japanese Aerospace Exploration Agency as

324 alternative microbial contamination monitoring system for the ISS (Ott et al., 2014). For the

325 analysis of crew health related genetic modifications, qPCR technologies have been

326 successfully tested on-board the ISS within the projects Gene in Space and Wet-lab2, devoted

327 to definition of a robust, user-friendly nucleic acid extraction and sequencing approach

328 aboard ISS (Boguraev et al 2017; <u>https://www.genesinspace.org/;</u>

329 <u>https://www.nasa.gov/mission\_pages/station/research/experiments/1913.html;</u>

330 <u>https://www.nasa.gov/ames/research/space-biosciences/wetlab-2</u>). Moreover, the RAZOR EX

331 PCR, launched on Space-X 9 (July 2016) within Water Monitoring Suite project will allow

332 performing direct PCR amplification from water samples.

333 In all current space applications, however, the selected target-based detection and

334 quantification approaches require the use of disposable materials and labour intense

335 protocols, which will inevitably reduce their long-term applicability in space conditions. The

336 general precautions used on Earth to limit contamination risks, including the most stringent

337 procedures applied in clean rooms (Rettberg et al., 2019), might also represent a limiting

338 practical issue in the small close spacecraft environment, since the high sensibility of PCR-

based detection increases the chance of amplifying carry over contamination, with the

340 consequent production of false positive results. This is crucial for LAMP owing to the limited

341 accessibility to degradation of DNA products, while a major limitation of qPCR is also the

342 occurrence of inhibitors that can be co-concentrated or extracted along with nucleic acids

from the target microorganisms (Gibson et al., 2012). The presence of qPCR inhibitors
introduces a number of problems, ranging from low amplification efficiency and reduced
assay sensitivity to complete reaction failure and false negative results (Radstrom et al.,
2008).

347

348 Sequencing-based "-omics" approaches for microbial community characterization 3.3 349 The High-Throughput Sequencing (HTS) encloses a popular suite of technologies, 350 methodological approaches, and data elaboration workflows used to characterize the 351 phylogenetic composition of the total microbial community in different aquatic matrices. The 352 taxonomic classification is based on the huge amount of sequences generated by either a 353 portion of the cellular nucleic acid content or the whole genome, through the so-called 354 amplicon and shotgun sequencing approaches, respectively (Peabody et al., 2015). 355 The amplicon analysis represents the extension of sequencing based methods of organisms' 356 classification, defined by specific protocols under the general name of "genetic" or 357 "molecular barcoding" (Hebert and Gregory, 2005), which is based on the information 358 carried by a single conventional marker, such as the 16S rRNA for microorganisms. 359 The shotgun sequencing consists in generating millions of DNA fragments of different 360 lengths from a starting pool of genomes. These fragments cover a quote of the original 361 genomes inversely proportional to the genome lengths, and taxonomic assignment may be 362 quantitative according to the proportion of fragments classified for each taxon. The amplified 363 products are sequenced, controlled for quality, and assembled in longer contigs. The obtained 364 sequences are classified at different taxonomic levels, according to the best match with a 365 reference sequence database and following different similarity criteria (Segata et al., 2012). 366 Currently, the most used sequencing platforms are characterized by different technical 367 principles and include Roche 454 GS-FLX (pyrosequencing), Illumina MiSeq and HiSeq

368 (reversible terminator sequencing by synthesis), Ion PGM (semiconductor based sequencing
369 by synthesis), and the nanopore GridION and MinION<sup>™</sup> (Check Hayden, 2015; Clooney et
370 al., 2016; Ghanbari et al., 2015; Glenn, 2011).

371 Performance evaluation and standardization of HTS workflows are still limited for water 372 microbial community characterization. The DNA extraction procedures and primers sets were 373 reported to influence the assessment of the bacterial community composition in drinking 374 waters, with a generally effective representation of the core abundant taxa at total cell abundance levels of 10<sup>3</sup>-10<sup>5</sup> cells/ml (Brandt and Albertsen, 2018). Moreover, the method 375 376 sensitivity was detrimentally affected owing to the occurrence of contaminating bacteria in 377 extraction kits and laboratory reagents (Salter et al., 2014). 378 HTS-based technologies have been already tested under microgravity conditions and directly 379 on the ISS (Carr et al., 2020; Castro-Wallace et al., 2017; McIntyre et al., 2016). The most 380 suitable candidate technology to be transported and mounted on the ISS was the MinION 381 pocket-size device (Figure 2), which can provide rapid identification up to 300 kb with single 382 strand and 60 kb with double strands reads (Jain et al., 2016). The integrity and activity of 383 nanopores, in which DNA/RNA molecules pass through during base reading, were not 384 adversely affected by storage conditions, launch, cosmic radiations or handling in 385 microgravity, and reusability of flow cells was warranted. Specific indications to avoid air 386 bubbles interference at the nanopores were implemented (Castro-Wallace et al., 2017; 387 Rizzardi et al., 2016). Moreover, MinION performances in sequencing a mixture of genomic 388 DNA from a virus, a bacterium, and of mammal mtDNA, were comparable to those of 389 MinION, Illumina MiSeq and PacBio RSII run on ground. Accuracy respect to MiSeq and 390 RSII was slightly inferior (89% identity to the reference genome, respect to > 99%), but 391 sufficient for sequence analysis. Sequencing reads were successfully assigned to the correct 392 reference sequences in 90% of cases. Likely, with the novel improved versions of flow cells,

393 the molecular detection of aquatic microorganisms will improve notably (Kilianski et al., 394 2015). Tests also demonstrated that real-time metagenomics analysis was also possible by 395 using a laptop device that can be integrated in the ISS hardware. Considering genomic data 396 analysis, one of the least demanding platforms has been tested successfully simulating flight 397 conditions and hardware availability onboard of the ISS, demonstrating its realistic 398 applicability even on a laptop base (Castro-Wallace et al., 2017). Provided the adaptation of 399 hardware and instruments to the spatial and logistic limitations onboard, both the shotgun and 400 amplicon approaches, can be considered as suitable tools for microbial community

401 characterization and pathogen detection on the ISS.

402 The amplicon approach requires straightforward PCR amplification protocols, downstream 403 bioinformatics, and database searches. Nevertheless, this technique is prone to fail to detect 404 unknown or highly divergent 16S rRNA gene sequences, which may be associated to novel 405 or previously undetected pathogens. Moreover, taxonomic assignment is highly sensitive to 406 the gene region selected as a marker and to the assignment method (Claesson et al., 2010; Liu 407 et al., 2008; Tremblay et al., 2015), while accuracy of quantification can be affected by 408 the variability in the number of gene copies shown by microbial taxon and even between 409 individual cells of the same species (Větrovský and Baldrian, 2013). Finally, a common 410 scientific view of suitable markers for viruses is still missing (Chakraborty et al., 2014). 411 In comparison to amplicon analysis, the shotgun sequencing offers a wide genome coverage 412 of organisms, allowing the detection of unknown contaminants and the possibility to screen 413 for both bacteria and viruses. In the field of water quality monitoring, the shotgun sequencing 414 was used, for instance, as the benchmark method to assess the occurrence of potential 415 multiple waterborne pathogens, novel indicators for human sewage contamination, and 416 microbial safety of drinking water in relation to the efficiency of reclamation treatments 417 (Chao et al., 2013; Lu et al., 2015; McLellan and Eren, 2014; Newton et al., 2015). Although

the possibility to obtain up to a few thousands of base pairs for each read improved the microbial identification specificity, non-trivial disadvantages include the initial skimming of the huge number of produced reads, their quality check and assemblage in contigs useful for classification, which require high computational effort and time (Tan et al., 2015).



423

422

424 **Figure 2**. The NASA astronaut Peggy Whitson performing the Genes in Space investigation

425 on the ISS using the miniPCR and MinION. Credits: NASA

426

# 427 4 Real-time technologies for early warning microbial monitoring

428 The total contamination assessment (total microbial burden) through the accurate

429 quantification of microbial cell abundance and viability in waters circulating is a recognized

430 necessity on space crewed vehicles (Morris et al., 2012). Different early-warning real-time

431 methods that target parameters at the single-cell level (e.g., cellular biomolecules, membrane

432 integrity, enzyme activity, substrate uptake) have been developed for water monitoring, but

433 only few can efficiently operate in flight-like and space settings.

#### 435 **4.1 ATP-metry**

An option for real-time monitoring of biological contamination in water samples is the 436 437 analysis of adenosine triphosphate (ATP) cell content. The analysis is carried out through the 438 chemical and/or enzymatic extraction of this molecule from microbial cells, followed by the 439 measurement of light emission derived when the dissolved ATP, in presence of magnesium, 440 reacts with the luciferine (substrate) - luciferase (enzyme) complex. The emitted light 441 intensity is linearly related to the ATP concentration, easily measurable using a luminometer, 442 and gives virtually instant information (within minutes) of the metabolically active microbial 443 population. This peculiarity makes it suitable as an early-warning approach for measuring 444 bacterial contamination and monitoring water treatment efficacy in near real-time (Hammes 445 et al., 2010). The method is robust, easy to perform, and suitable to detect both cultivable and 446 uncultivable cells, with better estimates of total active microorganisms compared to 447 heterotrophic plate counts. The concomitant use of traditional cultivation-based approaches 448 showed levels of cultivable cells order of magnitude lower that those estimated by ATP 449 (Siebel et al., 2008; Zhang et al., 2019).

450 The ATP analysis was performed on pre-flight and post-flight ISS water samples and revealed a biological contamination ranging between 0 (drinking water) and 4.9x10<sup>4</sup> cells/ml 451 (humidity condensate) (Bacci et al., 2019; La Duc et al., 2004). Recently, the viable microbial 452 453 contamination on-board ISS was reliably monitored on surface samples by intracellular ATP 454 measurement (Perrin et al., 2018; Venkateswaran et al., 2003). ATP assays were also proven 455 effective in monitoring microbial contamination on surfaces from an inflated lunar/Mars 456 analogous habitat during long-term human occupation (Mayer et al., 2016). The total (ATP 457 content from both dead and live microbes) and viable microorganisms (intracellular ATP content) were in the range of  $10^5$ - $10^6$  relative luminescence unit (RLU)/m<sup>2</sup>. These results 458 were in line with ATP data measured directly on-board within the experiment T2 carried out 459

460 during Euromir-95 mission (Guarnieri et al., 1997). Recently, the ATP-metry was selected for 461 the real-time monitoring of the biological contamination on board ISS within the H2020 462 European project BIOWYSE (Biocontamination integrated control of wet systems for space 463 exploration, http://biowyse.eu), aimed at developing a compact, automatic, and microgravity-464 compatible on-board systems for the prevention, monitoring and control of microbial load in 465 waters and on wet surfaces (Figure 3). A humid area sampler was recently developed and 466 patented (ref. 102018000009137, dated 03/10/2018) (Detsis et al., 2018; Guarnieri et al., 467 2019).

468 Today, commercial kits for quantitative ATP-metry are available to monitor the microbial 469 biomass level in water and to validate cleaning and disinfection procedures, with a wide 470 number of bulk and intracellular ATP measurements on microbial communities reported from 471 natural and engineered aquatic environments (Abushaban et al., 2019; Fillinger et al., 2019; 472 van der Wielen and van der Kooij, 2010; Vang et al., 2014). Although the apparent 473 advantages, major technical limitations are related to the low sensitivity and result 474 reproducibility at low cell concentration and sample volume ( $< 100 \mu$ ), along with the 475 susceptibility to environmental conditions (e.g., pH, temperature, occurrence of enzyme 476 inhibitors). Some drawbacks of the method are partly circumvented by using external and 477 internal standards and by operating under controlled reproducible settings. Therefore, the 478 correlation between intracellular ATP content and microbial cell counts will rely on a robust 479 cross-calibration with results from other reference methods (Amalfitano et al., 2018b; 480 Hammes et al., 2010). It has been calculated that the average ATP-per-cell content is approximately 1.75 x 10<sup>-10</sup> nmol/cell or 6.87 x 10<sup>-17</sup> g ATP/cell (Zhang et al., 2019). 481 482 However, the amount of per-cell ATP can be significantly influenced by the phylogenetic 483 affiliation and the cellular physiological status, with diverse ATP-content reported between

484 either eukaryotic or prokaryotic cells from the same cultures (Bajerski et al., 2018; Yaginuma485 et al., 2015).

486

# 487 **4.2** Flow cytometry

488 Ubiquitously applied from the bio-medical research to environmental sciences, flow 489 cytometry (FCM) is considered an unparalleled high-throughput technology for single cell 490 analysis (Robinson and Roederer, 2015). This generic technology allows the measure (-491 metry) of the optical properties of cells (cyto-) transported by a liquid sheath (flow). 492 Following sample intake, a pressurized laminar flow is generated and suspended particles are 493 individually forced to cross a light source excitation point for scanning and evaluation. Light 494 scatter and fluorescence signals (either with or without a secondary staining with fluorescent 495 dyes) are detected at the single-cell level and instantly converted into digital information to 496 be shown on multidimensional plots, along with real-time data analysis and statistics

497 (Shapiro, 2005).

498 Flow cytometry is included in the roadmaps of national space agencies worldwide and 499 deemed as a necessary technology for defining and monitoring spaceflight-associated 500 requirements since early 80's. Owing to the high versatility for diagnostic medicine (e.g., 501 hematology, immunology, and physiology), this technology was so far retained as a prime 502 asset for health monitoring and clinical laboratory diagnostics for astronauts, in view of the 503 upcoming deep-space exploration missions (Crucian and Sams, 2012). A flow cytometry 504 platform was already successfully tested on-board the ISS to monitor and understand the 505 physiological adaptations of astronauts to microgravity (Dubeau-Laramée et al., 2014; Phipps 506 et al., 2014) (Figure 3). Advanced developments of the prototype were based on 507 commercialized flow cytometer with significant additional engineering modifications, mostly 508 aimed to generate laminar particle flow (distinct from the standard sheath fluid based

509 method), and to reduce the significant amount of liquid biohazardous waste and energy 510 operating requirements. The laminar flow within the flow chamber was found to be 511 dramatically altered by microgravity. In turn, the unavoidable elimination of the fluid 512 mechanical setting for particle hydrodynamic focusing was proven to significantly reduce 513 liquid waste and the total system operational load, in terms either of instrument size and 514 weight or energy consumption (Cohen et al., 2011; Crucian and Sams, 2005). 515 Notwithstanding the recent system design improvements, peripheral blood cells has been 516 retained as the only target for first studies in space environments (McMonigal and Crucian, 517 2015). Since a flow cytometer can provide support to a wide range of scientific applications 518 (e.g., biology, microbiology, and environmental science), it is needless to point out that a 519 spaceflight compatible machine could satisfy the unmet flight requirements for water 520 monitoring and treatment to complete future long-duration missions in closed healthy 521 environments.

522 The high sensitivity for scanning very small objects (i.e., from virus-like-particles to 523 prokaryotes, to pico- and micro-eukaryotes) and the wide detectable cell concentration range (generally between  $10^2$  and  $10^7$  cells ml<sup>-1</sup> without concentrating or diluting the samples) 524 525 represent unmatched features of this technology for water monitoring. A broad suite of assays 526 is available for microbial quality assessments in natural and engineered aquatic ecosystems 527 (Amalfitano et al., 2014; Boi et al., 2016; Gasol and Morán, 2015; Van Nevel et al., 2017). 528 Successful applications are reported to provide early warning indications of unexpected water 529 contamination events through the rapid detection of a variety of microbiological threats, 530 including pathogenic and potentially toxic microorganisms (Vital et al., 2010; Weisse and 531 Bergkemper, 2018; Yang et al., 2010). Moreover, flow cytometric measurements are suitable 532 to evaluate the efficiency of various industrial microbial bioprocesses (e.g., food and pharmaceutical preparations) (Díaz et al., 2010), and the performances of engineered systems 533

534 for water treatment (Besmer and Hammes, 2016; Safford and Bischel, 2019). Remarkably, 535 FCM was also useful to determine the presence of nonliving organic and inorganic 536 substances, including nano- and micro-sized particles, suspended solids, flocs and aggregates 537 of various origins (Aulenta et al., 2013; Casentini et al., 2016; Liu et al., 2016). On-going 538 instrumental developments for on-site applications are directed to install flow cytometers on 539 either mobile units (e.g., ships and vehicles) or fixed locations (e.g., treatment plants, marine 540 buoys, off-shore platforms), with the possibility for automatic programmable staining of 541 aquatic microorganisms and remote data transfer (Buysschaert et al., 2018; Pomati et al., 542 2011; Thyssen et al., 2007). 543 Leaving aside the high costs of all sophisticated systems, the need for specially trained staff is 544 likely to represent a major drawback in the daily scheduled monitoring practices. Moreover, 545 the reproducibility of cytometric data was reported to be adversely affected by changing cell 546 staining protocols (e.g., fixatives, type of fluorescent dye), incubation conditions (e.g., 547 temperature, time to analysis), instrumental settings (e.g., fluidic and signal amplification 548 systems), and the source water (e.g., from natural or engineered systems) (Nescerecka et al., 549 2016; Prest et al., 2013). Further work is also required to establish user-independent strategies 550 for gating and data handling (Amalfitano et al., 2018a; Koch et al., 2014). One more specific 551 challenge is the lack of phylogenetic resolution. Despite the enumeration of targeted taxa may 552 rely on specific fluorescence staining procedures (Couradeau et al., 2019; Manti et al., 2011; 553 Neuenschwander et al., 2015), the cytometric information characterizing different 554 subpopulations of microbial cells is generally ataxonomic. A direct link between cytometric 555 fingerprinting and microbial diversity, assessed by 16S rRNA gene amplicon sequencing, 556 was demonstrated in recent studies (Props et al., 2017, 2016). However, the computational 557 workflows are convoluted and still under development (Rubbens et al., 2019). 558



Figure 3. Integrated breadboard of the ATP-metry system, developed within the European
H2020 project BIOWYSE. The hardware was designed (upper left photo) and built (upper
right photo) to fly within the European Drawer Rack Mark 2. The photo below shows the
Canadian astronaut Chris Hadfield holding the flow cytometry platform Microflow1,
successfully deployed and tested on the ISS (Credits: BIOWYSE project consortium and
NASA).

566

# 567 5 Comparative analysis of space-relevant technological features

568 The procedural workflows of each of the mature technologies, herein entitled for water

- 569 microbial monitoring in space, comprise major steps and provide different types of results.
- 570 The overall time-to-results can vary from minutes to days, largely relying on user-dependent
- 571 manual operations and technological solutions for automation (figure 4).



572

Figure 4. Workflows of mature technologies entitled for microbial water monitoring during
space missions. Cultivation-, nucleic acid-based, and real-time methods are suitable to
identify and quantify the waterborne microorganisms, ideally present in a drop of ISS water
and visualized by epifluorescence microscopy on a filtration membrane (upper left pictures).
Comparative levels of readiness, analytical performances, and practical issues were reported
along with major procedural steps, timing, and types of achievable results.

580 Both current and future applications in space will be necessarily bounded by the definition of

581 novel standards of microbial quality (i.e., other than those applied on Earth by cultivation-

582 dependent approaches) (Amalfitano et al., 2018b), but also by a compromise between the best 583 analytical performances and detrimental practical issues to cope with during spaceflights. 584 The implementation of successful on-board workflows will critically rely on the overall water 585 cycle management (Pickett et al., 2020), while fundamental analytical aspects can be 586 constrained by the minimal amount of water available for routine monitoring activities. For 587 example, technological accuracy and result reproducibility are directly linked to workable 588 water volumes and the constitutive occurrence of microbial targets in the sample (e.g., 589 microbial cell abundance, per-cell gene copies, cell viability). Volumes required for accurate 590 and reproducible analyses will reasonably range between few tens of microliters (e.g., for 591 flow cytometric assessments) and hundreds of milliliters (e.g., for cultivation- and nucleic 592 acid-based methods), as also reported in terrestrial studies (Safford and Bischel, 2019). 593 Notably, sample filtration, a major pre-treatment step for ATP-metry and nucleic acid-based 594 methods, can selectively concentrate the microbial biomass and target microorganisms, while 595 minimizing the water loss for analytical needs. The filtering surfaces are also suitable for 596 visual inspection (e.g., by microscopy), storage, and comparative analysis (e.g., on Earth). 597 However, filtration can influence the composition of the dissolved organic matter of the 598 permeate water, possibly triggering microbial regrowth in the downstream distribution 599 system (Park et al., 2018), and it will also necessitate additional crew time for manual 600 operations (e.g., filter substitution, regeneration, disposal). 601 Sufficient data are not yet available from real space conditions to implement protocol details 602 within the procedural workflows. Although the selected technologies were demonstratively 603 applied under either simulated or real microgravity conditions, their own critical advantages 604 and limitations will require full reconsideration for flight-like and space applications.

From the one hand, analytical benefits will necessarily include the time needed to achieve

606 results (i.e., speed to results), the accuracy and flexibility in detecting specifically-selected

microbial targets (i.e., identification depth) with reproducible consistent results (i.e., 607 608 reproducibility), and the number and multiple type of achievable results (i.e., herein named 609 multiparametricity). On the other hand, the operating conditions can be particularly stringent 610 and limiting the direct applicability in space, unless addressing critical requirements such as 611 reagent usage, waste production, operator skills, and crew time (table 1). 612 Using a simplistic pairwise comparison, we sought to emphasize that few selected 613 technological features have to be consciously retained from the methodological proof-of-614 concept level up to the device deployment, instrumental demonstration, and routine use in the 615 on-board housekeeping program. Although the comparative scores were assigned 616 subjectively (table S1), it is likely evident that the stringent requirements of microbial water 617 monitoring in space cannot be met by a single technological solution. 618 619 Table 1. Advantages and limitations of the most promising approaches for microbial 620 monitoring in space settings. The selected technologies are flexible (i.e., applicable to 621 different microbial targets in samples of various origin), suitable for miniaturization and

622 automation with limited maintenance, and already tested in flight-like conditions. A

623 comparative score was arbitrarily assigned through a pairwise comparison matrix for each of

624 selected space-relevant technological features, including major analytical performances (i.e.,

625 speed to results, identification depth, reproducibility, multiparametricity – green marks) and

626 practical issues to cope with in space (i.e., reagent usage, waste production, operator skills,

627 crew time – red marks).

628

Technology	Advantages	Limitations	Analytical performances/ Practical issues	Comparative Scores	Applications in space
Plate Cultivation	<ul> <li>No pretreatment</li> <li>Detection limit 1 cell/100 ml</li> <li>Low equipment requirements</li> </ul>	<ul> <li>No direct quantification</li> <li>Risks of contamination</li> <li>Reliant on cultivation conditions</li> </ul>	Speed to results Identification depth Reproducibility Multiparametricity Reagent usage Waste production Operator skills Crew time		La Duc et al., 2004 Morrison et al., 2017 Pierson et al., 2013
LAMP	<ul> <li>Detection limit ~10<sup>2</sup> gene copies/ml</li> <li>Unaffected by template conc.</li> <li>No effects of inhibitory compounds</li> </ul>	<ul> <li>No direct quantification</li> <li>Risks of contamination</li> </ul>	Speed to results Identification depth Reproducibility Multiparametricity Reagent usage Waste production Operator skills Crew time		Ott et al., 2014
qPCR	<ul> <li>Target-specific and quantitative</li> <li>Detection limit ~10<sup>3</sup> gene copies/ml</li> </ul>	<ul> <li>Pretreatment</li> <li>processing</li> <li>Affected by</li> <li>template and</li> <li>inhibitory</li> <li>compounds</li> <li>Reliant on PCR</li> <li>related issues</li> </ul>	Speed to results Identification depth Reproducibility Multiparametricity Reagent usage Waste production Operator skills Crew time		Boguraev et al., 2017 Parra et al., 2017
High Throughput Sequencing	<ul> <li>In-depth phylogenetic resolution</li> <li>Specific for unknown non targeted microorganisms</li> </ul>	<ul> <li>Pretreatment processing</li> <li>Reliant on PCR related issues</li> <li>Complex data interpretation</li> </ul>	Speed to results Identification depth Reproducibility Multiparametricity Reagent usage Waste production Operator skills Crew time		Castro-Wallace et al., 2017 McIntyre et al., 2016
ATP-metry	<ul> <li>Real-time data (&lt; 5 min)</li> <li>Detection limit ~0.1 pg/ml</li> </ul>	<ul> <li>Unspecific detection</li> <li>Destructive analysis</li> <li>Risks of contamination</li> </ul>	Speed to results Identification depth Reproducibility Multiparametricity Reagent usage Waste production Operator skills Crew time		Guarnieri et al., 1997 La Duc et al., 2004
Flow cytometry	<ul> <li>No pretreatment</li> <li>Quantitative and near-real time data (&lt; 20 min)</li> <li>Detection limit ~10<sup>2</sup> cells/ml</li> </ul>	<ul> <li>Complex data interpretation</li> <li>Clogging issues</li> </ul>	Speed to results Identification depth Reproducibility Multiparametricity Reagent usage Waste production Operator skills Crew time		Dubeau- Laramée et al., 2014 Phipps et al., 2014

#### 631 6 System miniaturization and future challenges

632 An important point is that the monitoring technologies can be mission-dependent, but only 633 those instruments that minimize crew involvement in their end-to-end operation are likely to 634 be applicable on the long-term (Karouia et al., 2017). Overall, the selected devices have to be 635 compact, suitable for automation, low power-consuming, and virtually invisible except when 636 needed (Limero and Wallace, 2017). The ground-based counterparts have already been 637 miniaturized using microfluidics, but the deployment of monitoring-effective tools onboard 638 spacecrafts will also require substantial reengineering and instrumental customization. The 639 instruments deployed so far have not been yet permanently used in spaceflight water 640 monitoring, yet numerous examples of successful international projects and commercial 641 partnerships dedicated to the human space exploration let argue that critical space waterrelated tasks can be operatively accomplished at reasonable costs and times (< 5 years) 642 643 (Karouia et al., 2017). 644 Considerable progress has been made in miniature the onboard instrumentation to assess the 645 water microbiological contamination. This also includes microbial cultivation systems, such 646 as the AquaPad developed by the French CNES space agency (Augelli, 2018). The

647 microfluidic chips that allow cell isolation and incubation have been developed for ground-

648 based applications, and appear suitable for space uses. For example, the encapsulation of

649 single cells from a mixed microbial community into small droplet compartments of a water-

650 in-oil emulsion can offer further opportunities for physiological studies and viability assays

652 populations isolated into plugs or micro-Petri dishes (Boedicker et al., 2008; Boitard et al.,

(e.g., metabolic by-products diffusion, resistance to toxicants, enzymatic activities) on clonal

653 2015).

651

Considering the ability of microfluidic systems to efficiently conduct measurements on small
 volumes of complex fluids without the need for a skilled operator, lab-on-a-chip technologies

656 and portable diagnostic devices have gained increased popularity for sensing a wide range of 657 water parameters and microbial pathogens even in the most remote settings (Mairhofer et al., 658 2009). More recently, new qPCR platforms based on microfluidic technologies have been 659 developed allowing the simultaneous analysis of numerous genes and samples in volume 660 chambers of few nanoliters, placed at high density on a chip (Ricchi et al., 2017). Being less 661 sensitive to inhibitors than qPCR, the digital PCR (dPCR) is mainly applied to monitor gene 662 transcriptions in microbial cells without the need of a standard curve for gene copy 663 quantification (Devonshire et al., 2016). 664 The system miniaturization was also considered the most appealing trait for space 665 applications of the nanopore DNA sequencer MinION (Castro-Wallace et al., 2017). By

assembling miniaturized and lab-on-a-chip solutions, the Water Monitoring Suite developed

by NASA represents so far the best performing custom-built device applied successfully on

the ISS to monitor different water quality properties. Along with the HACH colorimeter and

the Organic Water Module, respectively used for inorganic and organic chemical

670 assessments, the hardware suite also includes the PCR-based instrument RAZOR EX, with

671 customized sample pouch kits containing all pre-loaded reagents necessary for sampling,

sample preparation, and real-time PCR

673 (https://www.nasa.gov/mission\_pages/station/research/experiments/explorer/Investigation.ht

674 <u>ml?#id=1847</u>). However, as space exploration progresses toward extended missions to cis-

675 lunar space and Mars, PCR-based and multi-omics instruments onboard spacecraft should not

676 be considered in separation from other technologies needed for the in-flight microbiological

677 research (Karouia et al., 2017).

678 In particular, microfluidic platforms developed for the direct cell detection showed promising

679 perspectives because of the potential for precise and easy-to-use analytical procedures. While

680 enhancing analytical performances, the system miniaturization presents also the advantages

681 of reduced consumption of reagents and the ability to integrate monitoring and isolation 682 procedures within a single device (Auroux et al., 2002). A simple microfluidic system was 683 successfully tested for rapid and semi-automated bacterial enumeration in freshwaters and 684 promising outcomes suggested its applicability to drinking waters under both ground and 685 space conditions (Yamaguchi et al., 2014). Moreover, both sensitivity and specificity of ATP-686 metry and FCM can be also improved by miniaturizing the core systems down to scales 687 closer to the ones of microorganisms. A micro-fluorescence-activated cell sorting (µFACS) 688 was used to sort out cells of interest by changing the flow direction after cell detection (Fu et 689 al., 1999). An integrated platform that combines two different force fields in a single 690 microfluidic device (Dielectrophoretic-Magnetic Activated Cell Sorter - iDMACS) was 691 applied for simultaneous sorting of multiple bacterial targets (Kim and Soh, 2009). More 692 recently, a high-throughput Raman flow cytometer was developed on a microfluidic chip for 693 the label-free molecular fingerprinting at the single-cell level (Hiramatsu et al., 2019). 694 The onboard laboratory miniaturization included also the fluidic components (e.g., pumps, 695 valves, electronics), thus paving the way to the use of advanced biosensors for screening food 696 safety and water quality in space (Roda et al., 2018). Following the proofs of concept and 697 wearable technologies suited to monitor astronauts' health, the biosensing diagnostic 698 instrumentation, most reasonably based on electrochemical and optical detectors, was argued 699 as a secondary future option for in-flight water biochemical analyses (Choi et al., 2018; 700 Limero and Wallace, 2017; Nelson, 2011). In particular, the amperometric biosensors were 701 proven sensitive to monitor different water analytes, chemical contaminants (e.g., pesticides, 702 organophosphates, carbamates), and numerous microbial biomarkers successfully targeted to 703 detect the major microbiological agents, food- and water-borne pathogens (e.g., E. coli, 704 Salmonella, L. monocytogenes, C. jejuni, B. cereus, M. smegmatis) (Grieshaber et al., 2008; 705 Velusamy et al., 2010). However, despite providing concrete benefits for health services and

our life on Earth and beyond, the spectroscopic and biosensing devices are still at a low level
of technological readiness for in-flight applications (García-Descalzo et al., 2019; Own et al.,
2019; Roda et al., 2018), thus falling out of the scopes of this review.

709 In general, a critical aspect of all microfluidic and lab-on-a-chip solutions is bound to the 710 very limited system reusability. Most miniaturized devices for terrestrial applications are of 711 single-use and, in many cases, based on disposable cartridges that cannot be safely stowed 712 onboard during long-term space missions. When cells are delivered to different system 713 compartments for collection, there is no clear approach for removal the analyzed samples 714 without compromising the system functioning and risking contamination. Clogging issues 715 may also interfere with the analyses due to the processing of large sample volumes. Reusable 716 systems have been tested, but their applications in space are likely constrained by the risk of 717 sample carryover, reduced analytical performance upon extended reuse, and the resources 718 required for cleaning and reactivation procedures. In this regard, the technological 719 development is highly demanding and still far from being accomplished in a near future.

720

## 721 **7** Conclusions

Current technologies for water microbial monitoring can satisfy the needs of long-term
 space exploration missions at reasonable costs and times, although substantial
 instrumental reengineering has to be considered.

The suitable methodological applications at a high technology readiness level (i.e., at
 least validated in a flight-like environment) will require reduced space to be allocated and
 can potentially provide rapid and specific responses regarding the in-flight occurrence of
 the microbiological contamination.

The advanced biomolecular characterization of water samples from the ISS is promoting
 a better understanding of the onboard levels and patterns of microbial contamination, thus

- contributing to the development of space bound technologies for the rapid and specificidentification of microorganisms of health concerns.
- As space exploration progresses toward longer missions, PCR-based and multi-omics
   approaches can be complemented by real-time technologies needed for the in-flight
   microbiological research and suitable for the early-warning microbial monitoring of space
   waters.
- 737

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