

1 **Next-Generation Food Research: Use of Meta-Omic**
2 **Approaches for Characterizing Microbial**
3 **Communities Along the Food Chain**

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24 **Abstract**

25 Microorganisms exist along the food chain and impact the quality and safety of foods in both
26 positive and negative ways. Identifying and understanding the behaviour of these microbial
27 communities enables the implementation of preventative or corrective measures in public
28 health and food industry settings. Current culture-dependent microbial analyses are time-
29 consuming and only target specific subsets of microbes. However, the greater use of culture-
30 independent meta-omic approaches has the potential to facilitate a thorough characterisation
31 of the microbial communities along the food chain. Indeed, these methods have shown
32 potential in contributing to outbreak investigation, ensuring food authenticity, assessing the
33 spread of antimicrobial resistance, tracking microbial dynamics during fermentation and
34 processing, and uncovering the factors along the food chain that impact food quality and
35 safety. This review examines the community-based approaches, and particularly the
36 application of sequencing-based meta-omics strategies, for characterizing microbial
37 communities along the food chain.

38

39 **INTRODUCTION**

40 Microorganisms along the food chain from farm to fork influence food quality and safety.
41 Historically, culture-based techniques have been used extensively to characterise these
42 microbes. However, with the development of molecular methods and high-throughput
43 sequencing technologies, culture-independent techniques have become more relevant to food
44 microbiome analysis. This has resulted in a corresponding shift from the investigation of
45 specific taxa or groups of food microorganisms to a broader community-based analysis
46 (Cocolin and Ercolini 2015). These methods are based on the extraction of nucleic acids,
47 proteins, and/or metabolites, allowing for the detection and characterisation of microbes
48 within an environment without the need for culturing. As the occurrence of and interactions
49 between microorganisms impact the quality and safety of food, a deeper understanding of
50 these processes allows for early interventions to adverse food safety events, ensuring optimal
51 food quality, and identifying the source of desirable or undesirable microorganisms.

52

53 Despite being labor-intensive and time-consuming, culture methods are still the methods
54 employed most regularly in the food industry (Dwivedi and Jaykus 2011, Sohler et al. 2014).
55 However, one of the biggest limitations is that these approaches frequently detect only a

56 fraction of the microbes that are present in the sample as they rely on the isolation and growth
57 of single microbes on culture media whose metabolic and physiological requirements can be
58 reproduced *in vitro*. This not only overlooks the portion of microbes that are viable but not
59 culturable (VBNC) but also fails to consider the relationships within the community of
60 bacteria present in the sample (Cao et al. 2017). Uncultured microorganisms are estimated to
61 account for up to 99% of the microorganisms in many environments, meaning that the use of
62 traditional culture methods causes a gross underestimation of the microbial population
63 (Handelsman 2004). Although this level of underestimation may not be as considerable in
64 food systems, nevertheless, because the microbiomes of food and food processing
65 environments are composed of complex, dynamic microbial communities, meta-omic
66 approaches have the potential to provide a more accurate and greater understanding of these
67 communities.

68

69 In this review, the contribution of microorganisms to the quality and safety of food and the
70 traditional approaches to microbial characterisation are briefly described. The main focus of
71 the review is on sequencing-based meta-omic approaches and their contribution to
72 understanding microbial community dynamics in food, food-associated environments and
73 along the food processing microbiome. Other non-sequencing-based meta-omic approaches
74 are also mentioned in brief.

75

76 **IMPORTANCE OF MICROORGANISMS THROUGH THE FOOD CHAIN**

77 **Quality: Flavor, Texture, Fermentation and Spoilage**

78 Food quality is often associated with physical parameters such as pH and moisture content,
79 which can influence the growth and survival of microorganisms within a food and the food
80 chain. Food spoilage is a process or change that renders a product undesirable or
81 unacceptable for consumption, which is impacted by both the food's intrinsic characteristics
82 and the extrinsic environment (Blackburn 2006). It is a complex process, whereby food
83 undergoes biochemical changes, often due to microbial activity according to ecological
84 determinants (Nychas and Panagou 2011). Some common spoilage bacteria include
85 *Pseudomonas* spp., *Shewanella* spp., *Bacillus* spp., *Clostridium* spp., lactic acid bacteria
86 (LAB), and Enterobacteriaceae (Blackburn 2006). Ultimately, different bacteria cause
87 varying quality problems in different types of food, with some examples presented in Table 1.

88 Furthermore, despite their slower growth rate, yeasts and molds are able to exploit many
89 ecological niches in food systems and can utilise substrates and tolerate extreme conditions
90 that are not possible for bacteria (in't Veld 1996, Petruzzi et al. 2017). Common spoilage
91 yeasts include species of *Zygosaccharomyces* (in high sugar foods), *Saccharomyces* (a cause
92 of gassiness and turbidity in wines), or *Candida* (cause off-flavors in meat and dairy
93 products) and common spoilage molds include *Zygomycetes* (in produce with high water
94 content), *Penicillium* spp. (cause rot in fruits), and *Aspergillus* (in grains, spices and nuts)
95 (Sahu and Bala 2017).

96

97 However, microbes can also improve food quality by changing its intrinsic characteristics.
98 This is evident in fermented foods, where the activity of microbes can improve their
99 organoleptic and nutritive qualities in addition to extending shelf life. Fermented food
100 microbes can either be introduced spontaneously (from the raw materials or production or
101 processing environments) or inoculated as starter cultures and, over time, can produce
102 enzymes, volatile compounds, and antimicrobial molecules, such as organic acids, fatty acids,
103 hydrogen peroxide, diacetyl, and bacteriocins, which can help to slow down or prevent the
104 growth of spoilage and pathogenic microbes (Reis et al. 2012). Although the spontaneous
105 introduction of microbes is of specific relevance to this review, here we briefly provide an
106 overview of some of the most important microorganisms in fermented foods in general.

107

108 LAB are among the most important microbes in the production of several fermented foods
109 (Hatti-Kaul et al. 2018). This is reflected in the natural adaptation of many LAB to fermented
110 food environments but has been complemented by many years of research to better
111 understand and enhance their contribution to product safety as well as organoleptic,
112 nutritional, and health properties (Leroy and De Vuyst 2004). Different species of LAB have
113 been used in dairy products (cheese and fermented milks), meats (sausage), fish, vegetables
114 (sauerkraut and pickles), soy sauce, cereals (sourdough), and alcoholic beverages (wine)
115 (Leroy and De Vuyst 2004). Another group of bacteria associated with fermentation are
116 acetic acid bacteria, which mainly consist of *Acetobacter* and *Gluconoacetobacter*. This
117 group of bacteria play important roles in coffee, cocoa and vinegar fermentation because of
118 their ability to oxidize carbon substrates (Schwan and Ramos 2014). *Bacillus subtilis* and
119 *Bacillus licheniformis* are important for the industrial-scale fermentation of soybeans as they
120 grow rapidly, resulting in short fermentation times (Schallmeyer et al. 2004). Yeast can also

121 play an important role in the production of many fermented foods. *Saccharomyces cerevisiae*
122 is used in alcoholic fermentation, and yeasts are ultimately used in many indigenous
123 fermented foods as they are acid tolerant, able to grow at high temperatures, and are present
124 in many environments (Schwan and Ramos 2014). In Asia, indigenous foods fermented with
125 yeast, such as miso, soy sauce and wines, are commonly consumed (Aidoo et al. 2006).

126

127 **Safety: Pathogens and Microbial Antagonism**

128 Despite the value of fermented food and other microbes in contributing to food quality and
129 safety, with respect to food safety, microbes are frequently regarded negatively, with
130 foodborne pathogens responsible for foodborne illness and outbreaks across the globe
131 annually. The consumption of contaminated food causes an estimated 4,500 deaths annually
132 in Europe (World Health Organisation 2017). The causative agents of foodborne outbreaks in
133 Europe in 2019 were bacterial pathogens (26.4%), bacterial toxins (19.3%), viruses (10.7%),
134 and parasites and other agents (3.6%), and 40% of reported outbreaks had unknown causative
135 agents (European Food Safety Authority and European Centre for Disease Prevention and
136 Control 2019). Common pathogenic bacteria include *Bacillus cereus*, *Campylobacter jejuni*,
137 *Clostridium botulinum*, *Clostridium perfringens*, *Cronobacter sakazakii*, *Escherichia coli*,
138 *Listeria monocytogenes*, *Salmonella* spp., *Shigella* spp., *Staphylococcus aureus*, *Vibrio* spp.,
139 and *Yersinia enterocolitica*. Viruses such as norovirus and hepatitis E as well as parasites,
140 including *Toxoplasma gondii* and *Trichinella spiralis*, are also common causes of outbreaks
141 and have been recently reviewed (Bintsis 2017). There are various ways that pathogenic
142 microorganisms can enter the food chain. They can be inherent to the raw ingredients, or
143 introduced along the processing line via equipment, food handlers or packaging materials,
144 among other routes. Microbial communities can also be present in the form of biofilms,
145 which are microbial communities that adhere to solid surfaces and may contain pathogenic
146 and spoilage species that can persist on surfaces in food-processing facilities (Coughlan et al.
147 2016). Once attached, these biofilms can be difficult to remove as they are embedded in a
148 polymeric matrix and cells in the biofilm may be resistant to disinfectants or antimicrobials,
149 particularly in mixed-species biofilms (Yuan et al. 2020). Research efforts on control
150 strategies to prevent biofilm formation and remove existing biofilms are ongoing to overcome
151 this challenge in the food industry. Food safety management systems, including hazard
152 analysis and critical control points, and risk assessment principles have been widely
153 implemented to prevent foodborne illnesses and outbreaks and control the spread of

154 pathogens along the food chain. However, these management systems are reliant on having a
155 thorough understanding of the microorganisms present and the risk they may pose.

156

157 Although microbes are often viewed negatively from a food safety perspective, some have
158 been useful in biocontrol or biopreservation. Microbial antagonism has been applied in the
159 food industry through the use of bacteriocins, phages, and more (Jordan et al. 2014).

160 Bacteriocins, which consist of antibacterial peptides, have been used to target spoilage and
161 pathogenic bacteria in food and in turn prolong the shelf life and improve the safety of food
162 (Galvez et al. 2008). Bacteriocins from LAB such as nisin has been approved for use in foods
163 and is most commonly used in foods such as meat, dairy and vegetable products (Jordan et al.
164 2014). Bacteriophages or phages are virus predators of bacteria that have shown great
165 promise as they are naturally occurring and control for specific pathogenic bacteria without
166 impacting the quality and microbiota of foods (O'Sullivan et al. 2019). Phages have been
167 applied to a range of foods at various stages from farm to fork to eliminate common
168 pathogenic bacteria such as *Campylobacter jejuni*, *Salmonella* spp., *E. coli* O157:H7, and
169 more (Vikram et al. 2020). Additionally, biofilms from some species can aid in improving
170 food safety by outcompeting undesirable bacteria. Some LAB strains were found to exhibit
171 antagonistic properties against unwanted bacteria, act as a natural barrier, and alter biofilm
172 formation of spoilage microbes (Ouali et al. 2014). In food, the microbial community and the
173 interactions between microbes play essential roles in food quality and safety.

174

175 **TRADITIONAL APPROACHES TO CHARACTERISATION OF MICROBES**

176 As noted above, culture-based assays have historically been used for the detection,
177 enumeration and isolation of viable foodborne pathogens or spoilage microbes in food and
178 environmental samples (Dwivedi and Jaykus 2011). In general, samples are first
179 homogenized and then often undergo enrichment steps (pre-enrichment and selective
180 enrichment), followed by selective or differential plating to distinguish from other microbes
181 present and, finally, confirmation with biochemical, serological, or other methods. Pre-
182 enrichment is used to recover injured cells and dilute inhibitory compounds in food samples,
183 whereas selective enrichment increases the concentration of a target pathogen while
184 suppressing the growth of other microflora (Dwivedi and Jaykus 2011). These conventional
185 methods are inexpensive but time consuming and labour-intensive and can take from 2-3
186 days to a week for inoculation, isolation and confirmation depending on the targeted

187 microorganism (Mandal et al. 2011). Furthermore, owing to the possible presence of VBNC
188 bacteria, false negatives may occur, which could mean the unsuccessful detection of
189 pathogens or spoilage bacteria in food.

190

191 With the need for more timely detection of bacteria for foodborne outbreak response, rapid
192 methods that replace conventional plating steps with faster immunology or molecular-based
193 approaches have been investigated in depth and adopted more widely in recent years (Wang
194 and Salazar 2016). High levels of sensitivity and specificity are needed for food pathogen
195 detection (Feng 2007). Immunology-based methods like enzyme-linked immunosorbent
196 assay (ELISA) based on the specific binding of antigens with antibodies, have shown
197 potential but their lack of sensitivity and relatively high limit of detection [$10^3 - 10^5$ colony
198 forming units (CFUs)/mL] has meant that enrichment is generally first required before
199 detection. Nevertheless, when ELISA is coupled with nanotechnology, its application for
200 food analysis has achieved greater sensitivity, specificity and stability (Wu et al. 2019).

201

202 Compared to traditional culture methods, nucleic acid-based techniques such as polymerase
203 chain reaction (PCR), quantitative PCR (qPCR) and loop-mediated isothermal amplification
204 (LAMP) require shorter time and through multiplexing, can facilitate concurrent detection,
205 real-time monitoring and quantification of multiple microbial targets (Liu et al. 2017, Tao et
206 al. 2020). Development and optimization of each assay is important, as complex food
207 matrices may hinder nucleic acid extraction and contain inhibitors that may interfere with
208 reactions in the assay. Particularly when multiplexing, primer design is crucial, as primer sets
209 require similar annealing temperatures for a successful assay (Wang and Salazar 2016).
210 Unfortunately, these methods, like immunology-based assays, often still require enrichment
211 or concentration steps because of their limit of detection ($10^3 - 10^4$ CFUs/ mL), and as
212 pathogens are often in low concentrations in foods, direct detection is difficult (Ceuppens et
213 al. 2014, Wang and Salazar 2016). Another rapid method used largely in clinical settings is
214 matrix assisted laser desorption ionization-time-of-flight mass spectrometry (MALDI-TOF
215 MS), which analyzes signals from ribosomal proteins after ionization and time of flight
216 detection that are distinctive for each strain, allowing for rapid microbial identification (de
217 Koster and Brul 2016). Although these rapid methods are generally preferable to culture-
218 based approaches, thorough validation is required by industry and regulatory bodies before
219 routine adoption.

220

221 Although culture-based and rapid methods are useful in identifying microbes in complex
222 food- or food environment-related samples for public health and commercial purposes, they
223 create an unbalanced emphasis on specific microorganisms and, despite multiplexing, still
224 only capture a small percentage of the microbial community as a whole (Fleet 1999). As
225 microorganisms exist in communities, it is important to study them as such because the
226 growth, survival, and activity of one species or strain may impact or be associated with the
227 presence of another. Furthermore, from a practical perspective, approaches that could
228 theoretically allow the simultaneous identification of all pathogens and spoilage microbes in a
229 sample from the food chain could have a disruptive positive influence.

230

231 **META-OMIC APPROACHES: COMMUNITY APPROACHES**

232 Advances in technologies have provided the opportunity for faster and superior
233 characterisation of food chain microbiomes, with a shift toward replacing or supplementing
234 culture-dependent methods with culture-independent, molecular-based methods (Sohier et al.
235 2014). Whole-genome sequencing has successfully complemented culture-dependent
236 methods by providing deeper discrimination of microbial strains than previous typing
237 methods, with regulatory bodies in the United States and Europe now including it as a tool
238 for pathogen typing and antimicrobial resistance (AMR) surveillance (Rantsiou et al. 2018).
239 In *E. coli* O157:H7 outbreak investigations, genome sequencing stood out from other typing
240 methods, providing insights that enabled improved epidemiological case and cluster
241 identification, geographical origin tracking, and information of potential emerging strains
242 (Jenkins et al. 2019).

243

244 In contrast, culture-independent methods including the use of DNA sequencing technologies,
245 have enabled identification and characterisation of multiple microbes in foods or along the
246 food chain at the same time while also bypassing the need to culture microbes (Cocolin and
247 Ercolini 2015). Some current community-based approaches are shown in Figure 1.

248

249 **Sequencing-Based Meta-Omic Approaches: Metagenetics, Metagenomics and** 250 **Metatranscriptomics**

251 Metagenetics, also known as amplicon sequencing, metataxonomics, metabarcoding and
252 sometimes, 16S metagenomics or 16S rRNA gene sequencing, is a targeted approach that

253 involves the amplification of marker genes from mixed genomic DNA by PCR, followed by
254 direct sequencing and alignment against a reference database to identify the taxonomic
255 composition of whole microbial communities (Franzosa et al. 2015). The 16S ribosomal
256 RNA (rRNA) gene is most frequently used in the identification of bacteria, as it is universally
257 found in bacteria, and the gene contains nine hypervariable regions, some or all of which can
258 be targeted through amplification and sequencing to identify the corresponding bacterial
259 taxonomy. A similar approach can be applied to fungi, through targeting the 18S or 23S
260 rRNA genes or the internal transcribed spacer (ITS) regions of the rRNA operon.

261

262 Shotgun metagenomics, commonly referred to as metagenomics, is the untargeted genomic
263 analysis of a population of microorganisms by sequencing the entire DNA sample extracted
264 from a mixed microbial community (Quince et al. 2017). This method involves fragmentation
265 of the sample DNA, followed by preparation of a library that is sequenced, with the resulting
266 data analysed to provide information on both taxonomic composition and functional potential
267 of the entire microbial community. Due to the untargeted nature of metagenomics,
268 information relating to all categories of microbes, including bacteria, viruses, archaea, and
269 single-celled eukaryotes like fungi, can be derived from the sample (Quince et al. 2017),
270 assuming the DNA extraction method is appropriate. The lack of an amplification step
271 removes the bias that metagenetics may have and has greater sensitivity, enabling taxonomic
272 classification up to the strain level. Another advantage of shotgun metagenomics is the
273 potential for the recovery, if sufficient sequencing depth is applied, of metagenome-
274 assembled genomes (MAGs), which can provide more genomic information, revealing
275 functional and safety-related properties of specific taxa (Bowers et al. 2017), and allow for
276 the investigation of strain-level diversity in food-related microbial species such as LAB
277 (Pasolli et al. 2020).

278

279 Metatranscriptomics relates to the untargeted sequencing of total mRNA isolated from a
280 sample, which allows for the identification of transcriptionally active microbes in the sample,
281 and may provide further insights into the potential functional characteristics of the microbial
282 community. This approach reveals the microbes that are viable and, indeed, most active
283 within a community while also enabling a deeper understanding of how microbial
284 communities in complex food microbiomes or food-related environments interact with each
285 other. This approach can also be used to look at *in situ* gene expression in food, collecting
286 information on the metabolic activities potentially related to food fermentation and/or

287 spoilage that are currently expressed in a food ecosystem. An additional advantage of
288 metatranscriptomics is the ability to detect RNA-based viruses, including foodborne
289 pathogens such as norovirus (Lewis et al. 2020).

290

291 **Other Meta-Omic Approaches: Metaproteomics and Meta-metabolomics**

292 Other non-sequencing community-based methods, i.e., metaproteomics and meta-
293 metabolomics, also have the potential to be used in food microbiome studies. Metaproteomics
294 is the large-scale study of the entire protein complement produced by microbial communities
295 within a sample at a given time point, which can aid in linking genomic and transcriptomic
296 data to biological function, deepening the understanding of phenotypic changes as conditions
297 change (Soggiu et al. 2016). Metaproteomics provides information on the microbial
298 communities and their abundances and functions, the interactions within the community, the
299 changes in community metabolism and physiology, and the substrate utilization, carbon
300 sources and assimilation pathways of the microbes in the sample (Kleiner 2019). Mass
301 spectrometry is used for metaproteomics and its application in the food microbiome has
302 mainly been in the characterisation of fermented foods such as fermented soybean and cheese
303 (Soggiu et al. 2016, Xie et al. 2019). Meta-metabolomics, however, involves the use of
304 chemistry, biochemistry, and bioinformatics to detect and analyse small weight metabolites in
305 samples and provide insights into microbial phenotypic characteristics. There are two
306 categories of meta-metabolomic analyses: untargeted, which focuses on the detection of as
307 many groups of metabolites as possible, and targeted, which focuses on a specific user-
308 selected group of metabolites under determined conditions (Li et al. 2020). Researchers in
309 this field typically use either mass spectrometry or nuclear magnetic resonance to evaluate
310 food ingredients, quality, safety, authenticity, and traceability (Kim et al. 2016). The
311 integration of metaproteomics and meta-metabolomics with other omic approaches has been
312 used to provide novel insights and link genomic information with phenotypes (Kim et al.
313 2016, Pinu et al. 2019).

314

315 **Current Challenges in Sequencing-Based Meta-Omic Approaches**

316 Despite their promise, these sequencing-based approaches have challenges to overcome to
317 achieve wider application. The major challenge for these meta-omic approaches is the lack of
318 standardization, causing variation in results because of the use of different extraction
319 methods, sequencing platforms, databases, and bioinformatics tools. To highlight this point,

320 we refer to several studies that have found differing conclusions depending on the approaches
321 taken for analysis, thereby highlighting the need to identify those that provide the greatest
322 accuracy and, ultimately, their use in a standardized manner (Lewis et al. 2020, McHugh et
323 al. 2021, Walsh et al. 2018, Yang et al. 2020).

324

325 Similarly in terms of analysis, the fact that results are typically presented in terms of relative
326 abundances may lead to misinterpretations, as an increase in the relative abundance of one
327 taxon results in the concurrent decrease of others. For this reason, it is necessary to quantify
328 microbial communities by using complementary methods and efforts to do so have included
329 digital PCR, qPCR, flow cytometry and culture. It is notable that the use of synthetic
330 standards in sequencing has produced varying results for different methods, again
331 highlighting a need for caution during analysis (Galazzo et al. 2020).

332

333 Furthermore, for metagenetics, there is a difficulty when endeavoring to compare outcomes
334 across different studies arising because of a lack of consistency relating to the hypervariable
335 region of the 16S rRNA gene targeted (Claesson et al. 2010). Additionally, the focus on one
336 marker gene can cause other issues. In particular, the operon copy number for 16S rRNA
337 genes differs across taxa, which may inadvertently affect quantitative estimation. Single-copy
338 target genes like *recA*, *rpoB*, and *gyrB* have been suggested as alternatives, but their use is
339 limited because of their relevance to specific taxa of bacteria only and/or the absence of
340 databases that are sufficiently populated (Ogier et al. 2019, Poirier et al. 2018)

341

342 From the perspective of methodology, extracting DNA/RNA of sufficient concentration and
343 quality is essential for sequencing, which may be challenging in some circumstances, such as
344 from environmental swab samples taken from areas with a low microbial load (De Filippis et
345 al. 2020). Amplification methods such as multiple displacement amplification (MDA) have
346 been applied to generate DNA of sufficient quantities and the inclusion of controls have been
347 investigated to reduce contamination, but these methods can lead to biases (Marine et al.
348 2014, McHugh et al. 2021).

349

350 For both standard metagenetic and metagenomic sequencing, there is no differentiation
351 between DNA extracted from living and dead organisms within a microbiome, which is of
352 key importance with respect to food or food environment microbiome studies. Propidium
353 monoazide (PMA) and the previously more commonly used ethidium monoazide (EMA) are

354 DNA-binding dyes that selectively bind to accessible DNA present in the matrix, essentially
355 binding to DNA from dead bacteria and other cells and preventing its amplification during
356 library preparation (Nocker et al. 2006). Treatment with PMA before DNA extraction thereby
357 selects for the subsequent sequencing of DNA from viable cells. The successful application
358 of PMA with sequencing allowed the selective analysis of viable cells during milk processing
359 and cheese manufacturing (Erkus et al. 2016, Kable et al. 2019). However, its performance
360 can be influenced by the microbial community and sample biomass (Wang et al. 2021).
361 Research on the application of these dyes with the use of internal standards could provide
362 insights that may allow for quantification of live and dead cells, and optimization of this
363 treatment in different food or environment matrices. Another option is the sequencing of
364 RNA in place of or alongside DNA, as measuring RNA copies targets the active microbial
365 fraction, which allows the differentiation of viable and non-viable microbes (Mira Miralles et
366 al. 2019), although the instability of mRNA can again provide challenges. Further research on
367 the discrimination of live and dead cells is required, particularly for the application of these
368 sequencing-based approaches for food-related samples.

369

370 It is also important to note that for metagenetics in particular, the short reads generated by
371 some sequencing platforms, such as those developed by the market leader, Illumina, can be
372 limiting with respect to assigning taxonomy at the species level. Other sequencing platforms
373 that produce longer reads, such as those from Oxford Nanopore Technologies (ONT) or
374 Pacific Biosciences, may address this but lower read accuracy and higher sequencing costs
375 can be issues.

376

377 Although shotgun metagenomic sequencing overcomes many of the biases associated within
378 amplicon-based approaches, one of its biggest challenge is the reduced microbial sequencing
379 depth that occurs when randomly sequencing samples that contain high amounts of host
380 DNA. Although most studies remove the reads from host DNA during bioinformatic analysis,
381 a more efficient alternative is to deplete host DNA or enrich microbial DNA through various
382 chemical methods and commercially available kits (Marotz et al. 2018, Yap et al. 2020).
383 Similarly in metatranscriptomics, highly abundant rRNA can result in increasing costs and
384 complex downstream analysis. To overcome this challenge, rRNA depletion or mRNA
385 enrichment strategies before sequencing and/or post-sequencing removal during downstream
386 analysis have been adopted (Shakya et al. 2019).

387

388 Additionally, with the use of sequencing technologies, advanced computational power and
389 bioinformatics skills are necessary for their use, which add to the challenges when
390 considering the application of these approaches.

391

392 **APPLICATIONS OF SEQUENCING-BASED META-OMIC APPROACHES ALONG** 393 **THE FOOD CHAIN**

394 **Public Health Applications**

395 The sequencing-based meta-omic approaches mentioned above have contributed significantly
396 to the study of various diverse microbiomes, including by facilitating significant advances in
397 food microbiome research. From a public health perspective, these meta-omic approaches
398 have provided insights relating to pathogen detection, outbreak investigation, AMR
399 determination and food authenticity and source tracking.

400

401 **Pathogen detection and outbreak investigation.**

402 Meta-omic approaches are advantageous, as they bypass the need for culturing and
403 enrichment of pathogens from samples before identification and characterisation of putative
404 etiological agents. They also are able to reveal the presence of uncultured and hard to culture
405 microbes, which may be useful in surveillance, source attribution, risk assessment and
406 epidemiological analysis when traditional methods fall short (EFSA Panel on Biological
407 Hazards et al. 2019). Both metagenetic and metagenomic approaches have enabled the
408 detection and characterisation of pathogens in various foods, including vegetables, meat, and
409 dairy products (Aw et al. 2016, McHugh et al. 2018, Mira Miralles et al. 2019, Yang et al.
410 2016). Metatranscriptomics, although less widely applied because of the challenges in RNA
411 isolation, also has great potential for identifying viable pathogens in food (Yang et al. 2020).

412

413 Use of metagenomic approaches can extend beyond the food chain, where metagenomic
414 sequencing of patient stool samples collected during the outbreak in Germany of STEC
415 (Shiga toxin-producing *E. coli*) O104:H4 assisted the recovery of genomes of the outbreak
416 strain (Loman et al. 2013). Moreover, metagenomics is useful when a viral agent is the cause
417 of the outbreak or, in the case of multi strain outbreaks, it is able to discriminate and
418 characterise several strains, allowing them to be distinguished considerably faster than
419 traditional culture-based methods (Buytaers et al. 2020). Compared to metagenetics, which

420 may be more useful for low biomass samples because of the amplification of the target,
421 metagenomics facilitates more sensitive characterisation to the species level and further
422 investigation of the functional potential of microbes present (Grützke et al. 2019).

423

424 Despite the potential of these meta-omic approaches, they are currently not widely used. One
425 reason is the lack of harmonized methods and standardized, accredited workflows/pipelines
426 that would allow consistent detection and characterisation of outbreak-causing agents (EFSA
427 Panel on Biological Hazards et al. 2019). However, the usefulness of metagenomic analyses
428 can be enhanced when they are complemented with further quantitative molecular assays,
429 highlighting their effectiveness in determining pathogen contamination or outbreak events. A
430 big technical challenge that hinders greater adoption of meta-omic techniques as a routine
431 screening tool for pathogens is that these techniques are not always sufficiently sensitive
432 (Leonard et al. 2015, Lewis et al. 2020). With low numbers of pathogenic cells in samples,
433 substantial sequencing depth is required, particularly for shotgun sequencing, as samples
434 contain DNA from other microbes or contaminants such as animal, plant or human DNA
435 (Yang et al. 2016). With sufficient sequencing depth, shotgun metagenomics can be a faster
436 and more valuable tool that provides more information than current conventional workflows,
437 which permit linking food/environment outbreak-related samples with clinical samples
438 (Buytaers et al. 2020, Grützke et al. 2019, Li et al. 2020). Although the complexity of various
439 food matrices can be a challenge, this is not as great an issue for less biologically complex
440 matrices, such as water used in food production or some minimally processed foods
441 (Fernandez-Cassi et al. 2017).

442

443 **Identification of antimicrobial resistance-encoding genes.**

444 Over the past decades, AMR has been identified as a serious public health threat and because
445 of this, more tools have been published for the detection of genetic determinants of AMR
446 from sequencing data. Although whole-genome sequencing of cultured isolates is usually
447 utilized, metagenomic sequencing shows great potential for monitoring AMR, as it has out-
448 performed culture-based methods in quantifying resistance in swine herds (Munk et al. 2017).
449 Shotgun sequencing has shown success in the monitoring of AMR genes in the environment
450 from farm to slaughter (Noyes et al. 2016, Pitta et al. 2016). It has also been used to
451 understand the association between antimicrobial use and resistance and the effect of
452 processing on the resistome and virulome (Campos Calero et al. 2018, Mencía-Ares et al.
453 2020, Van Gompel et al. 2019).

454

455 As with other metagenomic approaches, sequencing depth and the presence of host DNA
456 should be considered, as they have been found to affect resistome profiling in environmental
457 and food samples (Gweon et al. 2019, Rubiola et al. 2020). Other challenges include the
458 difficulty in assigning ARG to their host species or strains, which may be addressed by
459 sequencing with long-read technology and the choice of reference resistance gene database,
460 where differences were found between gene variants from the same reference sequence from
461 different databases, reiterating the need for comprehensive databases and standardized
462 workflows (Doyle et al. 2020, Slizovskiy et al. 2020). It is also important to note that the
463 AMR data may not always be phenotypically relevant, as these genes might not be expressed
464 or the choice of bioinformatic tools can result in false positives or negatives (Doyle et al.
465 2020). From the perspective of gene expression, metatranscriptomics can potentially be
466 employed to complement the analysis (Wang et al. 2020). The analysis of the mobilome (all
467 mobile genetic elements of the microbiome) has also been paired with resistome analysis to
468 understand the potential spread of AMR genes and virulence factors through horizontal gene
469 transfer (Slizovskiy et al. 2020).

470

471 **Food authenticity.**

472 Food fraud is a global issue that has many consequences, including possible health risks,
473 economic losses, and hindering sustainability efforts. Metabarcoding has been used to
474 determine the authenticity and origin of honey, traditional Chinese medicines, fish, and more
475 (Carvalho et al. 2017, Coghlan et al. 2012, Khansaritoreh et al. 2020, Liu et al. 2020). The
476 basic concept is that the microbiome associated with a traditional food is closely linked to the
477 geographical origin and mode of production of the food as the microbes are typical of raw
478 materials and environment. Although there have been some successes, there are challenges
479 associated with using microbiomes as a means of determining the provenance of food. These
480 include the need for the existence of databases containing the components of the expected
481 microbiome of the food and the potential alteration of the microbiome due to storage or
482 processing conditions (Liu et al. 2020). Similar to other meta-omic applications, the reliance
483 on the completeness of reference databases together with the accuracy of food matrix
484 authentication are important to avoid inaccurate conclusions. Haiminen et al. (2019) found
485 both DNA and RNA shotgun sequencing to be accurate untargeted methods for food
486 authentication and contaminant detection, which has been applied by Kamilari et al. (2019) to

487 characterise Protected Designation of Origin (PDO) cheeses with complementary
488 metabolomics to define product origin differentiating factors.

489

490 **Other public health-related fields.**

491 Besides the food industry, other fields have also found benefits in the application of
492 community-based microbiome analysis methods. Community-based approaches have
493 contributed to the increasing knowledge of the indigenous microbial community and AMR
494 patterns in both healthcare settings and water systems that have provided evidence for the
495 greater need for surveillance (King et al. 2016, O'Hara et al. 2017, Zhang et al. 2017). In
496 hospital settings, meta-omic approaches have provided clues to the routes of entry and
497 relationships between pathogens and non-pathogens, as well as helped in environmental
498 surveillance to fight hospital-acquired infections and AMR (Comar et al. 2019, Rampelotto et
499 al. 2019). Similarly, when supplemented with other techniques, shotgun metagenomics was
500 effective in uncovering the presence of virulence factors and novel biomarkers of pathogen-
501 related species in drinking water distribution systems (Zhang et al. 2017). Additionally, on an
502 international scale, urban sewage and waste from aircraft flights have been cited as
503 economically and ethically acceptable approaches for continuous global surveillance and
504 prediction of AMR using metagenomics (Hendriksen et al. 2019, Petersen et al. 2015).

505

506 **Food Industry Applications**

507 Microbial communities exist throughout the food chain and understanding their dynamics and
508 the conditions that promote or hinder their growth would be useful for food safety and quality
509 purposes. Research efforts using meta-omic approaches have looked into foods, food-
510 associated environments, and food-processing steps, as presented in Figure 2, which are
511 elaborated in the following sections.

512

513 **Foods: fermented and non-fermented.**

514 One of the main applications of community-based approaches is in the study of fermented
515 foods. Previous reviews noted that most of the early studies on fermented foods employed
516 metagenetics to monitor the activity of microorganisms during fermentation (De Filippis et
517 al. 2017). In recent years, more studies have utilized metagenomics and metatranscriptomics
518 to understand the changes in microbial community diversity and activity during fermentation
519 in a broad range of foods, including vegetables, cheeses, and more (De Filippis et al. 2016,

520 Duru et al. 2018, Jung et al. 2013, Kim et al. 2020, Liu et al. 2020, Pham et al. 2019, Xiao et
521 al. 2020). Metatranscriptomic analysis revealed the changes in gene expression and metabolic
522 properties of LAB during fermentation of vegetables (Jung et al. 2013, Xiao et al. 2020).
523 Likewise from metatranscriptomic analysis of cheese, metabolic interactions within the
524 microbial community, and temperature-driven functional changes during ripening were
525 revealed (De Filippis et al. 2016, Pham et al. 2019). The use of both metagenomic and
526 metatranscriptomic analyses allowed for the detection of active microbes during fermentation
527 and of microbes responsible for biogenic amine production in fermented soy products (Kim
528 et al. 2020, Liu et al. 2020). This parallel approach was also useful in understanding the
529 dynamics of the microbial community during ripening, revealing the impact of temperature
530 on the microbial community and genes expressed (Dugat-Bony et al. 2015, Duru et al. 2018).
531 These are selected examples of studies within the continuously growing pool of research that
532 employ these methods to study the microbial consortia in fermented foods. Unsurprisingly, it
533 has been suggested that multiple meta-omic approaches facilitate the improved, efficient, and
534 sustainable production of fermented foods through detailed functional characterisation of
535 their microbiomes (Chen et al. 2017).

536

537 Although the number of studies using meta-omic approaches to study non-fermented foods is
538 considerably lower than that of fermented foods, those that have been completed highlight the
539 great potential of such approaches. Most of these applications have related to the
540 characterisation of food-associated environments or food-processing steps, which are
541 elaborated in the following sections. Other than those studies, there have been promising
542 studies involving the use of community-based methods to screen for spoilage or pathogenic
543 microorganisms. However, because of the complex nature of food samples and the frequently
544 low pathogen abundances, direct sequencing of DNA or RNA of food has, to date, been
545 found to be less sensitive than conventional culture-based or amplicon-based methods (Lewis
546 et al. 2020, Yang et al. 2020). It should also be noted that, even though both short and long-
547 read sequencing technologies have shown promise with respect to accurate classification of
548 microbes to the family and genus levels, not all approaches sufficiently classify to the species
549 or strain level needed for pathogen detection (Grützke et al. 2019). This is sometimes a
550 significant limitation, especially in terms of food safety, where identifying at only the genus
551 level may not be informative enough to understand the actual species present that could cause
552 food safety or quality issues along the food chain. The need for sensitive and specific tests
553 coupled with other challenges prove that these community-based approaches are currently not

554 applicable at the regulatory compliance level, but with further development that will be the
555 standard be in the future (Yang et al. 2016).

556

557 **Food-associated environments.**

558 Food-associated environments, from farm to processing facility, have repeatedly been found
559 to impact, both positively and negatively, the final product microbiome.

560

561 *Environmental factors.*

562 Microorganisms can enter the food chain at a number of different points. This includes the
563 crops and animals from which the foods are sourced/derived as well as environmental factors
564 such as soil, water, farming systems, pests, and climate conditions. Meta-omic approaches
565 have found that factors such as pasture systems, animal housing, airborne dust, irrigation
566 water, and several others can influence the microbiota diversity and composition of food
567 (Allard et al. 2019, Doyle et al. 2017, Wu et al. 2019). Besides diversity and composition,
568 meta-omic approaches used to characterise the resistome reveal that farm environments are
569 potential vehicles for AMR bacteria and genes, originating from dust and animal feces that
570 contribute to AMR spread and worker exposure (Luiken et al. 2020, Noyes et al. 2016). The
571 use of animal waste as fertilizer (manure/wastewater) can also cause the dissemination of
572 AMR bacteria and genes in the environment, which in turn affect the microbiota of crops
573 grown or animals raised on the land (Allard et al. 2019, He et al. 2019). Seasonality is
574 another contributing factor to the microbiota of the animal and plant environment. Seasonal
575 impacts were evident in certain products, like milk and beef, where the use of metagenetics
576 and metagenomics has revealed seasonal variations in the microbiota of final products
577 (Hwang et al. 2020, Kable et al. 2016, McHugh et al. 2020).

578

579 *Food-processing environments.*

580 Meta-omic techniques have been adopted in the characterisation of several environments
581 involved in the processing of foods such as meat (De Filippis et al. 2013, Hultman et al.
582 2015, Stellato et al. 2016), dairy (Anvarian et al. 2016, Doyle et al. 2017, Kable et al. 2016),
583 and alcoholic beverages (Bokulich et al. 2015, Bokulich et al. 2013, Wang et al. 2018). One
584 key observation from using meta-omic approaches for such studies is the presence of a
585 resident microbiome that persists within the processing environments and has the potential to
586 affect final food product quality and safety. This was highlighted in a recent review relating

587 to the use of high-throughput sequencing to characterise the dominant taxa found in both
588 processing environments and food products, which summarized the evidence that the
589 processing environment can act as a reservoir and source of microbial transfer to food (De
590 Filippis et al. 2020). This can be both beneficial and detrimental, with, for example,
591 beneficial effects apparent in fermented food production. In this regard, microbes in the
592 environment were found to contribute positively to the production of fermented vegetables,
593 wine, and Chinese liquor (Bokulich et al. 2013, Einson et al. 2018, Wang et al. 2018). In
594 contrast, spoilage or pathogenic microorganisms have been found on surfaces of various
595 dairy-, meat- and vegetable-processing facilities using different meta-omic approaches
596 (Hultman et al. 2015, McHugh et al. 2020, Pothakos et al. 2015, Stellato et al. 2016, Zwirzitz
597 et al. 2020). For example, *Pseudomonas* spp. was found in drain biofilms in cheese- and
598 salmon-processing plants (Dzieciol et al. 2016, Langsrud et al. 2016) and pathogens like
599 *Staphylococcus* and *Yersinia* were found on surfaces in milk- and meat-processing plants
600 (Hultman et al. 2015, Kable et al. 2019). Indeed, correlation of microbial communities in
601 biofilms, as determined by metagenetics, with environmental factors has been used to track
602 persistence over time, showing that bacterial communities were location-specific in meat- and
603 fish-processing plants (Rodríguez-López et al. 2020). Additionally, microbial co-occurrences
604 of pathogens with other microbes and microbial interactions within complex ecosystems can
605 be evaluated through meta-omic approaches, which may determine patterns that favour or
606 prevent the growth or survival of foodborne pathogens (den Besten et al. 2018, Illegghems et
607 al. 2015). This was investigated through 16S rRNA sequencing that examined interactions
608 between *Listeria* spp. and the microbiome within a food production facility, and identified
609 species that acted as apparent protagonists or antagonists that had impacts on the presence of
610 *L. monocytogenes* within the processing plant (Fox et al. 2014).

611

612 Handling can be a potential source of contamination or microbial transfer, whereby microbes
613 can be unknowingly transferred from surfaces to the food product. *Moraxella* spp., a
614 prominent meat-spoilage bacteria, was found on gloves of employees, which were identified
615 as a potential source of contamination using full-length 16S rRNA gene sequencing
616 throughout a pork-processing plant (Zwirzitz et al. 2020). Similarly, handling was identified
617 as a catalyst in the proliferation of spoilage bacteria in beef products after high-throughput
618 sequencing uncovered the origin of spoilage-associated bacteria from carcasses and their
619 persistence in the environment (De Filippis et al. 2013).

620

621 **Food-processing steps.**

622 Using meta-omic approaches to monitor the changes in food microbiomes during food
623 processing has been useful in understanding the impact of processes on the quality and safety
624 of foods. This has been studied through two approaches. One approach has involved profiling
625 the entire food-processing chain, where samples were taken from the start to the end of the
626 process and meta-omic methods were used to track the changes in microbial community
627 dynamics, which can facilitate the generation of mitigation measures. This whole-chain
628 approach often involves sampling of both food and environmental samples and has
629 highlighted areas where contamination or spoilage can potentially occur; e.g., in meat
630 processing, animal carcasses or hides were identified as possible sources of contamination
631 and measures taken during and after slaughter were found to be key in reducing bacterial load
632 and transmission of AMR genes to meat products (Calero et al. 2020, De Filippis et al. 2013,
633 Noyes et al. 2016, Yang et al. 2016). A similar approach to studying sausage production
634 showed that the emulsification step selected for gram-positive spoilage bacteria (Hultman et
635 al. 2015). Other investigations have highlighted the impact of storage, low temperatures, and
636 equipment on the milk microbiota in dairy processing (Falardeau et al. 2019, Kable et al.
637 2016, McHugh et al. 2020), whereas in breweries, food contact surfaces were noted as areas
638 that could allow transmission of spoilage bacteria or genes (Bokulich et al. 2015).

639

640 The other approach that has been taken when using meta-omic methodologies is process
641 focused, where specific processing steps that are often considered critical points in food
642 safety management systems are examined. Processes such as heat treatment, cold storage,
643 packaging, cleaning, and others have been studied to understand the microbial dynamics
644 during these processes and ensure their efficacy at eliminating or reducing growth of bacteria.
645 Metagenetics used to investigate heat treatments unsurprisingly found a reduction of bacterial
646 abundance and diversity in meatballs and cheese but it also affected the quality of the final
647 products (Kamilari et al. 2020, Li et al. 2021). Similarly, monitoring the ripening processes of
648 cheese using metagenomics and metatranscriptomics has provided a better understanding of
649 the temperature-driven differences in flavor development (De Filippis et al. 2016, Duru et al.
650 2018), whereas metagenetics, proteomics and complementary physicochemical and sensory
651 analysis revealed the efficacy of high-pressure processing in improving the quality and shelf
652 life of fish fillets and led to the identification of quality markers for further study (Tsironi et
653 al. 2019). For storage in particular, metagenetic and metagenomic analysis revealed cold
654 temperature storage is an area along the processing chain that allowed for the proliferation

655 and dominance of certain psychrotrophic spoilage microorganisms in meat and dairy
656 (McHugh et al. 2020, Stellato et al. 2016). Monitoring microbial dynamics to understand the
657 effect of storage temperature on the microbial community has been performed using
658 metagenetics coupled with sensory assessment or culture-dependent methods in sausage and
659 fish, which has resulted in the development of models to infer spoilage dynamics and
660 associations of bacterial species during storage (Benson et al. 2014, Zotta et al. 2019).
661 Modified atmosphere packaging (MAP) is currently used to extend the shelf life of various
662 foods like fresh and processed meat and seafood, fruit and vegetables, but optimization of the
663 gas composition is required to keep the product's quality. In the evaluation of MAP for
664 poultry, Wang et al. (2017) identified a shift in the bacterial community compared to other
665 packaging conditions using metagenomics, and Höll et al. (2020) used metatranscriptomics to
666 monitor the regulation responses of two spoilage bacteria to different atmospheric conditions.
667 Similarly, evaluations of shelf life of fish fillets in MAP and vacuum packaging at low
668 temperatures have been performed with metagenetics and sensory analysis or metabolomics
669 to understand the dynamics of spoilage bacteria over time (Jääskeläinen et al. 2019, Sørensen
670 et al. 2020). The efficacy of cleaning and disinfection has been investigated with
671 metagenetics, with evidence of bacterial diversity and abundance altered after cleaning in
672 dairy and pig facilities (Bridier et al. 2019, Dass et al. 2018). Similarly, RNA-based 16S
673 rRNA sequencing showed current cleaning practices with ozonation were effective and
674 caused shifts in potentially active microbiota in meat-processing plants (Botta et al. 2020). In
675 contrast, sanitation in salmon-processing plants, determined by metagenetics, was found to be
676 inadequate as *Pseudomonas* spp. persisted in biofilms on conveyor belts (Langsrud et al.
677 2016).
678
679 Ultimately, sequencing-based meta-omic approaches have been found to be effective tools in
680 identifying microorganisms along the processing chain, and routine implementation can help
681 to uncover the factors that influence microbial population dynamics (McHugh et al. 2020,
682 Zwirzitz et al. 2020). The numerous studies carried out to date show that there is great
683 potential for the use of meta-omic approaches in tracking microbial communities along the
684 food chain.

685 **SUMMARY POINTS**

- 686 1. Microorganisms are important contributors to the quality and safety of a food product
687 and they exist throughout the whole food chain.
- 688 2. As microbes exist in communities, it is valuable to study them as such. Meta-omic
689 approaches bypass the need for culturing and isolating microbes and allow for the
690 greater characterisation of microbial communities.
- 691 3. Metagenetics and metagenomics are two sequencing-based meta-omic approaches
692 that are already being used in the characterisation of foods, food-associated
693 environments and food processing microbiomes. Although only a few studies have
694 used metatranscriptomics, results show potential in assessing the dynamics of viable
695 microbes along the food chain.
- 696 4. The use of sequencing-based meta-omic approaches shows promise in better
697 characterisation of microbiomes along the food chain and would allow for greater
698 understanding of the factors contributing to food safety and quality. However,
699 standardized workflows/pipelines are necessary to allow for data sharing and
700 comparability and widespread adoption at a regulatory and industry level.

701

702 **FUTURE ISSUES**

- 703 1. With increasing adoption of these meta-omic approaches to uncover the microbiome
704 of food and food-related environments, there is a great need for standardized
705 workflows/pipelines for methodology and analysis.
- 706 2. Large amounts of data are generated by sequencing. This requires good data
707 management practices and systematic metadata documentation to facilitate data
708 sharing of research outputs. Additionally, bioinformatics expertise for the analysis of
709 the data generated is currently essential to draw accurate and correct interpretations
710 from the sequencing data. Future efforts will need to focus on accurate, automated
711 analytical tools.
- 712 3. As substantial parts of the analysis require referencing available databases, the results
713 from sequencing studies are only as good as these databases. Databases are currently
714 compiled mainly from human microbiome studies, as more research has been done in
715 that field, which may result in a bias toward human-related microbes. The ongoing

716 increase in microbiome studies on food and other fields should correct this imbalance
717 to enable better characterisation of microbiomes.

- 718 4. With the further development of assays to overcome the challenges of meta-omic
719 approaches, such as host DNA depletion and the ability to distinguish viable microbes
720 in the microbial community, there will be an even wider application of meta-omic
721 approaches for the characterisation of microbes along the food-processing chain.
- 722 5. From metagenomic data, the recovery of MAGs could make way for more single-
723 strain studies that can contribute to a greater understanding of the resident microflora
724 of food environments as well as the strains responsible for fermentation or spoilage in
725 foods. Additionally, increasing the number of studies into the functional properties of
726 microorganisms within food environments using metatranscriptomics or
727 metagenomics with complementary approaches like metabolomics can provide greater
728 insight into the active microorganisms and metabolic pathways involved in processes
729 along the food chain.
- 730 6. Portable sequencing devices from ONT have allowed for field/onsite sequencing
731 which has proven to be useful in clinical outbreak investigations and environmental
732 sampling. These portable devices could enable rapid detection of microbiological
733 contaminants or pathogens in food-production or food-processing environments.
734 Although some studies have explored this possibility, further comparisons with other
735 sequencing technologies and platforms are required to determine accuracy and
736 comparability (McHugh et al. 2021, Yang et al. 2020).

737

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739 The authors are not aware of any affiliations, memberships, funding, or financial holdings
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741

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748

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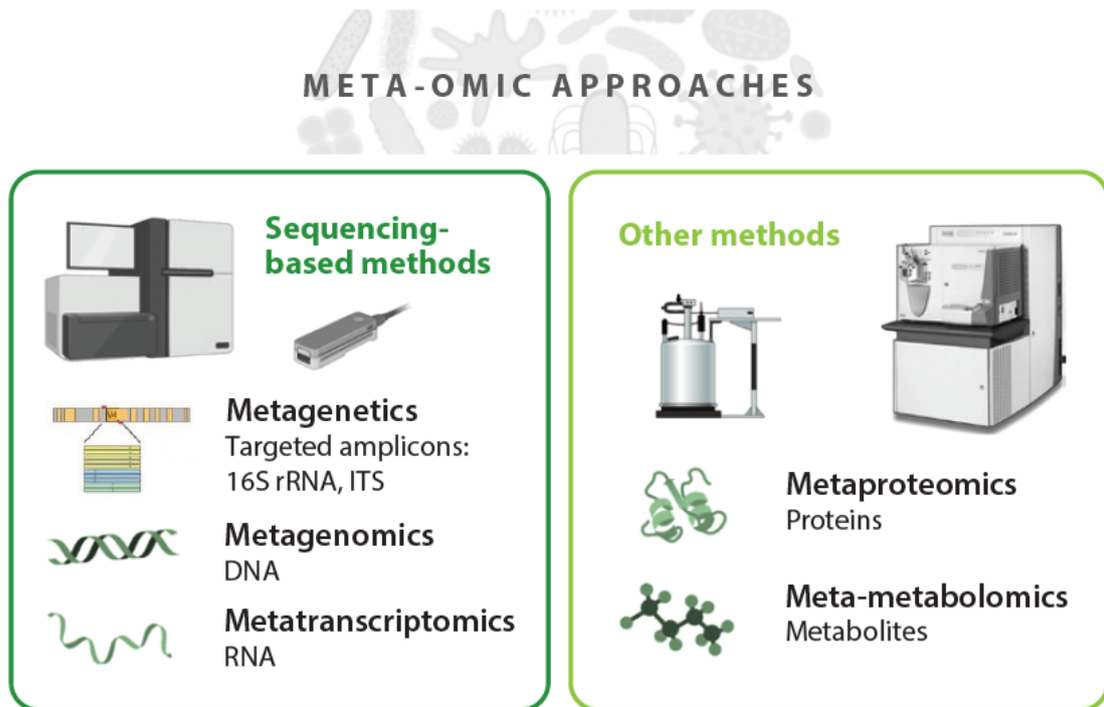
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1181 **TABLES**

1182 Table 1. Some classical examples of types of food spoilage that can be caused by different organisms.

Spoilage characteristic	Spoilage organism in food type	References
Off-odor and off-flavors	<p><i>Pseudomonas</i> spp., <i>Carnobacterium</i> spp., <i>Serratia</i> spp., <i>Leuconostoc</i> spp. and <i>Brochothrix thermosphacta</i> produce off-odors and off-flavors in meat, fish and poultry</p> <p><i>Shewanella putrefaciens</i> causes rancid and sulfurous odors and <i>Aeromonas</i> spp. produces a sour flavor in smoked salmon</p> <p>Various Enterobacteriaceae cause off-odors and off-flavors in preserved seafood products</p> <p><i>Citrobacter</i> and <i>Proteus</i> have been found to cause off-odors in poultry</p> <p><i>Candida</i> spp. and <i>Kluyveromyces</i> spp. cause off-odors and flavors in fermented dairy products.</p>	Blackburn (2006), Stohr et al. (2001), Fleet (2011)
Changes in texture	<p><i>Pseudomonas</i> spp. cause meat and poultry to become slimy/mushy due to the action of degradative enzymes</p> <p>LAB can cause poor texture in cheese</p> <p><i>Bacillus</i> spp. are able to cause ropiness in breads and bakery products</p> <p><i>Clostridium</i> spp. and <i>Bacillus</i> spp. cause softening in vegetables and fruit</p> <p><i>Erwinia</i> and <i>Penicillium</i> spp. cause soft rots in vegetables, leading to a mushy texture.</p> <p>The texture of cheese and yogurts is altered by <i>Candida</i> spp. and <i>Kluyveromyces</i> spp.</p>	Blackburn (2006), Nychas and Panagou (2011), Fleet (2011)
Discolouration	<p><i>Pseudomonas fluorescens</i> is able to cause blue coloration in cheese</p> <p><i>Carnobacterium viridans</i> causes green discoloration in cooked cured sausage</p>	Nogarol et al. (2013), Peirson et al. (2003)
Gas formation	<p><i>Clostridium</i> spp. cause gas formation resulting in bloating in canned or vacuum-packed goods and late blowing defects in cheese</p> <p>Enterobacteriaceae is responsible for gas production in salad products</p> <p><i>Saccharomyces</i> causes gassiness in wines</p> <p>Several yeast species cause swelling in juice packets</p>	Petruzzi et al. (2017), Sahu and Bala (2017)



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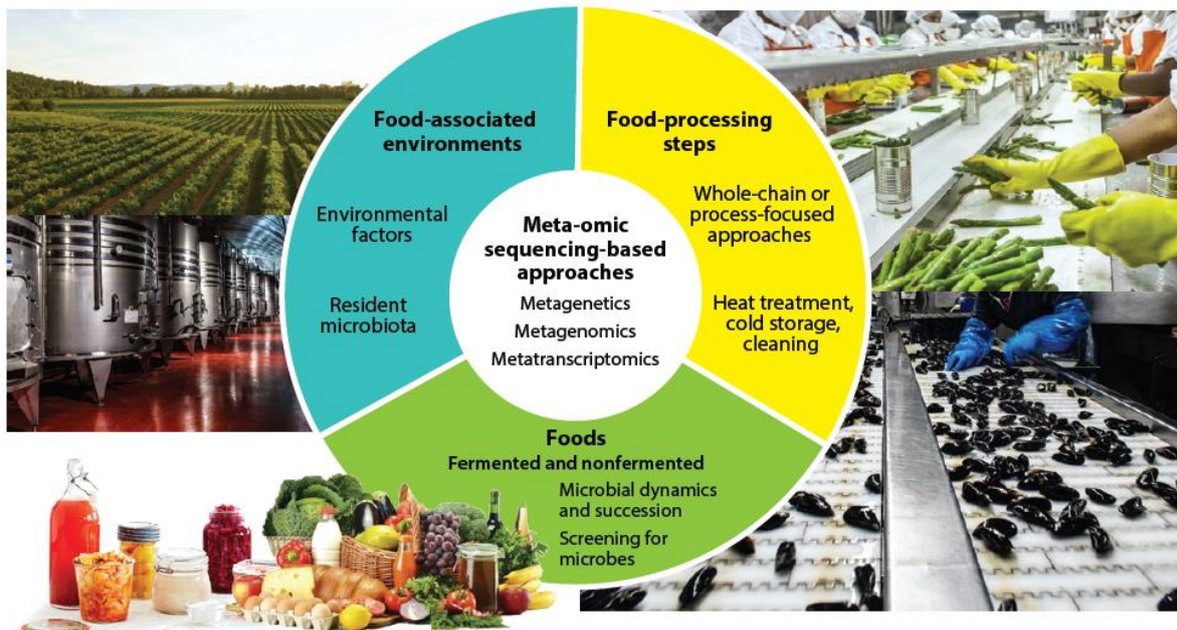
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1187 Figure 1. Current meta-omic approaches used in microbiome research. Sequencing-based
1188 approaches include Metagenetics, Metagenomics and Metatranscriptomics and other
1189 community-based methods include Metaproteomics and Meta-metabolomics, which are
1190 currently being used in human, environment, and food microbiome studies.

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CHARACTERIZATION OF MICROORGANISMS ALONG THE FOOD CHAIN



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1195 Figure 2. Current applications of meta-omic sequencing-based approaches along the food

1196 chain. Metagenetics, Metagenomics and Metatranscriptomics have been used in studies

1197 investigating the microbial community of food, food processing steps and food-associated

1198 environments.

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