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## 33 ABSTRACT

34 Expectations for the industrialization of cultured meat are growing due to the increasing 35 support from various sectors, such as the food industry, animal welfare organizations, and 36 consumers, particularly vegetarians, but the progress of industrialization is slower than 37 initially reported. This review analyzes the main issues concerning the industrialization of 38 cultured meat, examines research and media reports on the development of cultured meat to 39 date, and presents the current technology, industrialization level, and prospects for cultured 40 meat. Currently, over 30 countries have companies industrializing cultured meat, and around 200 companies that are developing or industrializing cultured meat have been surveyed 41 42 globally. By country, the United States has over 50 companies, accounting for more than 43 20% of the total. Acquiring animal cells, developing cell lines, improving cell proliferation, 44 improving the efficiency of cell differentiation and muscle production, or developing cell 45 culture media, including serum-free media, are the major research themes related to the development of cultured meat. In contrast, the development of devices, such as bioreactors, 46 47 which are crucial in enabling large-scale production, is relatively understudied, and few of 48 the many companies invested in the development of cultured meat have presented products 49 for sale other than prototypes. In addition, because most information on key technologies is 50 not publicly available, it is not possible to determine the level of technology in the 51 companies, and it is surmised that the technology of cultured meat-related startups is not 52 high. Therefore, further research and development are needed to promote the full-scale 53 industrialization of cultured meat.

54

55 Keywords: cultured meat; cultured meat industrialization; muscle satellite cell; myogenesis

## 56 Introduction

57 Cultured meat, also called in vitro meat or laboratory-cultured meat, is an edible tissue 58 produced by the isolation, proliferation, and differentiation of muscle satellite cells (MSCs) 59 obtained from a small amount of livestock tissue (Lee et al., 2021). The production of livestock products based on stem cell and tissue culture technologies is seen as a future 60 61 technology and an emerging industry that is not only resource-efficient but can effectively 62 address environmental degradation and the uncertainties associated with food security in the 63 face of a growing global population and dwindling natural resources (Risner et al., 2023). Several countries around the world have implemented or taken steps to create policies to 64 categorize cultured meat as cellular agriculture (Soice and Johnston, 2021). 65 66 A 2023 report by GlobeNewswire ascertained that the estimated value of the global 67 cultured meat market was USD 182 million in 2022 and will continue to grow, with a projected CAGR of 23.2% (Fig. 1). However, this is only 0.014% of the global traditional 68 69 meat market size of USD 1.28 trillion, which was reported in the same year (Phuong, 2023), 70 indicating that the cultured meat market is still small compared to the traditional meat market 71 (Fig. 1). Currently, the only marketable cultured meats that have been officially certified as 72 safe by the United States Food and Drug Administration (FDA) are cell-cultured chicken 73 from Upside Foods and GOOD Meat.

Following this official approval, several companies worldwide are seeking permission to
sell cultured meat. In July 2023, an Israel-based company, Aleph Farms, submitted a
regulatory approval application to the Swiss Federal Office for Food Safety and Veterinary
Medicine (Aleph Farms, 2023). Subsequently, in January 2024, Israel's Ministry of Health
(MoH) approved the sale of cultured beef from Aleph Farms, making it the third country to
offer cultured meat for sale and the first approval for a bovine species (Aleph Farms, 2024).
In October 2023, CellMEAT requested the Ministry of Food and Drug Safety (MFDS) of the

81 Republic of Korea certification for Dokdo shrimp (Lebbeus groenlandicus) cell culture as a 82 temporary food ingredient (CellMEAT, 2023). In December 2023, Food Standards Australia 83 New Zealand (FSANZ) announced new amendments to an application received from Vow 84 seeking approval of cultured quail (FSANZ, 2023). Likewise, research is underway around 85 the world to produce cultured beef, pork, lamb, turkey, foie gras, and various types of seafood 86 (oysters, lobster, shrimp, salmon, and tuna) using cell culture technology, and the 87 development of various materials and equipment for cultured meat production, including 88 adipocytes, supports, microcarriers, growth factors, and bioreactors, is gaining traction. Despite expectations, the full-scale industrialization of cultured meat has not yet been 89 90 achieved, and the timing of the industrialization of cultured meat remains unclear. In 91 addition, the terminology for cultured meat has not yet been standardized. The Food and 92 Agriculture Organization (FAO) and the World Health Organization (WHO) use the term 'cell-based food,' the United States Department of Agriculture-Food Safety and Inspection 93 94 Service (USDA-FSIS) uses 'cell-cultured meat,' and the U.S. Food and Drug Administration 95 (FDA) uses 'cultured animal cell material' (e.g., cultured Gallus gallus cell material) (FAO-96 WHO, 2023; FDA, 2023a; USDA-FSIS, 2023).

97 Therefore, this review analyzes the main issues related to the industrialization of cultured 98 meat, as well as research reports and media reports on the development of cultured meat to 99 date, with the aim to present the current technology, industrialization level, and prospects of 100 cultured meat.

101

# 102 Cultured Meat and Food Safety

103 Cultured meat production facilities are considered to be safer than conventional meat
104 production facilities against foodborne pathogens, such as *Salmonella*, *Campylobacter*,

105 Escherichia coli, yeasts, molds, and parasites because they are designed with enclosed

structures that can control the entry of external substances (Chriki and Hocquette, 2020). 107 However, potential threats from the cultured meat production process cannot be completely 108 ruled out. Among the anticipated food safety concerns are contamination with 109 microorganisms and prion proteins that may occur during the cell culture phase, residues of 110 antibiotics and cell freezing agents, the safety of cell lines (genetic manipulation and 111 excessive passage culture), exogenous recombinant growth factors, unknown allergens, and 112 the safety of support materials (Broucke et al., 2023; Ong et al., 2021). 113 Furthermore, it is crucial to adhere to the guidelines of food authorities, such as the FDA, when using a scaffold for the production of cultured meat. This includes following 114 115 regulations regarding the use of materials, solvents, cross-linking agents, inedible substances, 116 toxic compounds, allergens, and other related factors (Levi et al., 2022). However, challenges 117 remain in the commercialization of scaffolds due to the need to establish safety evaluation 118 and approval standards for solvents or cross-linking agents used in scaffold polymerization, 119 potential decomposition by-products of biodegradable scaffolds, physicochemical 120 modifications of synthetic polymer scaffolds, and recombinant proteins that improve cell 121 attachment efficiency (Bomkamp et al., 2022). In response to these concerns, several countries, such as those in Australasia and the 122 123 European Union (EU), Korea, Singapore, the United Kingdom, and the United States, have 124 taken steps toward establishing regulations and classification guidelines for cell-based foods or temporarily allowing them as food ingredients (EU, 2021; FDA, 2019; FDA, 2023b; FSA, 125 126 2023; FSANZ, 2023; MFDS, 2023; Singapore Food Agency, 2023; USDA-FSIS, 2023). 127 These regulations are overseen by national agencies in each country (**Table 1**). 128 As there are many threats to the safety of cultured meat, it is essential to establish a 129 "standard safety assessment procedure for cultured meat" that includes not only cell-cultured 130 chicken but also other major livestock species, such as beef and pork, or cell-cultured

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131	seafood, to ensure the safety certification and commercialization of cultured meat (Ong et al.,
132	2021). Furthermore, potential threats in cultured meat that are not yet well understood need to
133	be further investigated, and the safe production of cultured meat should be based on the use
134	of validated food ingredients in the product production process and appropriate regulations
135	(Bhat et al., 2015). Safe consumption of the product is a prerequisite for cultured meat to be
136	licensed and marketed as a new food, which requires standardization of the manufacturing
137	process or the development of manufacturing guidelines (Mariano et al., 2023). It may also
138	be necessary to evaluate the safety of the final product manufactured according to the
139	standardized process or manufacturing guidelines, which can be done using methods similar
140	to those used to evaluate new foods for authorization for human consumption (Lee et al.,
141	2023c). In general, the safe consumption of food is assessed by short-term and long-term
142	toxicity tests in laboratory animals.
143	Toxicity tests used to assess food for human consumption analyze the genotoxicity,
144	reproductive toxicity, hematotoxicity, hepatotoxicity, or allergenicity in comparison to
145	existing products. In assessing the safety of cultured meat for consumption, it may be
146	necessary to standardize or set guidelines for the following five processes:
147	- Cell acquisition
148	- Cell culture preparation
149	- Cell culture and muscle differentiation
150	- Cultured muscle acquisition
151	- Manufacturing meat products using cultured muscle
152	
153	Sustainability and Animal Welfare
154	According to the United Nations' World Population Prospects 2022 report, the world's

155 population is expected to reach 9.7 billion by 2050 from 8 billion in 2022, and a joint report

prepared by the Organization for Economic Co-operation and Development (OECD) and the
FAO (OECD-FAO, 2022) predicts that global meat consumption will increase by 15% by

158 2031 to keep pace with projected population growth. As a result, more land for growing feed

159 is needed to keep up with the trend of increasing meat consumption.

160 The global livestock industry has drawn increased attention in recent years because of the 161 magnitude of its environmental impact. Greenhouse gases from livestock production are

162 estimated to be 14.5% of global greenhouse gas emissions, and agricultural water use is

163 reported to be 29% of global water use, 98% of which is used for the production of animal

164 feed (Gerber et al., 2013; Mekonnen and Hoekstra, 2012). The environmental costs of

165 livestock production also include land degradation, eutrophication of lakes and rivers, lower

166 soil fertility, reduced biodiversity, increased exposure to zoonotic diseases, and accumulation

167 of livestock manure, which could contaminate surface and groundwater, and has been shown

168 to contribute to the transmission of zoonotic diseases and antibiotic-resistant bacteria

169 (Godfray et al., 2018; Morand et al., 2019; Xie et al., 2018; Young et al., 2014).

170 Cultured meat has been reported to involve 78-96% less greenhouse gas emissions, 99%
171 less land use, 82-96% less water, and 7-45% less energy use than conventional meat

172 production methods, depending on what meat product it is being compared to (Reis et al.,

173 2020; Tuomisto and Teixeira de Mattos, 2011). These data suggest that cultured meat could

be a key promotional tool to induce positive consumer perceptions of its environmental

benefits and engagement in environmental protection. Pakseresht et al. (2022) reviewed a

total of 43 articles and identified environmental and ethical concerns among eight major

177 factors determining the consumer acceptance of cultured meat. However, data quantifying the

178 climate and environmental impacts of cultured meat production is highly speculative, based

179 on forward-looking projections, and actual cultured meat production systems are often hidden

180 due to intense competition, leaving little detailed information available for analysis (Lynch

and Pierrehumbert, 2019; Tuomisto, 2019). Therefore, a systematic approach with a larger
sample size of cultured meat production technologies needs to be developed to assess the
environmental impact of cultured meat.

184 As global meat consumption is on the rise, the scale of farming and the number of animals slaughtered are expected to increase, and the religious, ethical, and environmental 185 186 controversies that arise from the slaughter process are likely to become more intense than 187 before (Heidemann et al., 2020). Over the years, continued efforts have been made to 188 improve the efficiency of the livestock industry for mass production, but equally important is prioritizing animal welfare and, accordingly, the movement to improve animal welfare, such 189 190 as developing standards for animal welfare certification and labeling schemes, is reported to 191 be increasing every year (Anomaly, 2015; Parker et al., 2017). Given that cultured meat 192 would reduce the need for raising livestock for slaughter, it can improve animal welfare 193 concerns (Hocquette, 2016), and studies of consumers suggest that the emotional benefits of 194 cultured meat in terms of animal welfare contribute to positive perceptions of cultured meat 195 (Bryant and Barnett, 2020; Lin et al., 2023; Rolland et al., 2020). Conversely, some 196 consumers have expressed concerns that cultured meat will affect the demand for industrial 197 animals, leading to a decrease in the number of live animals, which poses a potential threat to 198 traditional livestock farming, ultimately leading to a disruption of the balance between 199 animals and nature (Laestadius and Caldwell, 2015; Newton and Blaustein-Rejto, 2021). In 200 response to these concerns, scenario analysis studies have been conducted on the possibility 201 of cultured meat partially replacing traditional livestock farming, but cultured meat is still 202 considered to be at a technological plateau, requiring extensive research and large capital 203 investments to replace conventional meat production (Mateti et al., 2022; Moritz et al., 2023). 204

# 205 **Consumer Perception**

206 Cultured meat producers are emphasizing the benefits of environmental efficiency, 207 sustainability, eco-branding, and environmental costs to win over consumers and are actually 208 creating added value for cultured meat products by reducing the negative environmental 209 impact of product production and providing differentiated products that make consumers feel 210 like they are investing in environmental protection (Reis et al., 2020). In the review by 211 Pakseresht et al. (2022) mentioned above, 43 (17.7%) of 243 screened articles on cultured 212 meat development and technology concerned consumer attitudes, highlighting the scarcity of 213 studies exploring consumers responses to this technology. In a choice experiment using a 214 randomized group of 533 consumers, it was found that taste, health, price, animal welfare, 215 environmental impact, and social impact were the most important factors in determining the 216 purchase of a burger product and that only 11% of consumers would choose a burger made 217 with cultured meat if all burgers had the same price (Slade, 2018). However, when presented 218 with a positive framing of cultured meat, more than 66-70% of consumers were willing to try 219 or purchase cultured meat, and those who were willing to purchase had a favorable evaluation 220 of cultured meat, citing improvements in environmental and animal welfare as benefits of 221 cultured meat (Bryant and Barnett, 2020; Rolland et al., 2020; Wilks and Phillips, 2017; 222 Zhang et al., 2020). Furthermore, in a system dynamics model study to estimate the demand 223 for cultured meat, the price of the product had the greatest impact on the speed of promotion 224 and purchase decision-making for cultured meat, with low prices showing high demand 225 regardless of the promotion strategy, suggesting the importance of proper pricing in the 226 launch of cultured meat products (Skinner and Blake, 2023).

However, cultured meat is categorized as a novel food that is only available for purchase or tasting in limited quantities in a handful of countries, and all the consumer research published as of November 2023 is based on hypothetical product settings. Additionally, consumer response has been found to be largely consistent, and it is expected that the production of

affordable cultured meat to meet consumer satisfaction will be paramount. As alluded to
above, another consumer concern regarding the development of the cultured meat industry is
that it will negatively impact traditional livestock farmers (Wilks and Phillips, 2017). A
survey of the acceptance of alternative meat products among farmers and non-farmers found
that both farmers and non-farmers expressed concerns about the impact of cultured meat on
traditional livestock farming, with farmers reporting a lower preference for alternative meat
products than non-farmers (Crawshaw and Piazza, 2023).

238 Cultured meat also provokes ethical, cultural, and religious discussions. According to Islamic beliefs, halal means exception in Arabic, and whether cultured meat is halal is a 239 240 determining factor in Muslims' acceptance of cultured meat consumption (Hamdan et al., 241 2018). Muslims in the United Kingdom were less likely to try new foods than non-Muslims 242 due to uncertainty about halal status, but Muslims were found to be more likely to purchase 243 cultured meat than non-Muslims (Boereboom et al., 2022). Muslims in Singapore also 244 considered the safety and halal status of cultured meat before accepting it, and there was a 245 link between food safety and religious acceptance (Ho et al., 2022). To enter the kosher and 246 halal markets, cultivated meat must comply with specific standards and requirements, 247 including those related to its origin and method of production. In September 2023, Orthodox 248 Union Kosher, the world's largest and most influential kosher certification authority, certified 249 poultry products from SuperMeat as kosher, marking a major advancement for the food technology's acceptance under Jewish dietary law (Tress, 2023). At the time of writing, 250 251 Aleph Farms (the first to receive approval for cultured meat for a bovine species) is awaiting 252 a decision on kosher and halal certification of its beef steaks after seeking consultation from 253 several religious authorities.

Therefore, the strategies necessary for consumer acceptance of cultured meat must
 consider the positions of various sectors, such as government policy, food safety, traditional

256 livestock farming and cultured meat, and religious/cultural/ethical perspectives. Accurate 257 data and research are needed to compare the sustainability of the conventional meat industry 258 and cultured meat industry, not only to highlight the positive aspects of cultured meat but also 259 to consider the coexistence of traditional livestock farming and cultured meat (Bryant and 260 Barnett, 2018). However, market-based information on actual cultured meat technologies is 261 inconsistent, making it difficult to evaluate and analyze, and environmental impact analysis is 262 based on data with higher uncertainty compared to traditional livestock farming (Rodríguez Escobar et al., 2021). In addition, because most consumers' positive perception and 263 acceptance of cultured meat is based on trust in the government, it is necessary to establish 264 265 strict standards for food safety (Ho et al., 2022). 266 In conclusion, to assess the ideal sustainability of cultured meat, bridging the knowledge

and information gap is a must, and collaboration between relevant companies and researchers
is needed to integrate the entire production process and scenarios so that the environmental
impact of cultured meat can be reasonably predicted. Furthermore, the government should
take into account the proposed scenarios and establish regulations to enable consumers to
choose safe cultured meat.

272

## 273 Domestic and International Cultured Meat Companies

Information on domestic and international cultured meat companies as of 2023 is presented

in **Table 2**, with a total of 195 companies producing food-grade cultured meat-related

276 products in 35 countries. The largest number of cultured meat companies were identified in

the United States (53), followed by the United Kingdom (17), Israel (14), Singapore and

278 Canada (11), South Korea (10), Germany (9), the Netherlands and Japan (6), India, France

- and mainland China (5), South Africa, Argentina, and Australia (4), the Czech Republic (3),
- 280 Belgium, Switzerland, Spain, Austria, and Chile (2), and other countries (New Zealand,

281 Denmark, Russia, Malaysia, Mexico, Vietnam, Sweden, Iceland, Croatia, Turkey, and 282 Portugal). Furthermore, of the 307 product categories mentioned as being researched by 283 companies, the top 10 categories are Meat, Beef, Fish, Pork, Chicken, Seafood, Scaffold, 284 Culture media, Ingredients, and Others, accounting for 79.5% of the total, indicating that 285 current trends in company-level cultured meat research are centered on cultured meat (beef > 286 fish > pork > chicken = seafood), supports, and media (**Fig. 2**). However, it is necessary to be 287 cautious in identifying trends as there are many cases where the researchers do not clearly 288 mention the animal species under research and refer to it as Meat or Seafood.

289

## 290 **Production of Cultured Meat**

# 291 Muscle satellite cells (MSCs)

MSCs are muscle-derived adult stem cells that are responsible for the regenerative capacity 292 of muscle following damage to myofibers. MSCs are characterized by rapid proliferation in a 293 294 highly active state early in life, while the proportion entering a quiescent state increases with 295 age (Mesires and Doumit, 2002). Myofibrils are composed of structures surrounded by an 296 inner sarcolemma and an outer basement membrane, and the basal lamina, which is close to 297 the myofibrils, has been identified as an extracellular matrix (ECM) that is in direct contact 298 with MSCs and is involved in the maintenance of physiological functions and the 299 development of skeletal muscle (Holmberg and Durbeej, 2013; Zhang et al., 2021). The basal 300 lamina is composed mainly of type IV collagen, which plays a role in maintaining MSCs in a 301 quiescent state by sequestering various growth factors and signaling molecules involved in 302 their activation and proliferation (Kann et al., 2021; Sanes, 2003). Furthermore, quiescent 303 MSCs located in the niche between the basal lamina and myofibrils have a fusiform 304 morphology with little cytoplasm and organelles and have been shown to express MSCspecific genes, such as paired box protein 3 (Pax3) and Pax7, and myoblast determination 305

protein 1 (*MyoD*) at the beginning of quiescence or proliferation entry (Fu et al., 2015; Kuang
et al., 2006; Zhang et al., 2010).

308

### 309 *Gene expression and signaling pathways*

310 Understanding the regeneration process of MSCs is necessary for cultured meat 311 production, and the genes and signal transduction pathways that regulate proliferation and 312 differentiation that have been widely reported to date are shown in Fig. 3. Pax3 is considered 313 one of the important genes responsible for MSC survival during embryogenesis. It is also 314 purported to be involved in the formation and underlying development of early muscles by 315 affecting the expression of MyoD and myogenic factor 5 (Myf5) to regulate the development 316 of limb muscles (MyoD) and peri-spinal and intercostal muscles (Myf5) in early embryos (Kablar et al., 1997). Pax7 is an essential gene for MSC maintenance, and individuals with 317 318 Pax7 knockout show a decreased rate of muscle regeneration in muscle injury treatments and 319 difficulty in generating MSCs (Kuang et al., 2006). In addition, Pax7 has been found to act as 320 an antagonist of MyoD, resulting in an increased number of Pax7-positive cells in the 321 muscles of individuals with *MyoD* knockout (Kuang et al., 2006; Olguin and Olwin, 2004; 322 Seale et al., 2000).

323 Activation of MSCs is an early step in myogenesis. When a muscle is damaged, the 324 disruption of the basal plate and reorganization of the environment leads to interactions 325 between signaling molecules that were previously sequestered by the basal plate and MSCs, 326 leading to their activation (Li et al., 2018). Muscle formation is mainly regulated by 327 myogenic regulatory factors (MRFs) expressed in activated MSCs. Some representative 328 MRFs are MyoD, Myf5, myogenin, and muscle-specific regulatory factor 4 (MRF4, also 329 known as Myf6) (Kim et al., 2023a). Activated MSCs divide to produce satellite cell-derived myoblasts that continue to divide and proliferate before committing to differentiation and 330

331 fusing to form myotubes, which then mature into myofibers. When satellite cells are 332 activated, they initiate differential expression of MRFs depending on the asymmetry of cell 333 orientation after division (Kuang et al., 2007). Accordingly, it has been shown that if the 334 orientation of the cells formed after somatic cell division is on the myofibrillar side, they 335 upregulate origin regulatory factors, such as MyoD and Myf5, whereas cells on the basal 336 plate side do not express Myf5 and retain stemness (Kuang et al., 2007; Troy et al., 2012). 337 MyoD and Myf5 are genes that activate myogenin and MRF4 and participate in the late 338 stages of muscle formation by influencing the fusion of myoblasts and the initiation of their 339 final differentiation, leading to cell maturation and ultimately the formation of multinucleated 340 myotubes (Cornelison et al., 2000; Hawke and Garry, 2001; Punch et al., 2009; Smith et al., 341 1993). MyoD has somewhat overlapping roles with myogenin, but when myogenin is deleted, 342 MyoD is unable to take over its role, and individuals with myogenin deletion have been 343 shown to die at birth due to impaired skeletal muscle formation (Adhikari et al., 2021; 344 Nabeshima et al., 1993). It was also found that in C2C12 cultures with myogenin deletion, 345 myomaker and myomixer, two genes that regulate the fusion of skeletal muscle, were 346 significantly downregulated, leading to the inhibition of differentiation (Adhikari et al., 347 2021). MRF4 is an origin regulator that is predominantly expressed in fully differentiated 348 muscle fibers and plays a role in maintaining the MSC pool. It has been reported that deletion 349 of MRF4 can significantly reduce the number of Pax7-positive MSCs in postnatal individuals 350 (Lazure et al., 2020). 351 Signals that regulate the stemness of MSCs are known to include  $p38\alpha/\beta$  mitogen-activated

352 protein kinase (MAPK) or Notch. First, inhibition of p38 has been reported to induce self-

renewal of MSCs by blocking the MyoD expression pathway and maintaining Pax7

assume as the second se

proliferative state (Ding et al., 2018; Li et al., 2023; Troy et al., 2012). Among Notch

356 signaling components, Notch1 is activated upon muscle injury in vivo by binding to 357 myofilament ligands to induce cell cycle exit, Notch2 is activated in MSCs to maintain the 358 stemness of the MSC population by inhibiting differentiation, and Notch3 has been shown to 359 inhibit the p38 $\alpha/\beta$  MAPK pathway to suppress myocyte enhancer factor 2 (MEF2) expression 360 associated with differentiation (Conboy and Rando, 2002; Gagan et al., 2012; Jo et al., 2022). 361 It has been reported that activated MSCs are proliferation-induced and differentiation-362 inhibited by the phosphoinositide 3-kinase (PI3K)/Akt pathway or the extracellular signal-363 regulated kinase 1/2 (ERK1/2) pathway (Li et al., 2023; Mohammadabadi et al., 2021; Ohashi et al., 2015). Growth factors known to be involved in PI3K/Akt activation include 364 365 fibroblast growth factor (FGF), insulin-like growth factor (IGF)-1/2, hepatocyte growth factor 366 (HGF)/c-Met, epidermal growth factor (EGF), and interleukin-6/Janus kinase 2/signal transducer and activator of transcription 3 (IL-6/JAK2/STAT3). These factors have been 367 368 shown to act as activators of mammalian target of rapamycin complex 1 (mTORC1), which 369 can regulate the proliferation of muscle progenitor cells (Brandt et al., 2018; Holterman and 370 Rudnicki, 2005; Lu et al., 2017; Messersmith et al., 2021; Ohashi et al., 2015; Ornitz and 371 Itoh, 2015; Relaix et al., 2021; Rhoads et al., 2016; Wang et al., 2023a). Furthermore, it has 372 been confirmed that EGF and FGF are involved in the ERK1/2 pathway, one of the MAPK 373 family signaling pathways, which can activate myoblast proliferation and impair the initiation 374 and maintenance of differentiation (Li et al., 2023; Mohammadabadi et al., 2021; Ohashi et al., 2015). Additionally, the Wnt pathway can activate both mTORC1/2, with mTORC1 375 376 regulating metabolism in response to environmental factors (growth factors, amino acids, 377 energy, and stress) and mTORC2 involved in the maintenance of MSC populations through 378 phosphatase family pathways (Oh and Jacinto, 2011; Rion et al., 2019; Wei et al., 2019). 379 The p38 $\alpha/\beta$  MAPK pathway activates MEF2 and plays a major role in the differentiation of myoblasts. Myotube formation is inhibited when MEF2 is removed because of the 380

381	involvement of MEF2 in the proliferation and differentiation of MSCs (Chen et al., 2017;
382	Shao et al., 2022; Wang et al., 2018). Furthermore, when the mTOR pathway is inhibited by
383	rapamycin in MSC cultures, the expression of myogenic genes (Pax7, Myf5, and MyoG) is
384	inhibited, indicating that the mTOR pathway is essential for the proliferation and
385	differentiation of MSCs (Zhang et al., 2015). In addition, previous studies on MSC
386	differentiation have shown that Wnt1 and Wnt7a signaling, along with activation of the
387	Wnt/ $\beta$ -catenin pathway, increases $\beta$ -catenin to induce myogenic differentiation of
388	mesenchymal stem cells and activate the myogenic regulators Myf5 and MyoD to influence
389	skeletal muscle development (Eng et al., 2013; Zhu et al., 2022). Signals that inhibit
390	differentiation include ERK, myostatin, and protein kinase A (PKA), with myostatin reported
391	to inhibit muscle formation by co-inhibiting the Akt pathway and PKA reported to induce
392	proteolytic cleavage to produce factors that inhibit MEF2 signaling (Backs et al., 2011;
393	Mohammadabadi et al., 2021; Trendelenburg et al., 2009).
394	The nuclear factor of activated T-cells (NFAT) can activate signaling molecules that
395	regulate the fusion of myoblasts and myotubes, such as MEF2 and IL-4, by the calcineurin
396	and p38/MAPK pathways; however, PKA has been reported to prevent premature
397	differentiation of myoblasts by rephosphorylating MEF2 and NFAT while inhibiting their
398	differentiation (Horsley et al., 2003; Knight and Kothary, 2011; McKinsey et al., 2002; Stork
399	and Schmitt, 2002; Wu et al., 2007; Yue et al., 2023). In addition, it has been shown that
400	mTOR regulates the proliferation of MSCs but can also regulate myotube fusion by both
401	kinase-dependent and -independent pathways (Park and Chen, 2005).
402	In conclusion, an understanding of the various gene expression and signaling processes
403	within MSCs for cultured meat production is required, and further research is needed to
404	control and regulate cell cycle arrest and activation, proliferation, differentiation, and even
405	fusion.

# 407 *Obtaining muscle satellite cells (MSCs)*

408 MSCs can be obtained by biopsy of muscle tissue from living animals and by harvesting 409 muscle tissue from animals immediately after slaughter. The most used processes for 410 harvested muscle tissue are disinfection, removal of fat and connective tissue, fragmentation, 411 digestive enzyme treatment, sequential filtration, centrifugation, pre-culture, and finally, cell 412 recovery to obtain primary cells (Lee et al., 2021). The obtained primary cells are then 413 subjected to immunofluorescence staining or polymerase chain reaction (PCR) to determine 414 the proportion of MSCs from the proportion of progenitor regulatory factors in the primary 415 cells. Typically, Pax7 and MyoD are used to determine the purity of MSCs, and by 416 comparing their expression levels, the activation of the MSCs used in the experiment can be 417 determined (Ding et al., 2017; Kim et al., 2023a; Pasut et al., 2013). In addition, flow 418 cytometry methods, such as fluorescence-activated cell sorting (FACS) and magnetic-419 activated cell sorting (MACS), can be used to obtain pure MSCs labeled with MSC-specific 420 markers, which can then be proliferated to sufficient quantities for use in cultured meat 421 experiments and production (Ding et al., 2018; Gromova et al., 2015; Kim et al., 2023a; 422 Motohashi et al., 2014).

423

# 424 Culture of muscle satellite cells (MSCs)

The culture of MSCs has been performed since before the 1990s, and the methods can be broadly divided into two types: culture of isolated single strands of muscle fibers and culture of cells isolated from enzymatically treated muscle tissue (Anderson and Pilipowicz, 2002; Bischoff, 1986; Doumit and Merkel, 1992; McFarland et al., 1988). Fetal bovine serum (FBS) is a key ingredient added to the basal medium for culture, but the exact nature of FBS is still poorly understood, and commercialization of cultured meat is currently limited by the 431 need to replace FBS completely (Lee et al., 2022; Lee et al., 2023a). It is difficult to avoid the 432 ethical issues associated with the production of FBS, as more than 2 million bovine fetuses 433 derived from slaughtered mothers are used for FBS production each year (Lee et al., 2022). In 434 addition to ethical concerns, the high price of FBS has led numerous research teams to 435 investigate serum-free media as an alternative to FBS, and along with research to refine the 436 active ingredients of FBS, results support that serum can be effectively replaced with proteins 437 required for cell growth or a combination of various growth factors (Messmer et al., 2022; 438 Schenzle et al., 2022; Skrivergaard et al., 2023; Stout et al., 2022; Stout et al., 2023). Furthermore, to meet halal standards, the use of blood in cultured meat production is also 439 440 limited (Hamdan et al., 2018). However, challenges remain, such as the use of recombinant 441 growth factors in the preparation of serum-free media or chemically composed media and the 442 cost of expensive additives (Stout et al., 2022; Stout et al., 2023).

443 Once the medium in which the cells are to be cultured is prepared, the method of culturing 444 the cells must be chosen according to each cell type. Cell culture techniques for cultured meat 445 production can be broadly divided into adherent culture and floating culture, and among the 446 cells, MSCs and fibroblasts have been studied, as well as adipocytes (Bodiou et al., 2020; Ge et al., 2023; Humbird, 2021; Lee et al., 2021). Approximately  $10^{14}$  cells and 10,000 L of 447 culture medium are required to produce 1 t of cultured meat, assuming a cell density of  $10^7$ 448 449 cells/mL in the bioreactor (Guan et al., 2021). However, the larger the bioreactor size, the higher the stirring intensity needed to maintain a homogeneous environment in the vessel, 450 451 which can lead to shear stresses of a magnitude that can cause cell damage (Allan et al., 452 2019). In a modeling study of cultured meat production scenarios, it was emphasized that 453 optimal cell selection to reduce the consumption rate of medium, completely replace or 454 decrease the cost of growth factors, and increase the size of perfusable bioreactors are 455 necessary for mass production environments (Risner et al., 2021).

456 As such, cultured meat is a tissue engineering technique under investigation based on the 457 theory that the self-renewal ability of MSCs can be harnessed to produce dozens of times the 458 amount of muscle tissue from a small piece of muscle. Cultured meat is one of the promising 459 future technologies that can be used as an important source of meat for some countries 460 because it is less sensitive to climatic conditions than conventional meat production, but there 461 is a need to improve economic issues, such as cell acquisition, mass production and cost, and 462 the amount of culture fluid and energy required for production compared to real meat. 463 Additionally, research is being conducted worldwide to improve the qualitative limitations, 464 such as flavor, texture and structure, meat color, and nutritional content, which are different from those of real meat. 465

466

# 467 Recent Trends in Muscle Satellite Cell (MSC) Culture Technologies

468 Isolation

Bovine MSC isolation techniques for cultured meat production reported in 2023 are shown in **Table 3**. The goal of the isolation process is to obtain the raw material for cultured meat. The isolation techniques used can be broadly categorized into 1) enzymatic reactions and centrifugation to obtain MSCs and pre-culture and 2) flow cytometry to increase the purity of the MSCs.

First, an enzymatic reaction is performed to obtain primary cells from muscle tissue. The cells are minced to increase the surface area, and connective tissue is removed to facilitate the reaction. Enzymes used for MSC isolation include collagenase, dispase, trypsin, and pronase in various concentrations. Centrifugation is a method that uses centrifugal force and density gradients to remove unwanted tissue and isolate desired cells. In the isolation process of MSCs, the centrifugal acceleration was 76-1,200×g, and the time was generally around 5-15 min.

481 The cells obtained by enzymatic reaction and centrifugation are primary cells. Cell pre-482 plating or purification techniques, such as FACS and MACS, are employed to increase the 483 purity of MSCs. Pre-culture is a technique for isolating specific cells from a mixture of 484 different cell types, effectively increasing the purity of MSCs by exploiting differences in the 485 adhesion properties of primary intracellular fibroblasts and MSCs (Richler and Yaffe, 1970). 486 The preincubation time used in the isolation of bovine MSCs reported in 2023 was 1-3 h. 487 Fibroblasts begin to adhere 5 min after incubation and adhere to surfaces faster than MSCs, 488 indicating that a relatively high purity of Pax7- or MyoD-positive cells can be obtained using the preincubation process (Table 3) (Kim et al., 2022; Xu et al., 2018; Yoshioka et al., 2020). 489 490 In other studies, preincubation conditions have been shown to vary from 5 min to 24 h after 491 fibroblasts begin to adhere (Table 3). One of the effective methods for rat MSCs was preincubation for up to 10 min with shaking every 5 min (Yoshioka et al., 2020). For chicken 492 493 MSCs, it was up to 2 h with shaking every 8 min after 2 h of rest, indicating that the 494 preincubation conditions may also vary depending on species-specific cell characteristics 495 (Kim et al., 2022).

Even without the pre-culture step, the purity of the MSCs can be increased by using cell 496 497 sorting techniques, such as FACS and MACS. Some drawbacks of the flow cytometry-based 498 isolation process are that it requires expensive equipment and reagents, trained professionals, 499 and is cumbersome because of sorter-induced cellular stress (SICS), such as high-pressure 500 jets, high voltage, and laser exposure during the isolation process, and cytotoxicity that can 501 occur when using specific markers (Lopez and Hulspas, 2020). Although FACS has the 502 advantage of being able to separate cells based on their size or three-dimensional features 503 using fluorescent labeling, it has the disadvantage of expensive equipment and long analysis 504 times. MACS uses magnetic particles to sort cells more than four times faster than FACS and is less expensive, but it is difficult to apply to cells that are susceptible to magnetism orcannot be labeled (Gerashchenko, 2011).

507 Both FACS and MACS label cells with clusters of differentiation (CD), which are specific 508 markers of MSCs, and each uses a fluorescent agent for FACS and magnetic particles for 509 MACS. Specific markers for MSCs used for cell labeling are species-specific, but some 510 examples are integrin  $\alpha$ 7, vascular cell adhesion protein 1 (Vcam1), and differentiation 511 clusters, such as CD29 (integrin β1), CD34 (hematopoietic stem cell marker), CD56 (neural 512 cell adhesion molecule), and CD82 (4-transmembrane glycoprotein) (Castiglioni et al., 2014; 513 Uezumi et al., 2016; Yoshioka et al., 2020). After sorting the MSCs, they can then be 514 cultured to check the expression of Pax7 or MyoD to confirm the purity of the isolated 515 MSCs, and the proliferation and differentiation capacity of the cells can be assessed. 516 In conclusion, the cell biological characteristics necessary for the isolation of MSCs from 517 each animal species have not yet been fully identified, and comprehensive research is limited 518 by the lack of standardization of separation methods, which is an obstacle to industrialization. 519

## 520 Proliferation

MSCs obtained during the isolation process will multiply in number in a properly 521 522 conditioned growth medium. The proliferation process is directly related to the yield of 523 cultured meat, and various studies have been conducted to improve the proliferation 524 efficiency. First, the basal media commonly used for MSC culture are Ham's F-10, 525 Dulbecco's modified Eagle's medium (DMEM), and DMEM/F12, with bovine fetal serum 526 added to the media at a concentration of 10-20% (v/v) in most cases (Table 4). Basal media 527 is a solution of basic nutritional components (e.g., amino acids, glucose, lipids, nucleic acid 528 bases, inorganic salts, vitamins, buffers, pH indicators) formulated in a certain proportion 529 according to the culture conditions of the desired cells. In the culture of MSCs, the basal

medium and serum concentrations are known to be closely related to the cell proliferation
rate and myotube formation (McFarland et al., 1988). In a broiler MSC culture experiment
based on culture medium composition, DMEM was found to be more effective than McCoy's
5A medium in terms of proliferation rate and MRF expression (Flees et al., 2022). In
addition, the common view that a low glucose content is effective for chicken and bovine
MSC proliferation when using DMEM as basal medium was confirmed (Flees et al., 2022;
Zygmunt et al., 2023).

537 As adherent cells, MSCs require an ECM-based coating for proliferation and 538 differentiation. Representative ECMs used for bovine MSC proliferation have been shown to 539 be gelatin, collagen I, laminin, and Matrigel (**Table 4**). Integrin  $\alpha7\beta1$ , which is present in the 540 cell membrane of MSCs, binds to collagen and laminin, and laminin induces the proliferation and migration of satellite cells (Ö calan et al., 1988; Sanes, 2003). However, C2C12 cells 541 cultured on plates coated with ECM proteins had a better proliferation rate compared to 542 543 highly elastic coatings, such as collagen I/laminin/fibronectin hydrogels, which were not 544 conducive to inducing proliferation of MSCs (Palade et al., 2019). 545 Growth factors are cell signaling proteins. For MSCs, basic fibroblast growth factor (bFGF), vascular endothelial growth factor (VEGF), and HGF are commonly used in culture 546 547 (**Table 4**). Typically, bFGF is added to the proliferation medium at a concentration of 5-10 548 ng/mL. Accordingly, in bovine MSC cultures, a bFGF content of 10 ng/mL in the medium 549 led to a faster proliferation rate than when the bFGF content was 5 ng/mL (Zygmunt et al., 2023). In addition, the expression of various endothelial cell-derived growth factors (IGF-1, 550

- 551 HGF, bFGF, and VEGF) can stimulate MSCs to proliferate and regenerate muscle
- 552 (Yamamoto et al., 2020; Zygmunt et al., 2023).

553 However, research into cultured meat using FBS remains prevalent. Even when serum-free

554 media is used, various culture ingredients, such as basal media, coating agents, and

recombinant proteins (growth factors, hormones), are employed. To address this issue, the development of natural product-derived media that meet food regulatory requirements is underway. However, industrialization is inevitably delayed because companies and research teams cannot disclose it due to competitiveness.

559

## 560 Differentiation

561 MSC differentiation is commonly achieved by removing the proliferation medium and 562 replacing it with the differentiation medium once the cells have reached a sufficient cell density in the proliferation medium (Ding et al., 2017). The differentiation process typically 563 564 uses media containing 2% FBS or horse serum (HS) to induce serum starvation (Table 5). 565 Serum starvation is often chosen to induce differentiation of MSCs (Pirkmajer and Chibalin, 566 2011). Induction of differentiation in studies published in 2023 was mainly performed at cell 567 densities above 70% confluence, and the duration of differentiation varied by 1-10 d depending on the experimental conditions, but only one study was identified that varied the 568 569 serum concentration within the culture period (Table 5). 570 When a low-serum environment is used to induce differentiation of MSCs, extensive 571 changes occur at the transcript level, with upregulation of progenitor transcription factors and 572 markers associated with differentiation identified during the differentiation process (Dmitriev 573 et al., 2013; Messmer et al., 2022). Transient and mild levels of serum starvation (15% 574 serum, v/v) induce autophagy, which can promote cell metabolism and differentiation, but 575 5% serum starvation induces excessive autophagy, leading to cell death (Wang et al., 2023b). 576 A hypoxic environment (1-10% O<sub>2</sub>) in MSC culture can create conditions that mimic oxygen saturation in mature skeletal muscle. Moreover, a hypoxic environment (2% O<sub>2</sub>) 577 578 upregulates the myogenic regulators Pax7, Myf5, and MyoD, and intermittent hypoxic 579 exposure increases the expression of VEGF released from MSCs (Koning et al., 2011;

Nagahisa and Miyata, 2018; Urbani et al., 2012). The hypoxia-induced factor-1 (HIF-1)
signaling pathway, which is expressed in response to hypoxic conditions, is thought to be
involved in the regulation of myoblast proliferation and differentiation. Under a hypoxic
environment (1% O<sub>2</sub>), broiler MSC cultures exhibited a decrease in the level of MyoDpositive cells along with changes in the transcriptome profile (Jung et al., 2024; Li et al.,
2007).

However, serum starvation tends to be the preferred method for induction of differentiation compared to hypoxic environments, and the signaling pathways involved and their effects on differentiation remain poorly understood. Furthermore, the regulations regarding cultured meat are extensive and do not clearly differentiate between cultured meat with or without differentiated tissue, leading to confusion within the industry.

591

# 592 Conclusion

593 This study analyzed the current technology, industrialization level, and future prospects of 594 cultured meat by analyzing research reports and media reports related to the industrialization 595 of cultured meat. At present, major companies are not entering mass production except for 596 prototype development, and the reason they do not disclose related technologies is that they 597 do not have enough technological capabilities. Therefore, when investing in cultured meat 598 development companies, it is necessary to accurately assess the level of technology that the 599 company has or has acquired. Much of the focus is currently on cell acquisition technology, 600 cell line acquisition technology, and cell culture and muscle differentiation technology. While 601 the level of technology related to the industrialization of cultured meat has reached the stage 602 where prototypes can be produced, it is believed that it has not yet reached the stage where 603 production costs can be dramatically reduced and the product sold to the market.

604 Nevertheless, given the steady increase in the number and depth of studies related to the

- 605 industrialization of cultured meat and the increasing number of companies involved, it is
- 606 expected that the industrialization of cultured meat could begin in the not-too-distant future.
- 607

# 608 **Conflicts of interest**

- 609 The authors declare no potential conflict of interest.
- 610

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- 615

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- 1094 **Figure legends**
- 1095 Fig. 1. Global cultured meat and traditional meat market by year from 2022 to 2030.
- 1096 Reproduced from GlobeNewswire and Statista report (Edition 2023).
- 1097
- 1098 Fig. 2. Major product trends for cultured meat companies.
- 1099
- 1100 Fig. 3. Gene regulation and signaling pathways in myogenesis.

1101	Table 1. Countries with	h established regulations and	classification guidelines fo	r cell-based foods.
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<b>Regions/countries</b>	Departments	Policies/regulations	References
Australasia	Food Standards Australia	Cultured quail as a novel food (food standards code,	FSANZ, 2023
	New Zealand (FSANZ)	Applications No. A1269)	
European Union (EU)	European Parliament (EP)	European Parliament and of the council (No. 2015/2283)	EU, 2021
Korea	Ministry of Food and Drug	Temporary standards and recognition standards of	MFDS, 2023
	Safety (MFDS)	specification for food, etc. (No. 2023-507)	
Singapore	Singapore Food Agency	Requirements for the safety assessment of novel foods and	SFA, 2023
	(SFA)	novel food ingredients (revised on July 20, 2023)	
United Kingdom	Food Standards Agency	Cell-cultivated products (revised on November 16, 2023)	FSA, 2023
	(FSA)		
United States	United States Department of	FSIS responsibilities in establishments producing cell-cultured	USDA-FSIS, 2023
	Agriculture-Food Safety and	meat and poultry food products (No. 7800.1)	
	Inspection Service (USDA-		
	FSIS)		

	Food and Drug	Federal Food, Drug, and Cosmetic Act (U.S. Code: Title 21)	FDA, 2019; FDA,
	Administration (FDA)	Public Health Service Act (U.S. Code: Title 42)	2023b
		Fair Packaging and Labeling Act (U.S. Code, Title 15)	
1102			

Countries	Companies	Products
Argentina	Alt Meat	Beef
	BIFE	Meat
	Cell Farm Food Tech	Beef
	Granja Tres Arroyos	Chicken
Australia	Heuros	Beef, Growth factors
	Magic Valley	Lamb
	Smart MCs	Ingredients, Meat, Other
	Vow Food	Meat, Other
Austria	enGenes Biotech GmbH	Growth factors
	QUBICON AG	Bioprocessing, Equipment,
		Other
Belgium	Fishway BV	Fish
	Peace of Meat	Meat
Brazil	Ambi Real Food	Beef
	BRF	Meat
	Cellva Ingredients	Fat
	Embrapa Swine and Poultry	Chicken
	JBS	Beef
	Sustineri Piscis	Fish
Canada	Another Fish	Fish
	Appleton Meats	Beef
	Cell Ag Tech	Fish

## **Table 2. Cultured meat-related companies.**

	Evolved Meats	Meat
	Future Fields	Culture media
	Genuine Taste	Ingredients, Meat
	Meatleo	Beef, Ingredients
	Myo Palate	Pork
	Seafuture	Seafood
	The Better Butchers	Meat
	WhiteBoard Foods	Scaffolds
Chile	LiveMatrix Biotech	Beef, Fish, Tuna
	Luyef Biotechnologies, Inc.	Meat
Croatia	ANJY MEAT	Meat
Czech Republic	Bene Meat Technologies	Beef, Chicken, Pork
	Enantis	Growth factors, Meat,
		Ingredients
Czech Republic	Mewery	Beef, Culture media, Pork
Denmark	Meat Tomorrow	Beef, Pork
France	BioMimesys	Scaffolds
	Fudzs	Meat
	GOURMEY - Suprême SAS	Duck
	HCS Pharma	Scaffolds
	Vital Meat	Chicken
Germany	Alife Foods	Beef
	Bluu Seafood	Fish

	CellTec Systems GmbH	Bioprocessing, Equipment,
		Meat, Seafood
	Cultimate Foods	Fat
	denovoMATRIX	Beef, Culture media, Chicken,
		Duck, Pork
	Innocent Meat	Meat
	mk2 Biotechnologies	Ingredients, Meat, Seafood
	MyriaMeat	Beef, Pork
	Ospin Modular Bioprocessing	Bioprocessing
Iceland	ORF Genetics	Growth factors
India	Clear Meat	Culture media, Meat
	Klever Meat	Ingredients, Seafood
	MealTech Pvt. Ltd.	Chicken, Ingredients
	MyoWorks	Ingredients, Meat, Scaffolds
	Neat Meatt Biotech Pvt. Ltd.	Chicken, Fish
Israel	Aleph Farms	Beef
	Believer Meats	Meat
	Believer Meats	Meat
	BioBetter <sup>TM</sup>	Growth factors
	E-FISHient Protein	Fish
	Ever After Foods	Meat
	Forsea Foods	Fish
	Meatafora	Meat, Scaffolds
	Meatosis	Fish

	Mermade Seafoods	Seafood
	Profuse Technology	Growth factors, Meat
	Sea2Cell	Fish
	Steakholder Foods	3D printing, Beef
	SuperMeat	Chicken
	Wanda Fish Technology	Fish
Japan	DiverseFarm	Meat, Seafood
	IntegriCulture	Meat
	Nissin Food Products Co., Ltd.	Beef
	Organoid Farm, Inc.	Beef
	Shojinmeat Project	Meat
	Toppan Printing	3D printing
Mainland China	Avant Meats	Seafood
	CellX	Meat
	Jimi BioTech	Beef
	Joes Future Food	Beef, Pork
	NewDay Farm	Bioprocessing, Equipment,
		Pork
Malaysia	Cell AgriTech Sdn. Bhd	Meat, Seafood
Mexico	Micro Meat	Equipment
Netherlands	Cultured Blood	Culture media
	FoldChanges	Computational biology
	Magic Caviar	Seafood
	Meatable	Meat

	Mosa Meat	Beef
	Upstream Foods	Seafood
New Zealand	Оро Віо	Ingredients, Meat
Portugal	Cell4Food	Seafood
Republic of Korea	Baobab Healthcare	Seafood
	CellMEAT	Seafood, Shrimp
	CellQua	Seafood
	DaNAgreen	3D culture, Scaffolds
	KCell Biosciences	Ingredients, Meat
	SeaWith	Meat, Scaffolds
	Simple Planet	Meat, Seafood
	Space F	Meat
	TissenBioFarm	3D printing, Meat
	Xcell Therapeutics	Culture media
Russia	ArtMeat	Fish, Other
Singapore	Ambrosia Sciences	Meat, Seafood
	Ants Innovate	Pork
	Esco Aster Pte. Ltd.	Bioreactors
	Fisheroo	Fish
	Gaia Foods	Beef
	ImpacFat	Fish
	Meatiply	Chicken, Duck, Pork
	Shiok Meats	Crab, Fish, Shellfish
	SingCell	Meat

	Umami Meats	Seafood
	Wasna	Culture media
South Africa	Mogale Meat	Chicken, Meat
	Mogale Meats	Beef, Antelope, Other
	Newform Foods	Beef, Chicken
	Sea-Stematic	Fish
Spain	BioTech Foods	Beef
	Cubiq Foods	Meat
Sweden	Re:meat	Beef
Switzerland	Cultured Food Innovation Hub	Meat
	Mirai Foods AG	Beef
Thailand	Charoen Pokphand Foods	Meat
Turkey	Biftek	Beef, Culture media
United Kingdom	3D Bio-Tissues Ltd.	Pork, Culture media, Tissue
		templating
	Alt Atlas Ltd.	Beef, Chicken, Pork, Other
	Animal Alternative Technologies	Meat
	Biomimetic Solutions	Beef
	Bright Biotech	Meat, Growth factors,
		Ingredients
	CellRev	Bioreactors
	Cellular Agriculture Ltd.	Meat
	Extracellular	Meat, Seafood
	Higher Steaks	Pork

	Hoxton Farms	Fat, Other
	Ivy Farm Technologies	Pork
	LiquiBio	Meat, Seafood
	Moolec	Meat
	Multus Media	Culture media
	Quest Meat	Beef
	Roslin Technologies	Meat
	Uncommon	Beef
United States	Aqua Cultured Foods	Seafood
	Ark Biotech	Bioreactors
	Artemys Foods	Beef
	Atlantic Fish Co.	Seafood
	Balletic Foods	Meat
	BioBQ	Beef, Scaffolds
	BioCraft	Meat, Other
	Blue Ridge Bantam	Avian, Chicken
	Bluefin Foods, Inc.	Fish
	BlueNalu	Fish
	CellCrine, Inc.	Beef, Chicken, Pork
	Clever Carnivore	Beef, Chicken, Pork
	Cultured Abundance	Meat
	Cultured Decadence	Fish, Lobster, Shellfish
	CytoNest, Inc.	Scaffolds
	Defined Bioscience	Culture media

Eat Just - GOOD Meat	Meat, Chicken
Ecovative Design	Scaffolds
Edge Foods	Beef, Chicken, Pork
Excell	Meat, Scaffolds
Finless Foods	Fish, Tuna
Fork & Good	Meat
GenScript	Beef, Chicken, Fish, Pork,
	Tuna
iLabs	Bioprocessing, Equipment
Jellatech	Scaffolds
Kiran Meats	Beef
Lab Farm Foods	Chicken, Pork
Marinas Bio	Fish
Matrix F.T.	Meat, Microcarriers
MilliporeSigma	Bioprocessing, Equipment,
	Ingredients, Other
Mission Barns	Pork
Molecular Devices	Bioprocessing, Equipment,
	Other
Myodenovo	Meat
New Age Meats	Pork
NouBio	Culture media, Microcarriers
Novel Farms	Pork, Scaffolds
OceanTastes, Inc.	Shellfish, Other

	Ohayo Valley	Beef
	Omeat	Beef, Chicken, Fish, Pork
	Optimized Foods	Mushrooms
	Orbillion Bio	3D printing, Beef, Lamb
	Pearlita Foods	Oysters
	Provenance Bio	3D printing, Scaffolds
	Reel Foods	Seafood
	SciFi Foods	Beef
	Sound Eats	Fish
	SunP Biotech	3D printing, Scaffolds
	Triplebar Bio	Cell lines
	TruSpin Nanomaterials	Scaffolds, Other
	Umami Bioworks	Fish, Shellfish, Tuna, Other
	Upside Foods	Beef, Chicken, Duck
	Vivax Bio	3D printing
<u>~</u>	Wildtype	Fish, Salmon
Vietnam	Minh Phu Seafood	Shrimp

1104 Abbreviations: 3D, three-dimensional.

Muscles	Enzymes	Centrifugation conditions	Pre-plating	References
Longissimus	Pronase	500×g, 10 min	N/A	Kim and Kim, 2023
Biceps femoris	Pronase	300×g, 5 min; 1,200×g, 15 min	N/A	Kim et al., 2023a
Longissimus	Pronase	500×g, 10 min	N/A	Kim et al., 2023b
Longissimus thoracis	Collagenase II, Dispase II	N/A	3 h + 3 h	Lee et al., 2023b
Semitendinosus	Collagenase	N/A	N/A	Messmer et al., 2023
Top round	Collagenase mix	800×g, 5 min	N/A	Park et al., 2023
Semimembranosus	Collagenase, Trypsin	100×g, 5 s; 1,000×g, 10 min	N/A	Skrivergaard et al., 2023
Semitendinosus	Collagenase II	N/A	N/A	Stout et al., 2023
Longissimus thoracis	Collagenase, Trypsin	N/A	1 h	Tzimorotas et al., 2023
Longissimus lumbrorum	Collagenase D	76×g, 5 min	1 h	Uyen et al., 2023
Hind limb	Collagenase II, Trypsin	N/A	N/A	Zhang et al., 2023
Longissimus dorsi	Collagenase II	500×g, 10 min	1 h	Zygmunt et al., 2023

## **Table 3. Isolation methods for bovine muscle satellite cells published in 2023.**

1106 Abbreviation: N/A: not applicable.

Basal media	Sera	Growth factors	Coatings	References
DMEM	10% FBS	N/A	N/A	Kim and Kim, 2023
DMEM/F12	10% FBS	N/A	N/A	Kim et al., 2023a
DMEM	10% FBS	N/A	N/A	Kim et al., 2023b
Ham's F-10	20% FBS	bFGF	Collagen I	Kim et al., 2023c
Ham's F-10	20% FBS	bFGF	Collagen, Matrigel	Oh et al., 2023
Ham's F-10	20% FBS	N/A	Bovine collagen I,	Park et al., 2023
			Matrigel	
DMEM/F12,	Serum-free media, 20%	bFGF, HGF, Hydrocortisone,	Fibronectin, Laminin	Messmer et al., 2023
Ham's F-10	FBS	IGF-1, IL-6, ITSE, PDGF,		
		VEGF		
DMEM,	10% FBS, Serum-free	bFGF, Fetuin, ITS, HGF,	Matrigel	Skrivergaard et al., 2023
DMEM/F12	media	PDGF, Insulin		
DMEM	20% FBS	bFGF	Laminin, Vitronectin	Stout et al., 2023

## **Table 4. Proliferation methods for bovine muscle satellite cells published in 2023.**

LG-DMEM	10% FBS, 2% FBS,	N/A	Entactin-Collagen-	Tzimorotas et al., 2023
	Ultroser G		Laminin	
DMEM	15% FBS	N/A	Rat tail collagen I	Uyen et al., 2023
DMEM	20% FBS	bFGF	N/A	Zhang et al., 2023
LG-DMEM, HG	- 20% FBS	bFGF	Gelatin	Zygmunt et al., 2023
DMEM				

1108 Abbreviations: DMEM: Dulbecco's modified Eagle's medium; DMEM/F12: Dulbecco's modified Eagle's medium and Ham's F-12 Nutrient

1109 Mixture; LG-DMEM: low glucose-DMEM; HG-DMEM: high glucose-DMEM; FBS: fetal bovine serum; bFGF: basic fibroblast growth

1110 factor; HGF: hepatocyte growth factor; IGF-1: insulin-like growth factor-1; IL-6: interleukin-6; ITS: insulin-transferrin-selenium; ITSE:

1111 insulin-transferrin-selenium-ethanolamine; PDGF: platelet-derived growth factor; VEGF vascular endothelial growth factor. N/A: not

1112 applicable.

Basal media	Sera	Time (d)	References
SILAC DMEM Flex Media	2% HS	4	Kim and Kim, 2023
DMEM/F12	2% HS	1-4	Kim et al., 2023a
DMEM	2% FBS	1-4	Kim et al., 2023c
DMEM	2% HS	6	Lee et al., 2023b
DMEM/F12	Serum-free	3	Messmer et al., 2023
DMEM	2% FBS	4-5	Oh et al., 2023
DMEM	2% FBS	1-6	Park et al., 2023
Neurobasal	N/A	2	Stout et al., 2023
DMEM	2% HS	3	Uyen et al., 2023
DMEM	2% HS, 10% HS	3-7	Yun et al., 2023
DMEM	2% HS	1-5	Zhang et al., 2023
LG-DMEM, HG-DMEM	20% HS	3-10	Zygmunt et al., 2023

## 1113 **Table 5. Differentiation methods for bovine muscle satellite cells published in 2023.**

1114 Abbreviations: DMEM: Dulbecco's modified Eagle's medium; DMEM/F12: DMEM and

1115 Ham's F-12 Nutrient Mixture; LG-DMEM: low glucose-DMEM; HG-DMEM: high glucose-

1116 DMEM; HS: horse serum; FBS: fetal bovine serum. N/A: not applicable.

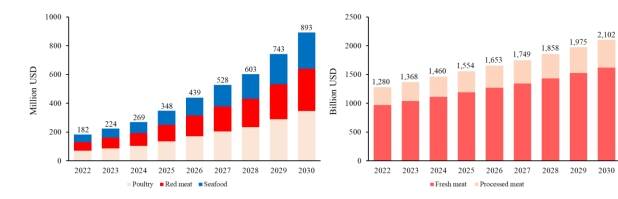


Fig. 1.

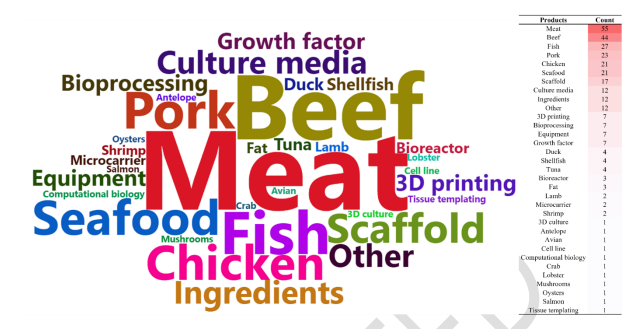


Fig. 2.

