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1 2	The Gut Microbiome and Pharmacology: A Prescription for Therapeutic Targeting of the Gut-Brain Axis
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Abstract

17 New frontiers for host-microbe interactions continue to emerge as our knowledge of the adult gut 18 microbiome in health and disease is continually supplemented and improved. Alterations in the gut 19 microbiota composition in irritable bowel syndrome (IBS) are now linked to symptom severity while population based evidence linking gut microbiome signatures to depression is an important 20 21 new landmark. The effects of drugs on gut microbiome composition is also becoming clearer. 22 Meanwhile, preclinical studies have delineated the influence of the gut microbiome at a structural 23 and activity level in distinct brain regions. Bacterial metabolites, such as tryptamine, can activate 24 specific receptors to impact gastrointestinal motility. These recent studies bring into focus the future implications for therapeutic targeting of the microbiome-gut-brain axis. 25

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Introduction

29 As our knowledge of the important role played by the gut microbiota in health and disease expands, new frontiers for host-microbe interactions continue to emerge. Recently, traditional concepts in 30 31 pharmacology and therapeutics have been challenged by reports outlining reciprocal microbiome-32 xenobiotic interactions and a growing appreciation that microbial metabolites might exert their 33 effects via receptor-mediated mechanisms. In this review, we first outline the most salient aspects 34 of the composition and function of the gut microbiome as a framework to understand the 35 importance of this virtual organ for gastrointestinal pharmacology and beyond. We then focus on 36 a number of key recently published articles illustrating the implications of important conceptual 37 advances that chart the scope and scale of microbial regulation of pharmacodynamics and pharmacokinetics in the gut-brain axis. This is considered within the context of the bidirectional 38 39 relationship between xenobiotics and our gut bacteria. Finally, we attempt to integrate these 40 observations to elaborate on the future implications for therapeutic targeting of the microbiome-41 gut-brain axis.

42 The Adult Gut Microbiome: A Metabolic Powerhouse

43 The adult gut microbiota is made up of trillions of microorganisms (bacteria, viruses, archaea, 44 yeasts and fungi) that reside in the gastrointestinal tract, contributing substantially to host 45 physiological homestasis. The community of bacteria is best studied with the highest density in the large intestine which according to recent estimates reaches 10¹³ bacterial cells in the human colon 46 [1,2]. The composition and function of this complex bacterial ecosystem is individual –specific 47 48 and impacted by a number of intrinsic and extrinsic factors, including diseases and drug use, diet, 49 age and lifestyle of the host [3-5]. Recent sequencing surveys confirm that the adult gut microbiota 50 is dominated from a compositional perspective by the phyla Firmicutes, Actinobacteria and 51 Bacteroidetes with lower relative abundances of Verrucomicrobia and Proteobacteria. There may 52 also be a core microbiota defined by 14 different genera with medication use in general contributing to microbiota compositional variation [3]. Our knowledge of the complexity of this virtual organ 53 54 continues to expand and through the use of sequencing approaches, metagenomic analysis and 55 bioinformatic pipelines. Pasolli and colleagues [6] have recently elegantly revealed the presence of 56 new microbial species on or in the host, including the gut, associated with westernized or nonwesternized lifestyles. In addition, many newly identified species-level operational taxonomic 57

units (OTUs) may be associated with disease states as their genome sequences were not previously
 captured in databases [7].

The aggregate genome of this community, the metagenome, far exceeds and complements the 60 metabolic capacity of the host genome. These microbial genes encode an array of metabolic 61 activities, providing the host with additional essential functional capacity, such as the digestion of 62 63 dietary fibers, which yields microbial metabolites important for host-microbe interactions. All 64 these recent reports continue to support the importance of the gut microbiota in human health, although there remains knowledge gaps surrounding the precise composition of a healthy gut 65 microbiome across the life span and more granular details on the molecular mechanisms 66 underpinning complex host-microbe interactions, particularly in the context of gastrointestinal 67 68 pharmacology.

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70 The Gut Microbiome in Disease

Shifts in the bacterial composition, structure or function in the gastrointestinal tract have been 71 72 associated with numerous disorders in the last few decades. As studies go beyond microbial surveys, the quality of the information derived from these studies continues to improve. For 73 74 example, it now appears that alterations in the gut microbiota composition in irritable bowel 75 syndrome (IBS) may include microbiota signatures associated with symptom severity [8]. In 76 particular, IBS symptom severity was negatively associated with microbial richness as well as the 77 presence of methanogens, and gut microbiota enterotypes characterized by enriched *Clostridiales* 78 or Prevotella species [8]. This confirms the importance for the gut microbiota in the development 79 of functional gastrointestinal disorders as well as chronic inflammatory diseases (see [9]).

80 With the increasing number of studies focused on the gut microbiota and mental health, 81 compositional alterations have also been highlighted in psychiatric and neurological disorders, such 82 as Alzheimer's disease [10], Parkinson's disease [11,12], autism spectrum disorders (ASD) [13], 83 schizophrenia [14] and depression [15]. In many cases, a causal role for these disease-associated microbiome configurations can be inferred from the transfer of behavioural phenotypes to animal 84 85 models via the microbiota [15,16]. In the case of IBS, this even extends to the transfer of specific psychiatric comorbidities such as anxiety [16]. More recently, the analysis of a large microbiome 86 87 population cohort enabled the identification of *Dialister* and *Coprococcus* spp as indicators of high quality of life, and revealed their depletion in depressive patients [17]. The results of this study 88

have advanced our knowledge further, providing the first population based evidence linking gut microbiome compositional signatures with a mental health disorder. It is therefore becoming increasingly important to consider the intestinal microbiota as a biomarker reservoir, in the development of new treatments and as a source of the side effects associated with particular hostdirected medications.

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96 The Gut Microbiome and Expanding array of Therapeutic Targets in the Gut-brain Axis:

97 While microbial signatures or alterations in the composition of the microbiota now appear to be 98 evident in various pathologies, the extent of, and mechanisms involved in, this communication 99 remain to be fully grasped. A variety of preclinical approaches, including the use of germ-free 100 animals (GF), have allowed the scope of influence of the enteric microbiota on the brain-gut axis 101 to be defined. Abdominal pain, underpinned by visceral hypersensitivity, is a core feature of 102 irritable bowel syndrome (IBS). Recently, it has been conclusively demonstrated that the gut 103 microbiota is required for normal visceral pain sensation, associated with increases in toll-like 104 receptor and cytokine gene expression in the spinal cord. This study also demonstrated that the 105 volumes of brain regions involved in pain processing such as the anterior cingulate cortex (ACC) 106 and periaqueductal grey, were decreased and enlarged respectively in GF mice. [18]. This is 107 consistent with previous studies which have demonstrated that the visceral hypersensitivity of IBS 108 patients can be transferred to GF rats via the fecal microbiota [19].

109 Microbial regulation of the transcriptional activity in different brain areas, such as amygdala, 110 prefrontal cortex or hippocampus, is now supported by several studies and often occurs in a sex-111 dependent manner [20-22]. Studies in GF animals also now implicate the gut microbiome in 112 appropriate regulation of microRNA (miRNAs; non coding RNAs that act through translational 113 repression to control gene expression) expression in brain regions implicated in anxiety-like 114 behaviours such as the amygdala and prefrontal cortex [22] or in memory and learning, such as the 115 hippocampus [21,23]. For instance, in the absence of a gut microbiota, the basal expression of 116 specific activity-related genes in the amygdala is altered, leading to the suggestion that a 117 hyperactivity of this brain structure might be at the root of the behavioural abnormalities associated with growing up germ free [24-27]. Whether this can be exploited therapeutically is an open 118

question but in support of this possibility, the behavioural consequences as well as the molecular signature of his hyperactivity can at least partially be reversed by the colonization of GF animals [25]. Of further interest is that fecal miRNAs of host or plant origin may have an important role in dictating microbiota composition, possibly by targeting regions in bacterial metagenomes [27-30] while fecal miRNA expression is also linked to gut microbiota fluctuations [31].

124 A recent study, based on a mouse model of autism (BTBR mice), highlighted a significant decrease 125 of two bile-metabolizing species: Bifidobacterium and Blautia. Moreover, this compositional shift 126 was associated with deficient bile acid and tryptophan metabolism, gastrointestinal dysfunction 127 and impaired social interactions [32]. These results support the concept that modulation of the gut 128 microbiota could be a promising strategy for the treatment of brain-gut axis disorders. In this 129 context, Burokas and his team assessed the effect of the administration of two prebiotics in a rodent 130 study: the gluco- and the fructo-oligosaccharides (GOS and FOS). Besides modifying the 131 expression of genes such as BDNF in the hippocampus, GOS and FOS also exerted anxiolytic and 132 antidepressant effects and reversed the behavioral and physiological impact of chronic stress 133 exposure [33]. The finer details of the mechanisms mediating these beneficial effects remains 134 unclear in many cases but substantial progress has been made in this area, particularly in the context 135 of pharmacodynamic interactions between microbial metabolites and the host.

136 The Gut Microbiome and Pharmacodynamics

137 Bacterial metabolites are considered likely to be key mediators of these host microbe interactions 138 with the possibility they can induce host cellular responses via their activity at G-protein-coupled 139 receptors (GPCRs) expressed either locally in the gastrointestinal tract or at more distal locations 140 [34]. One such example is tryptamine (a monoamine similar to 5-hydroxytryptamine (5-HT)), 141 metabolized by bacteria via tryptophan decarboxylation, which modulate colonic secretion via 142 activation of the 5-HT4 receptor (5-HT4R), a 5-HT receptor expressed in the colon of importance 143 for regulation of gastrointestinal motility [35-37]. Another receptor of importance in this regard is the aryl hydrocarbon receptor (AhR) and a reduction of the microbiota's ability to metabolize 144 tryptophan into ligands capable of activating this has been identified in metabolic syndrome [38] 145 146 and colitis [39], supporting the importance of this bacterial product in receptor-mediated host 147 homeostasis.

148 In other cases, microbial metabolites may alter the expression of key receptors to influence 149 gastrointestinal function. The most studied metabolites produced by gut bacteria are the short chain 150 fatty acids (SCFAs), derived from the fermentation of dietary fibers. For example, acetate 151 production can regulate the expression of 5-HT3 receptor expression to influence host secretory 152 patterns [36]. Beyond intestinal-located interactions and although well known for their direct interactions with the free fatty acid receptor (FFAR) 2 and 3 in the regulation of appetite and energy 153 intake, SCFA supplementation has recently been associated with antidepressant and anxiolytic 154 155 effects in mice. These effects were not present following exposure to a psychosocial stressor but 156 the SCFA treatment did alleviate stress-induced increases in intestinal permeability while the 157 stress-induced alterations in colonic gene expression of the SCFA receptors free fatty acid receptors 158 were unaffected by SCFA supplementation [40].

159 The gut microbiota can also secrete compounds able to translocate from the gut to the systemic 160 circulation, and to subsequently cross the blood-brain barrier. This applies to bacterial 161 peptidoglycan (PGN), a major component of the bacterial membrane, which is able to activate 162 neuronal pattern-recognition-receptors (PRR), leading to modulation of brain development during specific time windows, through an interaction with Pglyrp2 [41]. A deeper understanding of the 163 164 functional implications and regulation of bacterial-products could then constitute a relevant strategy for modulating host homeostasis, and potentially the development of new therapies in a 165 166 wide range of gut-brain axis disorders.

167 The Gut Microbiome, Pharmacokinetics and Toxicity

168 The study of pharmacokinetics has traditionally focused on the impact of the host on administered 169 drugs without due regard for the functional capacity of the gut microbiota. Orally administrated 170 drugs in particular represent a potential substrate for bacterial metabolism, which can lead to 171 intrapersonal variations in drug availability, efficacy or toxicity. One of the prospective drugs for 172 such a transformation was the immunosuppressant mycophenolate mofetil (MMF), which, despite its effectiveness, induces significant side effects. Nevertheless, treating GF mice with MMF 173 174 showed significantly reduced side effects [42], strongly implicating the gut microbiota in the 175 emergent adverse effects.

The bacteria inhabiting our gut have at their disposal a range of microbial enzymes able to modify 176 177 drugs and other xenobiotics. Tyrosine decarboxylase (TDC), expressed in particular by Enterococcus and Lactobacillus, was pointed out for its ability to interfere in the treatment of 178 179 Parkinson's disease through the inactivation of levodopa (L-DOPA) [43]. Moreover, it seems like 180 prolonged treatment with L-DOPA enhances tdc gene expression, leading to a less and less 181 effective treatment over time. F. Prauznitzii and Clostridiales, other specific enteric bacteria, have also been involved in the decrease of effectiveness of the immunosuppressant tacrolimus [44], 182 183 highlighting the potential negative effect of gut microbiota on an orally administered medical treatment. In a similar vein, a bioinformatic approach enabled the identification of tyramine oxidase 184 185 expressed by E. Coli as capable of binding amphetamine, leading to a potential modification of the 186 drug [45]. Together, these results substantiate the relevance of using new models in pharmacology, 187 that take into consideration microbial metabolism and the associated intra-individual variations. 188 After an adaptation for other xenobiotics, the pharmacokinetic model built by Zimmermann and 189 his team would hence represent an interesting basis to separate host and microbiome contributions 190 to pharmacokinetics and toxicity [46].

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192 Effects of drugs on the gut microbiome

193 While, as shown above, the microbiota can have negative effects on the pharmacological properties 194 of drugs, the reverse pattern is also valid: a large number of host-directed drugs across therapeutic 195 classes combined, can affect the bacterial growth of at least 1 strain in vitro [47]. Psychotropic 196 drugs have been particularly highlighted for their antimicrobial effects, causing alterations of the 197 microbiota as well as modifications of gastrointestinal function such as intestinal permeability in 198 vivo, and impacting on bacterial growth in vitro (Table 1) [48,49]. Earlier studies indicated that 199 olanzapine altered the composition of the gut microbiota [50]. Further studies focusing on this drug 200 showed that the microbiota was needed for drug-associated weight gain, a serious and common side effect of this antipsychotic treatment [51], as antibiotics attenuated the side effects in mice. 201 This was also true in germ-free animals and olanzapine has antimicrobial effects on the growth of 202 203 E. Coli and Enterococcus faecalis in vitro [52]. This opens up the possibility of targeting the gut 204 microbiota with, for example, prebiotics to try and limit these adverse side effects [53].

Alternative approaches allowing the modulation of the enteric microbiota, such as fecal microbiota 205 206 transplant (FMT), might also lend themselves to counterbalance, or at least limit, these adverse effects or indeed promote beneficial effects. Interestingly, the ketogenic diet (KD) which is used 207 208 to treat refractory epilepsy, appears to induce alterations in the microbiota which are necessary for 209 its anti-seizure effects [54]. Together with the FODMAP diet for control of IBS symptoms [55], 210 this is an example of a diet of reduced diversity which would not normally be considered beneficial for our gut microbes but which produce symptomatic improvements in the host. According to our 211 212 current knowledge of the gut microbiota and host-microbe interactions within the framework of pharmacokinetics and pharmacodynamics, the effects of a wide range of host-directed xenobiotics 213 214 on our bacteria community has to be more routinely considered in drug development pipelines.

215 Conclusion

216

217 Recent research has aided substantially our efforts to make sense of the microbiome-gut-brain axis in gastrointestinal pharmacology and beyond. This includes advances over compositional surveys 218 219 to important studies linking symptom severity to gut microbiome alterations in IBS, as well as 220 landmark population based evidence linking gut microbiome signatures to depression and quality 221 of life. Moreover, the increase in research linking the gut microbiome to neuropsychiatric disorders 222 from clinical studies is supplemented with preclinical approaches that implicate the gut microbiome 223 in regulating even the structure and activity of key brain regions. Meanwhile, traditional concepts in pharmacology will likely need to be redrawn to account for the reciprocal interactions between 224 225 our gut microbes and xenobiotics. This will have important implications for pharmacodynamics 226 and pharmacokinetic considerations during drug development. Our understanding of the molecular 227 mediators underpinning host-microbe interactions now includes an appreciation that microbial metabolites can impact on specific receptors to influence aspects of host physiology such as 228 229 gastrointestinal motility. It remains an appealing prospect that this knowledge can be harnessed 230 effectively for therapeutic targeting of the microbiome to influence gut-brain axis signaling using 231 interventions such as FMT, prebiotics, probiotics or postbiotics. Limiting the side effects associated 232 with psychotropic drugs such as antipsychotics via microbiome-based approaches is a further avenue of investigation with high potential. Effectively translating these promising recent 233 234 advances into the prescription pads of the future is an ambitious but important research objective.

235 **Conflict of Interest**

APC Microbiome Ireland collaborates with a number of industry partners including Dupont Nutrition Biosciences APS, Cremo SA, Alkermes Inc., 4D Pharma PLC, Mead Johnson Nutrition, Nutricia Danone and Suntory Wellness. GC has spoken at meetings sponsored by food and pharmaceutical companies including Janssen Ireland. This neither influenced nor constrained the content of this review.

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496 Figure Legends

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498 Figure 1: The Microbiome-gut-brain axis and Psychiatry

The composition of the gut microbiome is under the influence of various intrinsic and extrinsic 500 501 factors, such as the host genetics, age, and other lifestyle factors. The gut microbiome can recruit 502 the gut-brain axis, a bidirectional communication system between the brain and the gut, to influence 503 brain function and behaviour. Alterations in the composition and function of the gut microbiome 504 have been associated with a number of clinical psychiatric and neurological disorders while 505 preclinical approaches confirm the capacity of our gut microbes to exert behavioural and functional 506 effects of relevance to these brain disorders. Psychological stress exposure can also impact on the structure and function of the gut microbiome. 507

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509 Figure 2: Xenobiotics and Gut Microbiota Interactions

511 Orally administrated drugs are, after ingestion, in direct contact with the gut microbiome. The co-512 localization of bacteria and xenobiotics may result in reciprocal interactions. On one hand, many 513 xenobiotics have antimicrobial properties and can alter microbiota composition, diversity and 514 function, often in a manner that can be linked to the side effects of various medications. On the 515 other hand, the gut microbiota can metabolize the ingested drugs or indirectly alter their 516 metabolism by the host and this can result modification of availability, efficacy or toxicity of the drug in the organism. Many disease states are also associated with gut microbiome alterations, even 517 518 prior to drug use although the implication of this are currently unclear.

Table 1: Psychotropic drugs and their effects on the gut microbiome and intestinal physiology in preclinical studies 522

Drug	Disease	Observed effects	Reference
	Schizophrenia, major depression, bipolar disorder, obsessive- compulsive disorder	↑ bacterial richness and diversity	
Aripiprazole		↑ Firmicutes	[40]
		\uparrow the levels of acetate and isovalerate	[48]
		↑ distal ileum permeability	
		↓ E. Coli IAI1 & L. gasseri	[56]
Escitalopram	Depression/anxiety disorders	\downarrow <i>E. Coli</i> growth <i>invitro</i>	[48]
Escitatopiani		↑ distal ileum permeability	ومبا
	Depression/anxiety disorders	Inhibit L. rhamnosus and E. Coli growth	
		\downarrow Deferribacteraceae	[48]
Fluoxetine		↑ distal ileum permeability	
		↑ <i>Firmicute/Bacteroidete</i> ratio	[49]
×	Bipolar disorder, mood- stabilizer, major depression, schizophrenia	↑ bacterial richness and diversity	[48]
Lithium		↑ Actinobacteria et diminution Bacteroidetes	
	Schizophrenia, bipolar disorder	\uparrow level of <i>Firmicute</i> and \downarrow bacterial diversity in females	
		\downarrow <i>Proteobacteria</i> in males	[50]
Olanzapine		↓ Bacteroidetes	
		\uparrow <i>Firmicutes</i> and \downarrow <i>Bacteroidetes</i>	[51]
		\downarrow E. Coli and Enterococcus faecalis croissance in vitro	[52]
	Epilepsy, bipolar disorder, schizophrenia	\uparrow bacterial richness and diversity	
Valproate		\uparrow Actinobacteria and Firmicute - \downarrow Bacteroidete	[48]
		\downarrow propionate and butyrate levels and \uparrow isovalerate	
Venlafaxine	Depressive/anxiety disorders	↑ distal ileum permeability	[48]

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