Possibilities and limits for using the gut microbiome to improve captive animal health

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Abstract
Because of its potential to modulate host health, the gut microbiome of captive animals has become an increasingly important area of research. In this paper, we review the current literature comparing the gut microbiomes of wild and captive animals, as well as experiments tracking the microbiome when animals are moved between wild and captive environments. As a whole, these studies report highly idiosyncratic results with significant differences in the effect of captivity on the gut microbiome between host species. While a few studies have analyzed the functional capacity of captive microbiomes, there has been little research directly addressing the health consequences of captive microbiomes. Therefore, the current body of literature cannot broadly answer what costs, if any, arise from having a captive microbiome in captivity. Addressing this outstanding question will be critical to determining whether it is worth pursuing microbial manipulations as a conservation tool. To stimulate the next wave of research which can tie the captive microbiome to functional and health impacts, we outline a wide range of tools that can be used to manipulate the microbiome in captivity and suggest a variety of methods for measuring the impact of such manipulation preceding therapeutic use. Altogether, we caution researchers against generalizing results between host species given the variability in gut community responses to captivity and highlight the need to understand what role the gut microbiome plays in captive animal health before putting microbiome manipulations broadly into practice.

Keywords: Captivity, Gut microbiome, Animal, Wild, Conservation, Function

Introduction
In captivity, wild animals are subject to environments and lifestyles that they would not experience in their natural habitat as a result of significant human manipulation of the animals’ built environment, diet, healthcare, and social interactions [1]. These novel circumstances can lead to improved animal welfare and longevity in some species [2], but other species suffer from serious health problems under human care and human environments [2, 3]. There are diverse health problems which have been associated with captivity including issues with metabolism and digestion [4, 5], infection [6, 7], stress [8, 9], and reproduction [10, 11]. Reflecting its growing importance in other fields of organismal research, the microbiome has been proposed as a mediator of host condition under captivity [e.g. 12]. However, the extent, predictability, and drivers of microbiome changes under captivity remain unclear, limiting our ability to utilize microbial interventions to alleviate captivity-associated health problems.

Given the vast number of environmental variables that differ between captive and wild environments, it is difficult to predict how a particular species will fare in captivity [2]. The potential improvements (including veterinary care, freedom from predators, and increased availability of food [2]) and potential stressors (reduced range and mobility, artificial social groups, frequent proximity and contact with humans, etc. [3]) under captivity are numerous and hard to isolate from one another. Moreover, physical and behavioral differences between species may modulate the impact of certain changes and lead to...
varying health outcomes. For example, species that are typically solitary in the wild likely face different challenges than gregarious species when placed into captive environments with set social groups [13].

Microbial plasticity or heterogeneous microbial sensitivity could also contribute to differential animal responses under captivity. Diet and environment are known to shape the composition of the gut microbiome [e.g. 14, 15] with downstream effects on host metabolic, immune, and neurological systems in humans and animal hosts [16–18]. However, not all microorganisms respond similarly or to the same extent when exposed to new diets or housing conditions, and not all hosts will experience physiological effects as a result of microbiome plasticity [19, 20]. For example, domestication has been found to consistently impact the gut microbiota, but there is no singular domesticated gut microbiota [21]. Diet and captive habitats, as well as evolutionary change, can contribute to domestication related microbial shifts; for instance, wild horses kept in zoos show greater gut microbial similarity to domesticated horses than do wild animals in their native habitat [22].

Given the potential connection of gut microbial changes to downstream health effects and the high prevalence of gastrointestinal issues in captive animals (such as helicobacter gastritis seen in most captive cheetahs [23, 24], callitrichid wasting syndrome observed in various New World monkeys [4, 25], and Cryptosporidium infection that affects captive sifakas [26]), some have suggested including the microbiome in conservation management plans for captive animals [e.g. 27, 28]. These suggestions are inspired by strong effects of captivity observed in comparisons between wild and captive animal microbiomes. The generalizability of captivity effects on the microbiome and their connection to host health remain unclear, however, because to date there has not been a comprehensive review of the literature.

One of the particular challenges of studying health in response to the microbiome is that health is difficult to define and measure and varies on a species-by-species basis. Health extends beyond survival and reproduction and encompasses many axes of physical and mental well-being. In this paper we refer to health broadly, much like Fisher’s definition of animal welfare as “an evaluative term […] encompassing aspects of natural selection, coping, well-being, satisfaction of preferences, fulfilment of needs, and natural behaviour, or a combination of them” [29]. We emphasize that indicators of health and disease will vary between species, and there is no single definition of host health much like there is no single definition of a healthy microbiome [30]. For example, for a species struggling with gastrointestinal disease such as the red wolf [31], improved health may be defined as decreased instances of disease. Meanwhile, for a species that suffers from captive infertility such as the southern white rhinoceros [32], improved health may simply mean increased fecundity.

In this paper, we intend to help identify and address gaps in our understanding of the captive gut microbiome and in turn host health. We review the literature on how and why the gut microbiome responds to captivity, as well as what is known about the health consequences of these changes. To conclude, we highlight major outstanding questions in the field brought to light by our review and propose priority areas for future work, with an emphasis on research relevant to conservation efforts.

Review of current literature on the captive microbiome

The scope of captivity’s influence on the gut microbiome can be summarized with three broad questions. How does the microbiome change in response to captivity? Why does it change? And, do these changes affect overall animal health for the better or for the worse? We focus here on the gut microbiome, which has been the subject of the preponderance of work on captive microbiomes, but note that an important exception considered elsewhere [e.g. 33, 34] is the amphibian skin microbiome.

How does the gut microbiome change in response to captivity?

There have been more than 60 studies over recent decades characterizing the effects of captivity on the gut microbiome of host species spanning the animal kingdom. Though the term “captivity” can refer to a variety of habitats such as zoos, sanctuaries, labs, farms, or even households, most of these studies examine captivity in the context of zoos or sanctuaries and sometimes, but not always, focus on endangered species. Altogether, it is clear that there is no singular “captivity microbiome,” as differences in microbial diversity and community composition between captive and wild environments vary greatly among hosts (Table 1).

Many studies that measured microbial diversity report decreased gut richness or diversity in individuals in captivity compared to others in the wild [e.g. 22, 35, 36]. However, some studies have reported no difference in microbial diversity between the two environments [e.g. 37, 38], or even increased diversity in captive animals [e.g. 39–41]. The most comprehensive test of captivity effects to date demonstrates this heterogeneity persists even when sampling and sequencing methods are controlled for [42]. Of the 11 mammalian families represented in that analysis, only four showed significantly decreased bacterial diversity in captivity. Six other families showed no significant difference between the captive
| Author            | Year | Citation | Sample type | Species                                | Difference in captive/wild microbial diversity? | Difference in captive/wild microbial composition? | Functional implications discussed?                                                                 |
|-------------------|------|----------|-------------|----------------------------------------|-------------------------------------------------|-------------------------------------------------|-----------------------------------------------------------------------------------------------------|
| Alfano et al.     | 2015 | [113]    | rectal      | Koala                                  | Not reported                                     | N                                               | Not discussed                                                                                       |
| Allan et al.      | 2018 | [114]    | fecal       | Amargosa vole                          | No difference                                    | Y                                               | Discussed                                                                                           |
| Allan et al.      | 2018 | [114]    | foregut     | Amargosa vole                          | No difference                                    | N                                               | Discussed (metabolic function)                                                                     |
| Amato et al.      | 2016 | [5]      | fecal and intestinal | Various colobine species | No difference                                    | Y                                               | Discuss (metabolic function)                                                                       |
| Benno et al.      | 1987 | [115]    | fecal       | Japanese macaque                       | Higher in captivity                               | Y                                               | Discussed (metabolic function)                                                                     |
| Bik et al.        | 2016 | [116]    | rectal      | Bottlenose dolphin                    | Not reported                                     | N                                               | Not discussed                                                                                       |
| Borbon-Garcia et al. | 2017 | [73]    | fecal       | Andean bear                            | Lower in captivity                               | Y                                               | Assessed (predicted metabolic functions—KEGG pathways/PICRUSt)                                     |
| Cheng et al.      | 2015 | [72]     | fecal       | Tasmanian devil                        | Lower in captivity                               | Y                                               | Assessed (predicted metabolic functions—KEGG pathways/PICRUSt)                                     |
| Clayton et al.    | 2018 | [44]     | fecal       | Red-shanked douc                       | Lower in captivity                               | Y                                               | Assessed (predicted metabolic and antibiotic resistance functions—KEGG pathways/PICRUSt)           |
| Clayton et al.    | 2016 | [55]     | fecal       | Red-shanked douc, Mantled howling monkey | Lower in captivity                               | Y                                               | Assessed (predicted metabolic functions—PICRUSt)                                                   |
| De Jesus-Laboy et al. | 2011 | [117]   | fecal       | Goat                                  | Not reported                                     | N                                               | Assessed (Assess presence of antibiotic resistance genes)                                             |
| Delport et al.    | 2016 | [118]    | fecal       | Australian sea lion                   | Not reported                                     | Y                                               | Not discussed                                                                                       |
| Delsuc et al.     | 2013 | [119]    | fecal       | Various myrmecophagous mammals        | Not reported                                     | Y                                               | Not discussed                                                                                       |
| Eigeland et al.   | 2012 | [120]    | fecal       | Dugong                                 | Lower in captivity                               | Y                                               | Not discussed                                                                                       |
| Eisenhofer et al. | 2021 | [35]     | fecal       | Southern hairy-nosed wombat           | Lower in captivity                               | Y                                               | Discussed (metabolic function)                                                                     |
| Frankel et al.    | 2019 | [121]    | fecal       | 5 primate species                     | Lower in captivity                               | Y                                               | Discussed (metabolic function)                                                                     |
| Gao et al.        | 2019 | [87]     | fecal       | Tibetan wild ass                      | Lower in captivity                               | Y                                               | Discussed (immune function)                                                                       |
| Gibson et al.     | 2019 | [38]     | fecal       | Black rhinoceros                      | No difference                                    | Y                                               | Assessed (functional metagenomics—metabolic)                                                          |
| Greene et al.     | 2019 | [20]     | fecal       | Various lemur species                 | Not reported                                     | Y                                               | Discussed (metabolic function)                                                                     |
| Guan et al.       | 2016 | [122]    | fecal       | Sable                                 | Not reported                                     | Y                                               | Discussed (metabolic function)                                                                     |
| Guan et al.       | 2017 | [123]    | fecal       | Sika deer                             | Higher in captivity                               | Y                                               | Discussed (metabolic function)                                                                     |
| Guo et al.        | 2019 | [124]    | fecal       | Giant Panda                           | Lower in captivity                               | Y                                               | Assessed (functional metagenomics—metabolic and immune)                                              |
| Author                | Year | Citation | Sample type | Species                        | Difference in captive/wild microbial diversity? | Difference in captive/wild microbial composition? | Functional implications discussed? |
|-----------------------|------|----------|-------------|--------------------------------|-----------------------------------------------|-------------------------------------------------|-----------------------------------|
| Hale et al.           | 2019 | [125]    | fecal       | Snub-nosed monkey             | Lower in captivity                           | Y                                               | Discussed (metabolic and immune function) |
| Haworth et al.        | 2019 | [126]    | fecal       | Mountain goat                 | No difference                                | Y                                               | Discussed                         |
| Kong et al.           | 2014 | [127]    | fecal       | Red panda                     | Lower in captivity                           | Y                                               | Assessed (associated present OTUs with cellulose degradation ability (GenBank/Kimura—phylogenetic analysis based on 16S)) |
| Li et al.             | 2017 | [86]     | fecal       | Forest musk deer              | No difference                                | Y                                               | Discussed (metabolic function)     |
| McKenzie et al.       | 2017 | [42]     | fecal       | 41 mammal species             | Inconsistent between species                 | Y (except even-toed ungulates)                  | Discussed (metabolic function)     |
| Metcalf et al.        | 2017 | [22]     | fecal       | Przewalski's horse            | Lower in captivity                           | Y                                               | Not discussed                     |
| Milovic et al.        | 2020 | [128]    | fecal       | White-footed mouse            | Lower in captivity                           | Y                                               | Not discussed                     |
| Minich et al.         | 2021 | [78]     | fecal       | White-tailed deer             | Higher in captivity                          | Y                                               | Discussed (metabolic and immune function) |
| Moustafa et al.       | 2021 | [129]    | fecal       | Asian elephant                | No difference                                | Y                                               | Discussed (metabolic function)     |
| Nakamura et al.       | 2011 | [130]    | fecal       | Black howler monkey           | Lower in captivity                           | Y                                               | Discussed (metabolic function)     |
| Narat et al.          | 2020 | [58]     | fecal       | Chimpanzee                    | No difference                                | Y                                               | Discussed (metabolic function)     |
| Narat et al.          | 2020 | [58]     | fecal       | Western lowland gorilla       | Higher in captivity                          | Y                                               | Discussed (metabolic function)     |
| Nelson et al.         | 2012 | [39]     | fecal       | Elephant seal and Leopard seal| Higher in captivity                          | Y                                               | Discussed (immune function)       |
| Ning et al.           | 2020 | [88]     | fecal       | Amur Tiger                    | Higher in captivity                          | Y                                               | Assessed (functional metagenomics—metabolic) |
| Prabhu et al.         | 2020 | [131]    | fecal       | Gaur                          | No difference                                | Y                                               | Assessed (predicted metabolic and immune functions—PICRUSt/KEGG) |
| Rosshart et al.       | 2017 | [79]     | ileocecal    | House mouse                   | Not reported                                  | Y                                               | Assessed (transplant experiment with immune readouts) |
| Schwab et al.         | 2011 | [132]    | fecal       | Grizzly bear                  | Not reported                                  | Y                                               | Discussed (immune function)       |
| Sun et al.            | 2019 | [74]     | fecal       | Alpine musk deer              | Not reported                                  | Y                                               | Assessed (functional metagenomics—metabolic) |
| Sun et al.            | 2019 | [76]     | fecal       | Père David’s deer             | No difference                                | Y                                               | Assessed (predicted metabolic functions—PICRUSt/KEGG) |
| Tang et al.           | 2020 | [133]    | fecal       | Giant panda                   | Lower in captivity                           | Y                                               | Discussed (metabolic and immune function) |
and wild states, and one showed significantly increased diversity in captivity. A recent meta-analysis of 23 comparative studies lends further support to this ambiguity, documenting no systematic effects of captivity on gut microbial diversity [43].

Similar heterogeneity has been observed in comparative analyses of microbial community composition. With few exceptions, comparative studies typically show that overall microbial composition differs significantly between captive and wild populations (Table 1). Yet,

| Author               | Year | Citation | Sample type | Species                                    | Difference in captive/wild microbial diversity? | Difference in captive/wild microbial composition? | Functional implications discussed? |
|----------------------|------|----------|-------------|--------------------------------------------|-------------------------------------------------|--------------------------------------------------|----------------------------------|
| Tsukayama et al.     | 2018 | [40]     | fecal       | Kinda and grayfoot chacma baboon           | Higher in captivity                              | Y                                                | Assessed (functional metagenomics—abx resistance) |
| Uenishi et al.       | 2007 | [134]    | fecal       | Chimpanzee                                  | Not reported                                    | Y                                                | Discussed (metabolic function)   |
| Wasimuddin et al.    | 2017 | [135]    | fecal       | Cheetah                                     | No difference                                  | Y                                                | Assessed (predicted metabolic and immune functions—PICRUSt/KEGG) |
| Xiao et al.          | 2019 | [136]    | fecal       | 6 bat species                               | Higher in captivity                             | Did not compare                                  | Assessed (predicted metabolic functions—PICRUSt/KEGG) |
| Yan et al.           | 2021 | [9]      | fecal       | Pangolin                                     | Higher in captivity                             | Y                                                | Not discussed                    |
| Birds                |      |          |             |                                            |                                                 |                                                  |                                  |
| Oliveira et al.      | 2020 | [37]     | fecal       | Various raptor species                      | No difference                                  | Y                                                | Not discussed                    |
| San Juan et al.      | 2021 | [137]    | fecal       | Brown kiwi                                  | Lower in captivity                              | Y                                                | Not discussed                    |
| Scupham et al.       | 2008 | [138]    | cecal       | Turkey                                      | No difference                                  | Y                                                | Discussed (metabolic function)   |
| Ushida et al.        | 2016 | [6]      | cecal       | Japanese and Svalbard rock ptarmigan        | Not reported                                   | Y                                                | Discussed (metabolic function)   |
| Wiemann et al.       | 2011 | [36]     | cecal       | Capercaillie                                | Lower in captivity                              | Y                                                | Discussed (metabolic function)   |
| Xenoulis et al.      | 2010 | [41]     | cloacal     | 3 parrot species                            | Higher in captivity                             | Y                                                | Not discussed                    |
| Xie et al.           | 2016 | [139]    | fecal       | Red-crowned crane                           | Higher in captivity                             | Y                                                | Discussed (immune function)     |
| Reptiles             |      |          |             |                                            |                                                 |                                                  |                                  |
| Campos et al.        | 2018 | [140]    | fecal and rectal | Green turtle                            | No difference                                  | N                                                | Discussed (metabolic function)   |
| Garcia-De la Pena et al. | 2019 | [141]    | fecal       | Bolson tortoise                             | No difference                                  | Y                                                | Discussed (metabolic function)   |
| Sandri et al.        | 2020 | [142]    | fecal       | Aldabra giant tortoise                      | No difference                                  | Y                                                | Discussed (metabolic function)   |
| Tang et al.          | 2020 | [143]    | fecal       | Crocodile lizard                            | Higher in captivity                             | Y                                                | Assessed (predicted metabolic functions—PICRUSt/KEGG) |
| Amphibians           |      |          |             |                                            |                                                 |                                                  |                                  |
| Tong et al.          | 2019 | [144]    | intestinal  | Dybowski's brown frog                       | No difference                                  | Y                                                | Assessed (predicted metabolic and immune functions—PICRUSt/KEGG) |

Included is whether the paper reported a difference in microbial diversity and microbial composition, as well as whether functional implications of these differences were discussed or assessed. Papers were found with a directed review of existing literature including a Google Scholar search and consulting references cited in each paper collected. We retained only those studies which include gut microbiome measurements of at least one population of captive and one population of wild vertebrates.
there is no apparent consistency across host species as to whether certain bacterial taxa are enriched or depleted in captivity. While it is difficult to compare papers in which only a single host species was sampled because of differences in study design and data analysis, there are a few studies that have looked at microalgal composition in several captive species at once. In primates, two studies have documented a convergence toward human-like microbiomes in captivity [44, 45]. The 41-species comparison by McKenzie et al. documented no convergence and noted variable shifts in the relative abundance of bacterial phyla between host genera [42]. Finally, in a study involving multiple lemur species, Greene et al. observed cases where one or the other of the prominent bacterial genera Bacteroides and Prevotella increased in captivity depending on the dietary strategy of the host [20].

Beyond comparative work, there is limited data tracking the trajectory of microbial change in individual animals moved into captivity from the wild. Here too, though, variable microbial responses to captivity have been observed between host species (Table 2). Findings are inconsistent as to whether gut microbial diversity increases or decreases when animals are brought into captivity as well as how community composition changes. For example, Kohl et al. transplanted two closely related species of woodrats from the wild to captivity and noted striking differences between the species in the magnitude that microbial diversity decreased, as well as opposing trends in the relative abundance of bacteria in the phyla Proteobacteria and Firmicutes [19]. The gut microbiome can respond quickly upon introduction to captivity—for example Kohl and Dearing documented compositional changes in captive woodrats beginning within two weeks of capture [46]—but in some cases wild microbiome characteristics last much longer. A large proportion of the wild microbiome in rodents and lizards is retained in captivity through at least several months [19, 46, 47], and wild mouse microbial signatures can be transmitted through at least 10 generations of captive breeding in a laboratory facility [48].

A few researchers have conducted the inverse transplant experiment, tracking the microbiome as captive animals are reintroduced to the wild (Table 3). These studies provide some evidence that the captive gut microbiome shifts back to resemble the wild gut microbiome after reintroduction and can do so on the timescale of days to weeks [e.g. 49–51]. But, the observed shift back towards a wild microbiome may not completely erase microbial signatures of an animal’s captive origin, as seen in Przewalski’s horses even after 10 years post-release [22]. It is worth noting compositional shifts following reintroduction are likely highly dependent on the environment to which animals are transplanted. In addition to moving animals from captivity to the wild, Chong et al. transplanted Tasmanian Devils from one wild location to another and saw a compositional shift towards a microbiome that resembled native Tasmanian Devils of the new location [50]. The lack of a universal “wild” microbiome, even for a single species, is ubiquitous and logical given findings that some gut bacteria may be geographically restricted or fluctuate seasonally [30, 52–54].

### Table 2 Papers comparing gut microbiota before and after experimental transplantation from the wild to captivity

| Author                 | Year | Citation | Sample type | Species                  | Difference in captive/wild microbial diversity? | Difference in captive/wild microbial composition? | Functional implications discussed? |
|------------------------|------|----------|-------------|--------------------------|-----------------------------------------------|---------------------------------------------------|-----------------------------------|
| Mammals                |      |          |             |                          |                                               |                                                   |                                   |
| Edenborough et al.     | 2020 | [145]    | Fecal       | Angolan freetailed bat    | Higher in captivity                           | Y                                                 | Not discussed                     |
| Kohl and Dearing       | 2014 | [46]     | Fecal       | Desert woodrat           | No difference                                | N                                                 | Not discussed                     |
| Kohl et al.            | 2014 | [19]     | Fecal       | White-throated and Stephen’s woodrat | Lower in captivity                           | Y                                                 | Assessed (metabolic function—monitored ability to digest natural diet) |
| Schmidt et al.         | 2019 | [146]    | Fecal       | Deer mouse               | Lower in captivity                           | Y                                                 | Discussed (metabolic function)   |
| Other                  |      |          |             |                          |                                               |                                                   |                                   |
| Dhanasiri et al.       | 2010 | [147]    | mid and posterior large intestine | Atlantic cod | No difference | Y | Discussed (immune function) |
| Kohl et al.            | 2017 | [47]     | Fecal       | 3 lizard species         | No difference                                | Y                                                 | Discussed (metabolic function)   |

Included is whether the paper reported an increase or decrease in microbial diversity and microbial composition, as well as whether functional implications of these differences were discussed or assessed. Papers were found with a directed review of existing literature including a Google Scholar search and consulting references cited in each paper collected. We retained only those studies which include a wild population brought into captivity within a single lifetime.
What drives gut microbiome changes in captivity?

While it is clear that transitioning between wild and captive conditions can elicit changes to the gut microbiome, the ecological factors altered under captivity that specifically cause microbiome shifts often remain unspecifed. Captive diets can vary widely from natural diets [3, 55], and diet has been shown to have a strong influence over gut microbial composition in captive animals [21, 31, 51, 56]. Even minor changes in diet can lead to shifts in microbial composition, as evidenced in captive sifakas [57]. Environmental conditions beyond diet can also matter, with one study demonstrating that rehousing lab mice in a room with altered temperature and humidity, but the same diet, resulted in significant changes in gut microbial composition [49]. Additionally, microbial dispersal from abiotic environmental substrates (water, soil, plants, etc.), as well as from contact with conspecifics or other species (including humans), can influence the gut microbiome of captive animals and can vary between captive facilities [58]. Komodo dragons, for example, harbor gut microbes that are also present in their enclosures [59], while human handling has been implicated in the spread of microbes between mouse enclosures [48]. Animal social groups have a strong influence on gut microbial composition in the wild, likely affecting the microbiome by transferring bacteria between individuals through physical contact among social networks [e.g. 60–62]. It is reasonable to expect similar social dynamics transmit microbes among captive populations, although, beyond studies of cohousing [e.g. 63, 64], this is rarely explicitly considered.

It must be noted that diet and environmental exposures can be tightly linked, especially in wild environments, making it challenging to disentangle their microbial impacts. Diet has the potential to have both a probiotic and prebiotic influence on the microbiome as it exposes animals to environmental microbes in addition to providing direct substrates for bacterial growth [52, 65]. Animal diets also often vary as environmental conditions change. One study, for example, saw a strong effect of the amount of outdoor exposure on changes to the gut microbiome of colobine primates, but it could not be determined whether this was a result of the animals being exposed to new environmental microbes or eating a greater diversity of foliage [55].

Interaction with humans introduces additional stressors unique to the captive environment that can influence the microbiome. Most notably, administration of antibiotics reduces the natural flora of the gut and may even compound the effects of captivity by quickly removing large portions of the wild microbiota [66]. Isolation from natural pathogens like helminths through anthelmintic medicine or increased hygiene can induce changes to the microbiome by affecting the immune system [67, 68]. Additionally, direct physical interactions with humans, such as keepers or visitors, could both directly introduce human-associated microbes [44, 45] and cause stress, which itself can induce shifts in microbial composition [e.g. 69, 70].

Do gut microbial compositional changes in captivity affect microbiome function or host health? If so, how?

While the reviewed studies clearly show that captivity can affect the composition and diversity of the gut microbiome, characterizing these two metrics is insufficient to determine whether the gut microbiome plays a role in captive animal health as they do not give insight into whether bacterial functions change [71]. To date, there

Table 3 Papers comparing gut microbiota before and after experimental transplantation from captivity to the wild

| Author               | Year | Citation | Sample type | Species           | Difference in captive/wild microbial diversity? | Difference in captive/wild microbial composition? | Functional implications discussed? |
|----------------------|------|----------|-------------|-------------------|------------------------------------------------|-------------------------------------------------|-----------------------------------|
| Mammals              |      |          |             |                   |                                                 |                                                 |                                    |
| Bar et al. 2020 [49] |      |          | Fecal       | House mouse       | Lower in captivity                             | Y                                               | Discussed (immune function)       |
| Chong et al. 2019 [50]|      |          | Fecal       | Tasmanian devil   | Lower in captivity                             | Y                                               | Discussed (metabolic function)    |
| Leeuwen et al. 2020  |      |          | Fecal       | Deer mouse       | No difference                                  | Y                                               | Discussed (neurological function) |
| Schmidt et al. 2019  |      |          | Fecal       | Deer mouse       | Lower in captivity                             | Y                                               | Discussed (metabolic function)    |
| Yao et al. 2019 [75] |      |          | Fecal       | Giant panda      | Lower in captivity                             | Y                                               | Assessed (functional metagenomics) |

Included is whether the paper reported an increase or decrease in microbial diversity and microbial composition, as well as whether functional implications of these differences were discussed or assessed. Papers were found with a directed review of existing literature including a Google Scholar search and consulting references cited in each paper collected. We retained only those studies which include a captive population released into the wild within a single lifetime.
have been few studies that compare functional capacity of the microbiome in the wild and in captivity in addition to composition. Of these, most have predicted functionality using reference genomes related to taxa represented in 16S rRNA gene sequencing data [e.g. 44, 72, 73] and a few have conducted functional metagenomic sequencing [e.g. 38, 40, 74, 75]. Most show that predicted function differs between captive and wild microbiomes (but see [76] for a case in which no functional difference was predicted) and argue the altered functional capacity of the bacterial community may have a realized impact on host biology. However, altered functional capacity does not necessarily indicate changes in bacterial activity or host physiology [77].

There have been some instances, though, where microbial composition has been more directly tied to health outcome in captive animals. A handful of studies have correlated gut microbiome composition to specific disease phenotypes in captive animals, including chronic wasting disease in deer [78], gastrointestinal disease in wolves [31], and iron overload disorder in rhinoceroses [7]. That work provides support for a functional connection between microbial composition and health consequences in at least some contexts. Bolstering the idea that the wild microbiome specifically can be beneficial, fecal microbiota transplantation (FMT) of a wild microbiota community has been used in mammals to successfully reduce inflammation and improve survival against viral disease [79] and expand dietary niche [80], and probiotic supplementation improved body weight and fecal quality in cheetahs [81]. In most cases, however, the potential for microbial manipulations to lead to favorable health outcomes have not been explicitly tested, so the generalizability of these findings remains unknown. Microbial manipulations in captive livestock have also been used to try to address health concerns, although the mixed success of FMTs and dietary interventions in agricultural contexts underscores the likelihood that an altered microbiota will not cure all ills [82–85].

There are alternative interpretations of microbial differences between wild and captive animals that do not presume the captive microbiome is less favorable [e.g. 35, 86–88] or potentially detrimental [1]; these are important to consider in the absence of species-specific experimental evidence. First, microbiomes may have changed to match the new captive environment in a beneficial manner [89]. For example, natural selection could promote a microbiome better able to digest the captive diet while potentially reducing the animal’s capacity to digest a (no longer relevant) wild diet [21, 90]. Captive microbiome contributions to promoting growth or mitigating stressors may not fully outweigh the costs of a captive lifestyle, so captive animals may still be found to have worse health than wild individuals. Still, in these cases a return of a wild or ancestral microbiome to a captive animal would further diminish health in captivity. Second, microbiome changes under captivity may be constrained by tradeoffs such that the increase in some microbial functions, such as those relevant to captive environments, lead to the decrease or loss of other functions [77]. If so, re-establishing a wild microbiome may improve some aspects of health while reducing others. Third, the captive microbiome may be caused by, rather than causative of, poor health, in which case treating the microbiome would not alter the underlying condition. Finally, changes in the microbiome under captivity may primarily be neutral given functional redundancy of gut microbiota [91]. Given the scope of the current evidence that exists, we cannot presume to know what effect the captive microbiome has on the health of any given species. To ensure our understanding is sufficient for conservation applications, more research must be done to indicate under which circumstances and for which species microbial manipulations can improve health.

Future directions

There is no single answer to the question of whether the gut microbiome matters for host health in captivity in large part because there is no single captive microbiome. The microbiome’s impact will be particular to host species, diet, external environment, conservation status, and reintroduction plans, among many other factors that impact the gut microbiome and/or affect species health priorities. Insofar as there is a prospect of health-relevant compositional and functional differences of the gut microbiome under captivity, we must identify when reconstituting a wild microbiome is most likely to be a suitable route for improving animal welfare and conservation. (Although, even absent demonstrated health benefits, there may be cases where reconstituting a wild microbiome is a worthwhile goal, for instance where it provides more ecologically realistic models for host-microbe interaction research.)

Responsible applications of gut microbiome interventions to animal conservation practices will depend on knowing whether the captive microbiome is truly involved in modulating a given health concern. The gold standard causal evidence requires experimental microbiome manipulations in captive animals: changing the microbiome either through fecal transplants, antibiotics, or probiotics then measuring host phenotypes of interest to assess whether existing health problems are ameliorated [28, 79–81, 92]. Similar experimental tests could be employed to assess whether microbial manipulations prior to reintroduction improve survival in the natural environment. Of course, experiments for observing
Direct causality have many potential ethical and logistical limitations, particularly if concerning endangered species, so may not be possible in many cases.

Prior to proposing microbiome manipulations (either for experimental research or treatment purposes), alternate studies which go beyond documenting microbiome differences but not as far as microbial treatment could provide better corroboration of captivity-mediated microbiome health effects. Altering captive conditions to make the microbiome more wild-like (e.g., restoring diet complexity, increasing exposure to other individuals or species, diversifying environmental substrates, limiting handling; see Fig. 1) then tracking microbial and host health responses could identify beneficial wild-associated taxa or other critical ecological drivers of the captive microbiome.

If experiments in the species of interest are not possible, alternatives include gnotobiotic or in vitro research. In gnotobiotic studies, gut microbiota from different donors, such as wild and captive individuals, are transplanted into germ-free animals (often mice) whose phenotypes are then compared [93, 94]. These experiments have their own limitations including expense and limited facilities, as well as the reduced biological realism that comes from measuring a microbiome’s effect in a mismatched host [95], but they could be particularly useful for studying the microbiome of endangered species.

Similarly, in vitro culturing experiments could provide insight into microbial functioning of wild and captive gut microbiomes when live animal work is infeasible. Given that in vitro studies remove host biology, they cannot be used to relate gut microbial composition to host phenotype. However, they could allow researchers to characterize whole microbial communities under varying stimuli and measure functional responses relevant to host biology in a relatively cheap and more high-throughput manner [96, 97]. They may prove especially useful for testing potential manipulations (e.g., diet, microbial exposure) for promoting a shift from a captive to more wild-like microbiota.

More comprehensive observational studies that tie microbiome composition to host phenotype data in wild and captive settings could prioritize host species or health conditions which warrant further study [e.g. 7, 9, 31, 78]. Correlations between health status and the microbiome would not distinguish between cases where the captive microbiome was responsible for altered health and ones where it is a response to the unhealthy state or is curbing potentially worse states. They would, however, advance the field beyond characterizing the microbiome in wild and captive states and serve for hypothesis-generation to motivate future experimental work. Correlative studies have the benefit of being less invasive and lower cost than direct manipulations but do require a robust sample set.

**Fig. 1** Possible interventions for manipulating the captive animal microbiome. A Direct animal manipulations include administration of antibiotics, microbiome transplants, and physical handling. B Manipulations targeting environmental exposures include presence of other animals (other species, conspecifics, and recently-wild individuals), diet and food processing, water source, cleaning practices, visitor exposure, climate, and soil, plants, and their associated microbes. Icons have been adapted under a Creative Commons license (https://creativecommons.org/licenses/by/3.0/) at phylopic.org and thenounproject.com. The hand image was sourced from clipart-library.com under a Personal Use license.
In addition to collecting new data, researchers could leverage banked fecal samples and existing zoo or sanctuary records or larger databases like the Zoological Information Management System (ZIMS), Animal Records Keeping System (ARKS), Single Population Animal Records Keeping System (SPARKS), and Medical Animal Records Keeping System (MedARKS) [98].

Regardless of the approach chosen for a future study, choosing appropriate phenotypic markers to assess host health will be critical. Previous research on microbiome-mediated health impacts has primarily been conducted in animal models or humans [e.g. 99–101], so common markers and their benchmarks may not be entirely applicable to wild species or wild contexts (but see [102, 103] for biomarkers that may be applicable to wild animals). While transcriptomic or metabolomic data are frequently collected in biomedical research and can illuminate molecular mechanisms [77], data at that scale is often unnecessary for conservation work. Given the high cost and technical demands of “omics” measurements, alternatives such as stool consistency, parasitology, body size, or physical condition may be more appropriate. Indeed, some of these have been previously used to measure health phenotypes in wild or captive species [e.g. 31, 104–106], although not always with explicit ties to the microbiome. Pathological biomarkers targeted towards a particular health concern, for instance diagnosing or determining the severity of gastrointestinal issues, infection status, or neurological and stress disorders [e.g. 9, 107, 108], could be more informative but potentially more invasive or costly. Ultimately, markers should be chosen based on the species and health concern of focus, as well as the laboratory capabilities available. Furthermore, any research program studying the captive gut microbiota should be diligent about considering multiple axes of animal health when possible given the possibility for functional tradeoffs resulting from microbial compositional shifts [77].

Deciding how to implement these research strategies aimed at causation over correlation in studies of the captive gut microbiota will be highly dependent on logistical and scientific factors specific to a given research program. Limited resources, including access to animals, may come up against conservation priorities and severely restrict researchers’ choices. Fortunately, the methods presented above offer a wide range of options that may allow scientists to do functionally relevant research on the captive animal microbiome even when economic or logistical limitations preclude direct microbial manipulation experiments. Experts on specific animals will be best suited to decide which research strategies and phenotypic readouts will be most useful and feasible for achieving pressing conservation goals for the target species, as it is clear now that results will not be generalizable between taxa.

Once health links to the gut microbiome are identified for a given species, there are additional challenges in deciding whether and how to incorporate microbial manipulations into husbandry practices as this would require investments of both funding and labor at animal facilities. Again, species experts will be in the best position to balance wellness and conservation goals with the costs and challenges of microbial manipulations—particularly if functional tradeoffs are present. For example, if the captive microbiome proves beneficial in captivity but harmful when animals are reintroduced, the short-term wellbeing of an individual animal must be weighed against the long-term wellbeing of the species. To successfully reintroduce animals to the wild may require pre-seeding a wild microbiome, or pre-seeding may be an unnecessary expense if functionality recovers when the microbiome reverts to a more wild-like state on its own [22, 49–51]. It is likely that certain manipulations will be more appropriate or more costly for some species than others, and institutions such as zoos that manage a wide variety of species will likely need to employ a wide range of strategies to cope with this variation. Despite these challenges, the potential benefits of harnessing the influence of the gut microbiome to improve host health is undeniable and will certainly be a worthwhile research avenue for many species.

Conclusion
As more species are becoming impacted by threats such as climate change, human land use, and emerging infectious diseases [109–112], finding new ways to improve animal welfare in captivity may be critical for species success through captive breeding and reintroduction programs. Much work has been done in this century to characterize the differences between gut microbial composition of captive animals and their wild counterparts. As a whole, it is clear that there is enormous variation in how species’ gut microbes respond to the captive environment. Thus, there will likely be variation in how the captive microbiome influences host health, preventing a one-size-fits-all approach to managing animal health through the microbiome. Both captive microbiome research and eventual mitigation strategies will likely need to be handled on a species-by-species basis managed by those who are most familiar with their particular biology, but in all cases they should include broader surveys of health phenotypes that consider immune, behavioral, and reproductive health in addition to metabolic health.

This complexity raises important questions for how the field should move forward, such as what species
should be prioritized for research and what aspects of host health should be optimized. For example, should we begin with species that are known to have poor health in captivity or with those that face the greatest extinction threats? And is it worth improving metabolic health in captivity if it comes at the cost of immune health or metabolic functioning in the wild? The answers to these questions will likely be influenced by logistical barriers such as species availability and population sizes, but they merit careful consideration by scientists continuing captive microbiome research. As the field moves away from characterizing microbial composition and focuses more on linking microbial function to host phenotypes, it is crucial that researchers not only consider these questions, but acknowledge the limitations of their findings to avoid recommending widespread microbial manipulations before their effects are fully understood.

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