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FORUM

Interpretation of gut microbiota data in the ‘eye of the beholder’: A commentary and re-evaluation of data from ‘Impacts of radiation exposure on the bacterial and fungal microbiome of small mammals in the Chernobyl Exclusion Zone’

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Abstract
1. Evidence that exposure to environmental pollutants can alter the gut microbiota composition of wildlife includes studies of rodents exposed to radionuclides.
2. Antwis et al. (2021) used amplicon sequencing to characterise the gut microbiota of four species of rodent (Myodes glareolus, Apodemus agrarius, A. flavicollis and A. sylvaticus) inhabiting the Chernobyl Exclusion Zone (CEZ) to examine possible changes in gut bacteria (microbiota) and gut fungi (mycobiota) associated with exposure to radionuclides and whether the sample type (from caecum or faeces) affected the analysis.
3. The conclusions derived from the analyses of gut mycobiota are based on data that represent a mixture of ingested fungi (e.g. edible macrofungi, polypores, lichens and ectomycorrhizae) and gut mycobiota (e.g. microfungi and yeasts), which mask the patterns of inter- and intraspecific variation in the authentic gut mycobiota.
4. Implying that ‘faecal samples are not an accurate indicator of gut composition’ creates an unnecessary controversy about faecal sampling because the comparison of samples from the caecum and faeces confounds many other possible drivers (including different animals from different locations, sampled in different years) of variation in gut microbiota.
5. It is relevant also that Antwis et al.’s (2021) data lack statistical power to detect an effect of exposure to radionuclides on the gut microbiota because (1) all of their samples of Apodemus mice had experienced a medium or high total absorbed dose rate and (2) they did not collect samples of bank voles (M. glareolus) from replicate contaminated and uncontaminated locations.
6. Discussion of Antwis et al.’s (2021) analysis, especially the claims presented in the Abstract, is important to prevent controversy about the outcome of research on the biological impacts of wildlife inhabiting the CEZ.
INTRODUCTION

Animal gut microbial communities provide essential services for their host, such as interacting with the host's immune system (Pickard et al., 2017; Round & Mazmanian, 2009), defending against pathogen invasion (Rosshart et al., 2019; Pickard et al., 2017) and processing dietary material to provide important metabolites (Morrison & Preston, 2016; Sonnenburg & Bäckhed, 2016). As disruption to the gut microbiota can affect the health of the host, there is much interest in identifying features of the host or its environment that can impact the gut microbiota community composition (Zaneveld et al., 2017). Infection by parasites or pathogens (Kreisinger et al., 2015; Sabey et al., 2021), the level of biodiversity or habitat disturbance (Barelli et al., 2020), changes in diet and season (Guo et al., 2021; Lavrinienko et al., 2020; Maurice et al., 2015) and exposure to pollutants (Brila et al., 2021) are associated with a change in the gut microbiota of wildlife.

Evidence that exposure to pollutants impacts the gut microbiota extends to rodents inhabiting areas contaminated by radionuclides (Lavrinienko et al., 2020; Lavrinienko, Mappes, et al., 2018; Lavrinienko, Tukalenko, et al., 2018). Antwis et al. (2021) characterised the gut microbiota of four species of rodent inhabiting the Chernobyl Exclusion Zone (CEZ), a region surrounding the former nuclear power plant at Chernobyl, Ukraine, where substantial areas are contaminated by radionuclides. Samples were collected from bank voles (Myodes glareolus), where animals were caught from locations that represented a gradient of contamination, and from three species of mice (Apodemus agrarius, A. flavicollis and A. sylvaticus), where animals had experienced ‘medium’ (4–42 μGy/hr) or ‘high’ (>42 μGy/hr) absorbed dose rates. The CEZ was established to limit human exposure to radionuclides, but the wildlife inhabiting the CEZ provide the best-studied models of the biological impacts of exposure to environmental radionuclides (Beresford et al., 2016; Møller & Mousseau, 2006; Mousseau, 2021). Despite decades of research on wildlife within the CEZ, the effects of exposure to environmental radionuclides in wildlife remain a source of controversy (Beresford et al., 2016; Beresford, Horemans, et al., 2018; Kesäniemi et al., 2018; Møller & Mousseau, 2006; Mousseau, 2021). In their Abstract, Antwis et al. (2021) conclude

1a) ‘We provide novel evidence that host species determines fungal community composition’,
1b) ‘... fungi (Steccherinaceae and Strophariaceae) in the guts of bank voles that may serve as biomarkers of radiation exposure’,
(2) ‘...considerable variation between results for faecal and gut samples of bank voles, suggesting faecal samples are not an accurate indicator of gut composition’ and
(3) ‘associations between radiation exposure and microbiome composition of gut samples were not robust against geographical variation’.

These points have clear implications for our understanding of the effects of environmental radionuclides on wild animal gut microbiota, and for studies of exposure to pollutants on wildlife microbiota generally. However, statements (1a, b) need clarification as the fungal amplicon sequence data represent a mixture of ingested fungal material and authentic gut mycobiota. Conclusion (2) creates an unnecessary controversy as it neglects to account for confounding effects of multiple drivers of variation in the gut microbiota in the comparison of samples from the caecum and faeces. We discuss statement (3) to highlight the need for greater clarity about the power of Antwis et al.’s (2021) study design to identify the effects of exposure to radionuclides on the gut microbiota of small rodents.

MATERIALS AND METHODS

Antwis et al. (2021) used amplicon sequencing to characterise the bacterial and fungal components of the gut microbiota in four species of rodent (bank vole M. glareolus, wood mouse Apodemus sylvaticus, striped field mouse A. agrarius and yellow-necked mouse A. flavicollis) inhabiting the CEZ, Ukraine. Samples were collected by live trapping from two areas: (1) a contaminated site in the Red Forest and adjacent area, and (2) an uncontaminated site about 10 km south-west of the Red Forest (Figure 1). We re-analysed some of the amplicon sequence data used by Antwis et al. (2021) to examine the potential proportion of non-resident fungal sequence variants (SVs) in their data. As the data used in this article were downloaded from a public archive, we did not seek permissions for fieldwork or ethical approval for the work.

Full details about the sample data are provided in the original publication (Antwis et al., 2021). Briefly, data were obtained from GenBank (PRJNA594002) and processed in QIIME2 v.2020.6 (Bolyen et al., 2019), using CUTADAPT (https://github.com/marcelm/cutadapt) to remove primer/adaptor sequences and DADA2 (Callahan et al., 2016) to denoise the data. Taxonomy for SVs was assigned using the SKLEARN machine learning taxonomy classifier (Bokulich et al., 2018) against the UNITE v.8 (Nilsson, Larsson, et al., 2019) reference database. Because there are no reliable data that identify all species of fungi that could be ingested (either by direct consumption, or by association with other components of the diet) by bank voles.
and *Apodemus* mice in the CEZ, we used informed filtering to identify likely non-resident fungal SVs (see Lavrinienko, Scholier, et al., 2021). SVs were assigned to the major classes of microfungi using the information at the Microfungi Collections Consortium (www.microfungi.org/table1). We further classified fungal SVs according to guild or growth form based on assignments made by FUNGUILD v.1.2 (Nguyen et al., 2015). SVs categorised as plant pathogens, epi- and endophytes, lichens, mycorrhizae and wood saprophytes were assumed to be part of the ingested, non-resident fungal material in the gut, as were SVs with large fruiting bodies/growth forms (e.g. as gasteroid, pezizoid, tremelloid, etc.), which left the remaining microfungi and yeasts (and taxa with unclear growth forms but which were assigned as animal pathogens or SVs that lacked information about guild, e.g. because taxonomic resolution was not assigned below Phylum level) as candidate resident gut mycobiota. Thus, we make a contrast between data that can represent dietary items (principally macrofungi and lichens, plant-associated fungal pathogens, mycorrhizae or endophytes) and the remaining data as a candidate resident gut mycobiota (many microfungi and yeasts, taxa associated with animals and poorly known fungi).

Data were imported into PHYLOSEQ (McMurdie & Holmes, 2013) for analyses in R v.4.0.5 (R Core Team, 2020). Fungal data were rarefied to an even depth of 5,000 (all SV data, that resulted in a loss of 11 samples and 1,020 SVs), 3,000 (possible resident gut fungi, with a loss of 27 samples and 980 SVs) or 1,000 (possible dietary and diet-associated fungi, with a loss of 21 samples and 710 SVs) reads per sample. We calculated alpha diversity (observed number of SVs) and beta diversity (Bray–Curtis dissimilarity) in PHYLOSEQ. Variation in alpha diversity was assessed using pairwise Wilcoxon rank-sum tests with Holm correction. The ADONIS2 function in VEGAN (Oksanen et al., 2020) was used to examine the amount of variation in beta diversity explained by features of the data, such as host species (bank vole, wood mouse, striped field mouse and yellow-necked mouse), sampling year (2017, 2018) and total absorbed dose rate of radiation (μGy/hr). BETADISP function in VEGAN (Oksanen et al., 2020) was used to determine whether there were significant differences in dispersion among groups of samples.
3 | RESULTS

3.1 | Composition of the community of fungi detected in rodent guts

By classifying probable ingested fungi as macrofungi, plant pathogens, endophytic species and taxa associated with decaying wood, we separated the fungal SVs in rodent gut samples into 2,608 putative gut residents and 1,566 possible non-residents. This filtering procedure indicates that about a third of the read data could be derived from ingested fungi (Figure 2), with about 10%–15% of the reads assigned to macrofungi (e.g. Agaricomycetes).

The inclusion of all fungal SV data affects the analyses. Although interspecific differences in alpha diversity are apparent in unfiltered and filtered datasets, with bank voles having significantly fewer fungal SVs than species of Apodemus (pairwise Wilcoxon rank-sum test with Holm correction, $p < 0.05$ for all comparisons; Table 1, Figure 3), the inclusion of dietary material inflates the estimates of alpha diversity (Figure 3). Filtering the fungal SVs also impacts the pattern of beta diversity. Without SV filtering, bank voles differ from the three species of Apodemus mice, with samples from wood mice being somewhat different to those from striped field mice and yellow-necked mice (Figure 4a). There appears to be less interspecific variation in the gut mycobiota (Figure 4b) than among the ingested fraction of fungi, which is characterised by high inter-individual variation among samples of bank voles and little inter-individual variation among the samples of wood mice (Figure 4c). Filtering the SVs affects the explanatory power of ‘host species’ with slightly less interspecific variation among the samples of possible resident gut fungi (cf. values of $R^2$ in Table 2). Moreover, significant ($p = 0.001$) interspecific differences in the amount of dispersion were detected in the analyses based on the entire dataset and the suspected non-resident fungi, but not in the subset of possible resident gut fungi (Table 2, Figure 4).

![Figure 2](image)

**Figure 2.** Proportions of fungal classes identified in the gut and faecal samples from four species of rodent, separated by their possible resident (mycobiota) or non-resident (ingested) status in the host’s gastrointestinal tract.

| Class                          | Bank vole | Striped field mouse | Wood mouse |
|-------------------------------|-----------|---------------------|------------|
| (a) All fungi                 |           |                     |            |
| Striped field mouse           | 2.8e-05   |                     |            |
| Wood mouse                    | 3.9e-06   | 0.60                |            |
| Yellow-necked mouse           | 3.1e-11   | 0.60                | 0.68       |
| (b) Mycobiota                 |           |                     |            |
| Striped field mouse           | 0.0039    |                     |            |
| Wood mouse                    | 0.0018    | 1.00                |            |
| Yellow-necked mouse           | 3.2e-06   | 1.00                | 1.00       |
| (c) Ingested fungi            |           |                     |            |
| Striped field mouse           | 2.4e-06   |                     |            |
| Wood mouse                    | 2.3e-08   | 0.12                |            |
| Yellow-necked mouse           | 1.1e-15   | 0.10                | 0.94       |
4 | DISCUSSION

4.1 | Composition of the community of fungi detected in rodent guts

With our understanding of the processes that affect wild animal gut fungi (gut mycobiota) limited (Huseyin et al., 2017; Kong & Morris, 2017; Nilsson, Anslan, et al., 2019), it is important to carefully consider the possible sources of fungal material in samples from the animal gut (Lavrinienko, Scholier, et al., 2021). As amplicon sequencing enumerates all types of DNA (for the target region and taxon, such as a partial region of the 16S rRNA for bacteria or the ITS for fungi; Knight et al., 2018; Lavrinienko, Jernfors, et al., 2021) within a sample, the resulting SVs are derived from the authentic gut residents and any non-resident (ingested) material. A combination of (1) comparatively few fungal cells (compared with bacterial cells) in the vertebrate gut (Qin et al., 2010, Iliev et al., 2012) and (2) ingestion of fungi by many animals, for example by consumption of macrofungi or lichens (Abt & Bock, 1998; Fogel & Trappe, 1978) or intake of fungal plant pathogens, commensals/symbionts or the microfungi in fermenting or decaying material, raises the potential that amplicon sequencing-based studies of gut mycobiota will contain a substantial amount of non-resident gut fungi (Lavrinienko, Scholier, et al., 2021).

Antwis et al. (2021) did not differentiate among resident gut mycobiota or ingested fungal material. Evidence for an abundant fraction of non-resident fungal material in Antwis et al.’s (2021) analysis of ‘gut mycobiota’ is derived from their potential ‘biomarkers of radiation exposure’: the Steccherinaceae (members of the Polyporales that are a cause of white rot, often growing on wood) and the Strophariaceae (saprophytes within the Agaricales). As macrofungi associated with decaying material, these taxa are...
unlikely to be long-term residents of the mammalian gut. Our re-
analysis of Antwis et al.’s (2021) data identified a substantial frac-
tion of likely dietary fungi in their samples, including members of
the Agaricomycetes that are often actively selected for ingestion by
rodents (Abt & Bock, 1998; Fogel & Trappe, 1978). By re-analysing
Antwis et al.’s (2021) data, we show how (a lack of) filtering the
fungal SVs can impact the outcomes of analyses of gut mycobiota
(via alpha diversity, beta diversity and dispersion) and hence the
study conclusions. For example, there is a clear possibility that ra-
diation exposure impacts the diversity of fungi or other food items
in the environment that, in turn, affects the host’s (dietary) intake
of non-resident gut fungi, in addition, or alternate, to, any direct
effect of the environment on the rodent gut mycobiota per se.
Amplicon sequencing-based analyses of wildlife gut mycobiota thus
require an assessment of the ecology of host(s) and fungi to iden-
tify the possible source(s) of fungal SVs and make better informed
conclusions about ecological and evolutionary drivers of inter- and
intraspecific differences in gut mycobiota (Lavrinienko, Schollier,
et al., 2021).

4.2 Differences between microbiota detected in
samples from faeces and the caecum

The extent to which sample type or laboratory procedures impact
amplicon sequence data is an important issue for microbiota re-
search (Knight et al., 2018; Panek et al., 2018). By describing a cae-
cum sample as providing ‘...data on the true gut microbiome,’ Antwis
et al. (2021) create an artificial impression about data quality in mi-
crobiota studies as one may define, for example, a community of
microbes from the stomach, small intestine, caecum, colon and/or
faeces (Kreisinger et al., 2015; Martinez-Guryn et al., 2019; Videvall
et al., 2018), and make a distinction between the lumen and the gut
mucosa (Albenberg et al., 2014). Alternative sampling locations of
the gut microbiota are neither ‘true’ nor ‘false’ but are different rep-
resentations of natural variation in microbial assemblages that ac-
company the process of digestion and changes in the environment
(e.g. pH, oxygen tension, flow rate, etc., Martinez-Guryn et al., 2019)
along the gastrointestinal tract. While the composition of mi-
crobes in faecal samples often (Ingala et al., 2018; Velasco-Galilea
et al., 2018; Videvall et al., 2018; Zhou et al., 2020) differs from the
microbes identified from regions of the digestive tract, the com-
parison itself depends on the regions of the gastrointestinal tract
under consideration; for instance, faeces may recapitulate the mi-
crobiota of the colon, but not ileum or caecum (Videvall et al., 2018).
Moreover, when conducted with great care (e.g. standardising the
time between defecation and sample collection, and using consist-
ent and appropriate conditions to store the samples), faecal sampling
is an invaluable, non-destructive method of obtaining a representa-
tive sample of the gut microbiota that allows (and is a unique option
for) longitudinal sampling (Antwis et al., 2019; Johnson et al., 2019;
Lavrinienko et al., 2020).

The relevant issue about sample type is whether it interacts with
the treatment to produce a biased outcome. Antwis et al. (2021)
state ‘For bank voles, we observed differences in microbial commu-
nities associated with the gut and faeces, ... We also observed sig-
ificant differences in the relationships between radiation and gut/
faecal microbial families’. Unfortunately, Antwis et al. (2021) did not
use an appropriate study design that compares paired samples of
faeces and caecum from the same individuals collected at a simi-
lar time (cf. Ingala et al., 2018; Velasco-Galilea et al., 2018; Videvall
et al., 2018; Zhou et al., 2020). Rather, the variation in microbiota
that Antwis et al. attribute to ‘sample type’ (faeces or caecum) is con-
founded with variation derived from samples being collected from
(1) different animals, (2) different years, (3) different time points
(July/August versus August) and (4) different locations and hence

| TABLE 2 Effect of filtering fungal sequence variants (SVs) by their traits on the level of variation in beta diversity and dispersion in amplicon sequence data from samples of caecum and faeces from four species of rodent. (a) All data, (b) the likely resident gut fungi (mycobiota) and (c) the possible fungal SVs that were ingested as part of the host’s diet (non-resident fungi) |
|---|---|---|---|---|---|---|---|
| Beta diversity | | | | | | |
| | df | SS | R² | F | p |
| (a) All fungi | | | | | | |
| Species | 3 | 6.634 | 0.072 | 6.160 | 0.001 |
| Residual | 237 | 85.078 | 0.928 | | |
| Total | 240 | 91.712 | 1.000 | | |
| (b) Mycobiota | | | | | | |
| Species | 3 | 5.695 | 0.070 | 5.512 | 0.001 |
| Residual | 221 | 76.117 | 0.930 | | |
| Total | 224 | 81.812 | 1.000 | | |
| (c) Ingested fungi | | | | | | |
| Species | 3 | 7.745 | 0.088 | 7.264 | 0.001 |
| Residual | 227 | 80.676 | 0.912 | | |
| Total | 230 | 88.421 | 1.000 | | |
| Dispersion | | | | | | |
| | df | SS | R² | F | p |
| Species | 3 | 0.092 | 0.031 | 6.644 | 0.001 |
| Residual | 237 | 1.089 | 0.005 | | |
4.3 Associations between radiation exposure and gut microbiota composition

Antwis et al. (2021) note that associations between gut microbiota composition were not robust when the analyses were controlled for geographic distance, with sampling site explaining some variation in bacterial beta diversity. Hence, they conclude that ‘any variation in microbiome composition arising from proximity to the Chernobyl Nuclear Power Plant is more likely a habitat effect than a result of radiation exposure’. Because bank voles can disperse several kilometres per year (White et al., 2012), Antwis et al.’s (2021) animals with high absorbed dose rates represent samples from one cohort of animals within a single contaminated area (the Red Forest locality) rather than from separate cohorts inhabiting the different contaminated sites within the CEZ (see Figure 1). A lack of replication of sites with similar dose rate categories is a curious design for a study of radiation effects on wildlife, especially when the focal contaminated area is the Red Forest as this location is argued to comprise poor habitat (Beresford, Scott, et al., 2020). Studies of radiation effects that do not employ a replicated study-site design confound the treatment (radiation exposure) with location and thus are somewhat destined not to support the idea that exposure to radionuclides has no detrimental biological impacts: a lack of statistical effect can be interpreted that radiation exposure has little biological impact, while any apparent biological impacts can be dismissed as location-specific effects (e.g. the poor-quality habitat in the Red Forest) rather than exposure to radionuclides! Only by collecting samples from replicate contaminated and uncontaminated areas can location-specific effects be partitioned from impacts associated with radionuclide exposure (Jernfors et al., 2021; Kesäniemi, Jernfors, et al., 2019; Kesäniemi, Lavrinienko, et al., 2019). The additional discussion points raised by Antwis et al. (2021) that ‘Other studies of radiation effects in CEZ wildlife, including the microbiome studies of Lavrinienko et al. ..., also have their most contaminated sampling sites within the Red Forest...’ and ‘Any study that uses the Red Forest as a location for radiation effect studies on wildlife needs to consider the historical impacts of radiation and other stressors (e.g. wildfires) on this area...’ are somewhat misleading as they neglect to consider the consistent patterns associated with radionuclide exposure in gut microbiota samples from replicate contaminated and uncontaminated locations (Lavrinienko, Mappes, et al., 2018; Lavrinienko, et al., 2020). It is also relevant that Antwis et al.’s (2021) samples from Apodemus mice were derived from animals that had absorbed dose rates of \( \geq 20 \mu\text{Gy/hr} \); in fact, the data from 2017 consist almost exclusively of one category of absorbed dose rate (‘high’), and yet the authors included it in analyses as categorical predictor of three levels (0, 14 and 100 Apodemus in the ‘low’, ‘medium’ and ‘high’ dose rate categories respectively). The analysis by Antwis et al. (2021) potentially examines the response of Apodemus mice to medium and high absorbed dose rates but does not examine how exposure to radionuclides per se impacts the gut microbiota in different species of rodent. By contrast, comparable impacts of exposure to radionuclides in the gut microbiota of three (out of four) species of Apodemus are apparent in samples collected from the CEZ and the Fukushima accident sites (Lavrinienko, Hämäläinen, et al., 2021).

5 CONCLUSIONS

The literature about biological impacts of inhabiting the area surrounding the former nuclear power plant at Chernobyl is littered with arguments and counterarguments (Beresford et al., 2016; Beresford, Horemans, et al., 2020; Beresford, Scott, et al., 2020; Mousseau, 2021). Some of this discussion might be derived from simple misunderstandings in communication. For instance, Antwis et al. (2021) write ‘we present the first study of gastrointestinal (GI) tract microbiome composition in CEZ small mammals for which individual total absorbed dose rates have been estimated’. However, this statement either refers specifically to the analysis of samples from the caecum or they overlooked that previous studies of wildlife gut microbiota (faecal samples) estimated absorbed dose rates (Lavrinienko, Tukalenko, et al., 2018; Lavrinienko et al., 2020). Outside this use of language, however, several features of Antwis et al.’s (2021) study design and interpretation of data warrant more discussion.

Demonstrating that the inclusion of non-resident fungal SV data impacts amplicon sequencing-based analyses of ‘gut mycobiota’ has important implications for studies of wildlife gut mycobiota (Lavrinienko, Scholier, et al., 2021). A consequence of not identifying the probable resident fraction of the gut mycobiota is to promote misunderstanding about the drivers of variation in wildlife gut...
mycobiota (e.g. proposing macrofungi as biomarkers of a gut micro-
bial response to radiation exposure). The inclusion of all fungal SVs can
alter the pattern of interspecific differences in ‘gut mycobiota’
due to variation in ingested fungal material among host species. As
the ecologies and traits of many fungi are poorly known, it can be
challenging to define the authentic gut mycobiota, especially for
animals with a catholic diet like the bank voles. With information
about the species of fungi eaten and/or present in dietary items, it
could be possible to use informatic tools (e.g. SourceTracker,
Knights et al., 2011) to identify the ingested fungi. Without such
data, however, the analyses of wildlife gut mycobiota will depend on
the filtering decisions; for example, one may decide to focus on the
macrofungal fraction (rather than plant-associated fungi, such as en-
dophytes, ectomycorrhizae and/or pathogens) and/or omit SVs with
poor taxonomic resolution (e.g. those not assigned beyond Phylum
level) (see e.g. Lavrinienko, Scholier, et al., 2021). While such filter-
ing decisions can change with new insights into fungal biology, an
educated ‘best guess’ about fungal traits presents an opportunity
to stimulate new research (e.g. complementing analyses of dietary
niche or identifying novel life histories of fungi). Even with imperfect
information about fungal traits, informed filtering provides a more
detailed assessment of the biological signal in the data than simply
overlooking the ecology of fungi and animal hosts.

Understanding whether laboratory protocols and sample type
impact study conclusions is an important topic in microbiota re-
search (Ingala et al., 2018; Knight et al., 2018; Videvall et al., 2018;
Zhou et al., 2020). An apparent problem with interpreting some field
studies on organisms inhabiting the CEZ is derived from authors
overemphasising statistically significant relationships that have little
explanatory power (Beresford, Scott, et al., 2020). Given that many
sources of variation in gut microbiota composition were confounded
with the analysis of ‘sample type’, including a marked difference in
the range of absorbed dose rates received by animals sampled for
either faeces or caecum, it is hard to understand why Antwis
et al.’s (2021) concluded ‘considerable variation between results for
faecal and gut samples of bank voles, suggesting faecal samples are
not an accurate indicator of gut composition’ or ‘… of the effect of
stressors on the gut microbiome’. This interpretation of these data
creates an unwarranted controversy about the effect of sample type
on microbiota analyses generally, and radiation impacts on wildlife
gut microbiota specifically.

With reference to studies on wildlife inhabiting the CEZ, Beresford, Scott, et al. (2020) stressed the importance of defining ‘… the scientific question as clearly and as unambiguously as possible’. Reflection on Antwis et al.’s (2021) faecal samples reinforces this need for clarity, for example about how the distribution of data affects the conclusions. With faecal samples collected from a single area that did not include animals in the low absorbed dose rate cate-
gory (control animals), the Abstract, Introduction and Discussion by
Antwis et al. (2021) would benefit from clear statements and hypoth-
eses that explain (1) their sampling strategy to omit control animals
(animals that did not experience elevated absorbed dose rates) and
(2) that they examined the effects of absorbed dose rates of 20 μGy/
hr and above on Apodemus mice but not the impacts of radiation
exposure on gut microbiota. By contrast, with a sample of animals
from contaminated and uncontaminated areas, Antwis et al.’s (2021)
caecum samples from bank voles identified an association between
absorbed dose rate and some taxa within the gut microbiota.

It is relevant also that the studies by Lavrinienko, Mappes, et al. (2018), Lavrinienko, Tukalenko, et al. (2018), Lavrinienko et al. (2020) and Antwis et al. (2021) employed different sampling
protocols (Antwis et al. sampled animals after 1 week of pre-baiting
traps with oats and carrot/cucumber, whereas Lavrinienko et al. did
not pre-bait traps), stored their samples differently (use of etha-
nol or not), processed samples using different DNA extraction kits
(PowerFecal DNA Isolation kit and PureLink™ Microbiome DNA
Purification Kit) and collected samples in different years (2016, and
2017, 2018), from different months (Antwis et al. took samples during
late July and in August, whereas Lavrinienko, Mappes, et al. (2018),
Lavrinienko, Tukalenko, et al. (2018), Lavrinienko et al. (2020) col-
lected their samples during May-July) and from different locations
(Figure 1). It is essential to carefully consider the diversity of mech-
nisms that can elicit a change in gut microbiota (notably, season
associates with a change in gut microbiota of bank voles inhabiting
uncontaminated areas within the CEZ; Lavrinienko et al., 2020)
when attempting to replicate a microbiota study. Without an equiv-
alent study design, or clear explanation of why changes to the proto-
col were made, important drivers(s) of apparent differences among
studies could be overlooked.

A comprehensive discussion of the statements presented by
Antwis et al. (2021) in their abstract is important because the out-
comes of studies of the wildlife inhabiting the CEZ can be used to as-
sess the risks of radiation exposure and formulate policy. Independent
examinations of the biological impacts of radionuclide exposure are
needed to form robust conclusions, but these studies are informative
only when the comparison incorporates an appropriate study design:
failure to do so will only cloud our understanding of the biological
impacts of exposure to environmental radionuclides. For the reasons
outlined above, Antwis et al. (2021) made some strong assertions
that are hard to reconcile with their, or indeed others’, data and study
design(s). One unfortunate result of this attempt to stimulate debate
is yet another controversy that does not appear to be justified.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHORS’ CONTRIBUTIONS
All authors conceived the ideas; P.C.W. analysed the data; A.L. and P.C.W. led the writing of the manuscript, with all authors making critical contributions to the drafts and giving their final approval for publication.

DATA AVAILABILITY STATEMENT
Sequence data originally presented in the study by Antwis et al. (2021) (https://doi.org/10.1111/1365-2656.13507) are available from the NCBI SRA database under the accession number PRJNA594002.

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