Fecal microbiota changes in NZB/W F1 mice after induction of lupus disease

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The association between the gut microbiota and the development of lupus is unclear. We investigated alterations in the gut microbiota after induction of lupus in a murine model using viral peptide of human cytomegalovirus (HCMV). Three treatment arms for the animals were prepared: intraperitoneal injection of HCMVpp65 peptide, adjuvant alone, and PBS injection. Feces were collected before and after lupus induction biweekly for 16S rRNA sequencing. HCMVpp65 peptide immunization induced lupus-like effects, with higher levels of anti-dsDNA antibodies, creatinine, proteinuria, and glomerular damage, compared with mice treated with nothing or adjuvant only. The Simpson diversity value was higher in mice injected with HCMVpp65 peptide, but there was no difference in ACE or Chao1 among the three groups. Statistical analysis of metagenomic profiles showed a higher abundance of various families (Saccharimonadaceae, Marinifiaceae, and Desulfovibrionaceae) and genera (Candidatus Saccharimonas, Roseburia, Odoribacter, and Desulfovibrio) in HCMVpp65 peptide-treated mice. Significant correlations between increased abundances of related genera (Candidatus Saccharimonas, Roseburia, Odoribacter, and Desulfovibrio) and HCMVpp65 peptide immunization-induced lupus-like effects were observed. This study provides insight into the changes in the gut microbiota after lupus onset in a murine model.

Systemic lupus erythematosus (SLE) is a prototype of systemic autoimmune diseases characterized by persistent chronic inflammation and production of autoantibodies, particularly anti-dsDNA antibodies. It also results in dysregulation of cytokines, leading to severe and irreversible tissue injury1,2. Although the etiology is uncertain, genetic, environmental, hormonal, and epigenetic factors are associated with SLE development3–5. The mammalian gut is colonized by trillions of microorganisms that shape intestinal microbial diversity, collectively known as the microbiota6. The reciprocal interplay between the intestinal microbiota and the host immune system maintains tissue homeostasis7–9. Therefore, autoimmune diseases such as SLE may be associated with changes in the gut microbiota.

A recent cross-sectional study confirmed decreases in species richness diversity and taxonomic complexity in the feces of lupus nephritis patients compared with controls10. Other human evidence also links gut microbiota changes to the presence of serum antinuclear antibodies and changes in inflammatory cytokines associated with SLE progression11,12. In animal studies, altered microbial community structure and greater bacterial diversity have been reported in SLE13,14. Increased relative abundances of Lachnospiraceae and Rikenellaceae were reported to be associated with the severity of murine lupus, suggesting that the gut microbiota significantly influences the host immune system and effectively affects the development of SLE15,16. The reciprocal interplay between dietary tryptophan intake and microbial dysbiosis in lupus-susceptible mice could contribute to the exacerbation of lupus17. Pattern changes in intestinal microorganisms or the presence of specific bacterial genera in the gut are associated with immune responses related to lupus.

Human cytomegalovirus (HCMV), a virus linked to the development of SLE in humans, accelerates lupus-like disease in murine models18,19. Anti-dsDNA antibody production, proteinuria, and glomerular attack have been reported in mice that received CMVpp65 or its fragment18,20. In this study, we used HCMVpp65 peptide to immunize NZB/W F1 mice to induce lupus-like effects. We investigated the dynamics of the fecal microbiota associated with lupus-like effects in HCMVpp65-immunized mice, compared with mice treated with PBS or adjuvant only.

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Results

HCMVpp65<sub>422-439</sub> immunization induces lupus-like activity in NZB/W F1 mice. HCMVpp65<sub>422-439</sub> immunization induces anti-dsDNA autoantibodies and initiates glomerulonephritis in non-autoimmune prone mice<sup>21</sup>. To investigate alterations in fecal microbiota-associated viral peptide-induced lupus-like activities, we conducted HCMVpp65<sub>422-439</sub> immunization of NZB/W F1 mice at 12 weeks of age and evaluated the lupus-like effects (Fig. 1a). The experiment timeline is shown in Fig. 1a. The serum levels of IL-6, IFN-γ, IL-10, and IL-
17A were higher in HCMVpp65422-439-immunized mice at 12 weeks post-immunization (24 weeks of age, lupus group) compared with NZB/W F1 mice treated with PBS (control group) or adjuvant only (adjuvant group, Fig. 1b–e). Anti-dsDNA antibody, serum creatinine, and proteinuria levels were elevated in the lupus group at 12 weeks post-immunization (24 weeks of age, Fig. 1f–h). Moreover, the lupus group had larger spleens, more severe renal damage, and a higher glomerulonephritis score compared with the other two groups (Fig. 1i–k). These findings suggest that HCMVpp65422-439 immunization induced lupus-like effects.

**Microbial composition in NZB/W F1 mice.** To investigate the microbial community in NZB/W F1 mice, we analyzed fecal samples collected before immunization (predisease, 12 weeks of age) and at disease time points (lupus, adjuvant, and control groups, 24 weeks of age) by 16S rRNA gene sequencing. The alpha and beta diversities were measured to evaluate alterations in microbial composition. As shown in Fig. 2a, b, the lupus group had significantly higher Shannon’s ($P = 0.033$) and Simpson’s ($P = 0.008$) diversity index values (extent of microbial diversity), but there was no difference in the ACE or Chao1 index values (extent of microbial richness), compared with the control group. A higher Simpson’s index value was found in the lupus group than in the adjuvant group. Microbial diversity and richness did not differ significantly between the adjuvant and control groups (Fig. 2a–d). For beta diversity analysis, the pattern variations in the microbial community structure among the lupus, adjuvant, and control groups are listed in Table 1. The three-dimensional principal coordinate analysis (PCoA) plot showed distinctions in community composition among the four groups (Fig. 2e). PC1, PC2, and PC3 showed 9.93%, 17%, and 25.66% of the total variance in microbial species, respectively (Fig. e). PCoA of the Bray–Curtis distance matrix revealed differences in microbial community composition among the lupus, adjuvant, and control groups (Supplementary Fig. 2). Figure 2f, g shows the relative abundances of the top 10 bacterial phyla and genera. At the phylum level, Bacteroidetes (59.0%) was the most abundant in the control group, followed by Firmicutes (39.5%), Tenericutes (1.0%), and Patescibacteria (0.3%) (Fig. 2f). The lupus group, compared with the adjuvant group, had increased abundances of Firmicutes (53.5% vs. 45.7%), Patescibacteria (3.1% vs. 1.7%), and Proteobacteria (0.6% vs. 0.007%) and decreased abundances of Bacteroidetes (41.8% vs. 52%) and Tenericutes (0.7% vs. 0.9%). At the genus level, the genera (top 10) in the relative abundance of Lachnospiraceae NK4A136 group (f_ Lachnospiraceae), Oscillibacter (f_ Oscillibacteraceae), Ruminococcaceae UCG 014 (f_ Ruminococcaceae), Intestimonas, Candidatus Saccharimonas (f_ Saccharimonadaceae), Ruminiclostridium, Ruminiclostridium 9 (f_ Ruminococcaceae), and Anaeroplasma (f_ Anaeroplasmataceae) were observed in the...
four groups (Fig. 2g). The total percentages of the top 10 genera were 22.6%, 22.4%, 25.5%, and 31.0% in the predisease, control, adjuvant, and lupus groups, respectively.

**Altered microbial families and genera associated with HCMVpp65422-439 immunization.** The families and genera with significant changes in relative abundance are shown in Figs. 3 and 4, respectively. At the family level, the abundance of *Saccharimonadaceae* was increased in the adjuvant group compared with control group (Fig. 3a). The lupus group had higher relative abundances of the families *Saccharimonadaceae* (3.1%), *Marinifilaceae* (2.0%), and *Desulfovibrionaceae* (0.7%) compared with the adjuvant and control groups (Fig. 3b, c). The average *Firmicutes* to *Bacteroidetes* (*F/B*) ratio was increased in the lupus group (1.72 ± 0.49) compared with the control (0.72 ± 0.22) and adjuvant (0.89 ± 0.23) groups (Supplementary Fig. 3a).

At the genus level, the abundances of *Candidatus Saccharimonas*, *Turicibacter*, *Ruminiclostridium 5*, and *Ruminococcaceae UCG-009* were increased in the adjuvant group compared with the control group (Fig. 4a). The abundances of *Candidatus Saccharimonas* (*f_Saccharimonadaceae*), *Odoribacter* (*f_Oodoribacteraceae*), *Roseburia* (*f_Lachnospiraceae*), and *Desulfovibrio* (*f_Desulfovibrionaceae*) were higher in the lupus group than control group (Fig. 4b). Notably, *Candidatus Saccharimonas* (3.1%), *Roseburia* (2.3%), *Odoribacter* (1.9%), and *Desulfovibrio* (0.7%) had higher abundances in the lupus group compared with the other two groups (Fig. 4c).

The abundances of the families *Akkermansiaaceae* and *Lactobacillaceae* were decreased in the three groups at 12 weeks post-immunization, but the difference in *Lactobacillaceae* abundance was not significant (Supplementary Fig. 3b, c). Linear discriminant analysis (LDA) effect size (> 3) showed differential abundances between the lupus vs. adjuvant and lupus vs. control group comparisons (Supplementary Fig. 4).

**Functional prediction of microbial communities associated with lupus-like effects in HCMVpp65422-439 immunization.** We predicted the functional potential of microbial communities by phylogenetic reconstruction of unobserved states (PICRUSt) 22. Several pathways related to cell motility were differentially expressed between the adjuvant and lupus groups (Fig. 5). Also, genetic markers with significant discriminative power in cellular processes, including lysosomes, flagellar assembly, cytok skeleton proteins, bacterial motility proteins, and bacterial chemotaxis, were detected. Spearman’s rank correlation was used to infer the associations between the fecal microbial genera and lupus-like effects. Statistically significant positive correlations between microbial genera and lupus-like effects were identified (Fig. 6). All four lupus-like effects were correlated positively with *Odoribacter*, *Desulfovibrio*, and *Roseburia*. *Candidatus Saccharimonas* showed significant positive correlations with the creatinine level, anti-dsDNA IgG titer, and glomerulonephritis severity (Fig. 6).

### Table 1. Statistical differences in microbial community structure between two groups. Three different tests were performed using MRPP, Adonis, and Anosim and were based on the Bray–Curtis dissimilarity index. E-Δ, expected-delta.

| Groups                  | MRPP          | Adonis        | Anosim        |
|-------------------------|---------------|---------------|---------------|
|                         | E-Δ | P         | R² | P         | R     | P         |
| Adjuvant vs. control    | 0.36 | 1.8E−2 | 0.21 | 6.3E−2 | 0.11 | 3.0E−1 |
| Adjuvant vs. lupus      | 0.45 | 2.0E−3 | 0.30 | 9.0E−4 | 0.66 | 3.0E−3 |
| Control vs. lupus       | 0.45 | 1.3E−2 | 0.35 | 1.4E−2 | 0.58 | 1.9E−2 |

**Figure 3.** Relative abundance of fecal microbiota families. The abundances of family groups in the (a) adjuvant vs. control and (b) adjuvant vs. lupus groups analyzed by Welch’s *t*-test using STAMP software. (c) Families with higher abundances in the lupus compared with the adjuvant and control groups. *P* < 0.05 and **P** < 0.01.
Discussion
Alterations in the microbiota composition are related to the development of SLE\textsuperscript{23}. We found that fecal microbial alterations are associated with lupus-like effects in NZB/W F1 mice. HCMVpp65\textsubscript{422-439} immunization accelerated...
lupus progression and exacerbated glomerulonephritis. The lupus group had a significantly altered microbiota composition and bacterial community compared with the adjuvant and control groups. This suggests that the abundances of significantly altered microbial families (Saccharimonadaceae, Marinifilaceae, and Desulfovibrionaceae) and genera (Candidatus Saccharimonas, Roseburia, Odoribacter, and Desulfovibrio) in the lupus group may be associated with HCMVpp65 peptide-induced lupus-like effects in NZB/W F1 mice.

Significant increases in the abundances of the families Clostridiaceae, Lachnospiraceae, Ruminococcaceae, Desulfovibrionaceae, and Rikenellaceae and a decrease in that of Akkermansia muciniphila have been described. Moreover, the same group reported increased abundances of Lachnospiraceae, Ruminococcaceae, and Rikenellaceae and a decreased abundance of Lactobacillaceae in MRL/lpr mice at the disease stage. The higher relative abundance of the Rikenellaceae family in SNF1 lupus mice is associated with more severe lupus symptoms. Increased Lactobacillales in the gut of MRL/lpr mice was linked to improved renal function and an increased survival rate. The Lactobacillaceae and Akkermansiaeace families reportedly exert an anti-inflammatory effect by increasing the IL-10 level and decreasing that of IL-6 and play a crucial role in microbiota remodeling. In this study, the families Saccharimonadaceae, Marinifilaceae, and Desulfovibrionaceae were more abundant in the lupus group than the adjuvant group. The abundances of Akkermansiaeace and Lactobacillaceae were decreased in the three groups at 12 weeks post-immunization. Of note, increased abundances of Saccharimonadaceae and

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**Figure 5.** Metagenomic taxonomic profiling of microbial communities in the adjuvant and lupus groups by phylogenetic reconstruction of unobserved states (PICRUSt).

**Figure 6.** Spearman’s rank correlation between relative genera abundance and lupus-like effects in the lupus group. Heat map based on the correlation coefficient values. + P < 0.05 and + + P < 0.01.
Marinifilaceae are positively correlated with the levels of proinflammatory cytokines such as TNF-α, IL-1β, and IL-6. An association between an increased relative abundance of the Desulfovibrionaceae family and Th17 expression was found in patients with multiple sclerosis. A significant change in phylogenetic diversity and bacterial community structure were found in the lupus group, which may be related to an increased risk of inflammatory diseases.

At the genus level, we observed increased abundances of the genera Candidatus Saccharimonomas, Roseburia, Odoribacter, and Desulfovibrio in the lupus group. These microbial genera are linked to host immune regulation and autoimmune diseases. The increased abundances of Candidatus Saccharimonomas, and Roseburia in colitis are reportedly associated with reduced IL-17 and TNF-α expression. Moreover, Roseburia may promote regulatory T-cell differentiation and reduce the expression of IL-17 in inflammatory bowel disease. In contrast, the abundances of the genera Odoribacter and Desulfovibrio are increased in several autoimmune diseases, including rheumatoid arthritis, ankylosing spondylitis, and inflammatory bowel disease likely due to an increased level of IL-17 and number of CD4+ Th17 cells. Accordingly, we examined the serum levels of IL-6, IFN-γ, IL-10, and IL-17A at 12 weeks post-immunization (24 weeks of age, disease stage). The IL-6 and IL-17A levels were significantly higher in the lupus group compared with the adjuvant and control groups. This result suggests a correlation between elevated concentrations of proinflammatory cytokines and changes in the gut microbial composition of lupus mice.

Odoribacter encompasses several intestinal butyrate-producing species. Butyrate produced by microbial fermentation provides energy for epithelial cells, maintains the integrity of the intestinal barrier, and promotes the differentiation of regulatory T cells. Several species of the genus Odoribacter are reportedly decreased in abundance in patients with rheumatoid arthritis and in lupus animal models. In addition, the abundance of Desulfovibrio was decreased in the gut of MRL/lpr mice, and that of Blautia was increased. A recent study involved both an animal model and patients with anti-neutrophil cytoplasmic antibody (ANCA) positive vasculitis and showed increased abundances of Blautia and Roseburia. These findings suggest that the gut microbiota may play a role in the pathogenesis of autoimmune diseases.

This study has several limitations. Long term co-housing of mice may result in hybrid microbiota animals or cage effects. Taking advantage of cophagophy, co-housing gnotobiotic animals shortly before immunization can reduce bias in gut microbiota analysis. We were unable to eliminate the cage effect, despite co-housing of mice before lupus induction. Moreover, the small population and disease severity may have caused bias. However, our results are consistent with prior reports of the microbiota composition in murine lupus models. However, we observed significantly higher species and microbial community diversity. Therefore, specific bacterial community compositions may be linked to the development of autoimmune diseases.

NZBW F1 mice that received HCMVpp65 peptide developed a high anti-dsDNA antibody level, serum creatinine level, and proteinuria, which were accompanied by changes in bacterial populations at 24 weeks of age. Luo reported that the bacterial genera Clostridium, Dehalobacterium, Lactobacillus, Oscillospira, Dorea (Lachnospiraceae), Bilophila (Desulfovibrionaceae), and AF12 (Rikenellaceae) were present from the pre-disease stage (10–18 weeks of age) to the disease stage (23–33 weeks of age). The abundances of the families Lachnospiraceae and Desulfovibrionaceae were increased. Therefore, immunization of NZBW F1 mice with HCMVpp65 peptide not only accelerated lupus disease but also influenced the microbiota composition. Our results do not provide insight into the causality of the relationship between lupus acceleration and microbial changes, and so further studies are needed.

Methods

Synthetic peptides. The purity of synthetic HCMVpp65 peptide (GGLAMAGASTSAGRERKEN) was > 99%, as per the manufacturer’s guarantee (GenScript Biotech Corp, Piscataway, NJ). The HCMVpp65 peptide was prepared and stored according to the manufacturer’s recommendations.

NZBW/F1 mice and induction of lupus-like effects. The animal experiments were approved by the Institutional Review Board of the Chang Gung Medical Foundation (#2016062804 and #2018121402). NZBW/F1 female mice (3–5 weeks old) were purchased from Jackson Laboratory Co., Ltd. and housed under specific pathogen-free conditions in the animal center at Chang Gung Memorial Hospital. After 30 days of adaptive feed-
ing and co-housing, the NZB/W F1 mice were separated into the control group (n = 3), adjuvant group (injected with adjuvant only, n = 5), and lupus group (induced by HCMVpp65422-439, n = 7). The mice were housed in a specific pathogen-free room under a 12 h light/12 h dark cycle with stable humidity. The immunization schedule was as described previously. Immunization was performed from 12 to 14 weeks of age. On day 1, the NZB/W F1 mice received intraperitoneal injection of 100 μg HCMVpp65422-439 emulsified with complete Freund's adjuvant (Sigma-Aldrich, Catalog Number F5881). Boosting was performed using HCMVpp65422-439, in incomplete Freund's adjuvant (Sigma-Aldrich, Catalog Number: F5506) or PBS on days 14, 28, and 42. All experiments were performed in accordance with relevant guidelines and regulations, including the ARRIVE guidelines.

**Serum, urine, and stool collection.** Stool, urine, and blood were harvested from mice once every 2 weeks. For blood collection, mice were bled from the retro-orbital vein sinus, and plasma was collected by centrifugation at 13,000 rpm for 10 min at 4 °C and stored at −80 °C. Fresh stool and urine samples were collected from mice and preserved in micro-tubes, which were immediately transferred to liquid nitrogen.

**Measurement of cytokines.** Serum levels of IL-6, IFN-γ, IL-10, and IL-17A were measured using ELISA kits (Abnova, Taipei City, Taiwan; KA4983, KA4813, KA3070, and KA3074) according to the manufacturer's instructions.

**Evaluation of lupus-like effects in NZB/W F1 mice.** Lupus-like activity was evaluated by serum indices and kidney pathological analysis using ELISA and hematoxylin and eosin (H&E) staining. Briefly, the serum level of creatinine was measured using the ELISA Test Kit (MyBioSource, MBS751125 and MBS763433, San Diego, CA). The serum anti-dsDNA autoantibody titer was evaluated using the anti-dsDNA ELISA Kit (Inova Diagnostics, catalog number: 708510, San Diego, CA) according to the manufacturers' instructions. The proteinuria level was examined using a proteinuria strip (Meditestcombi 10 VET strip, MACHEREY–NAGEL, Allentown, PA). Glomerular abnormalities were evaluated as described previously. Briefly, the number of abnormalities in 100 glomeruli within a 5-μm-thick H&E-stained paraffin section of the kidney was recorded. Glomerular abnormalities were scored as follows: normal glomeruli (score 1), pure mesangial alterations (score 2), focal segmental glomerulonephritis (score 3), diffuse glomerulonephritis (score 4), diffuse membranous glomerulonephritis (score 5), and advanced sclerosing glomerulonephritis (score 6), based on the 1982 classification of the World Health Organization.

**H&E staining.** H&E staining was conducted according to the Cold Spring Harbor protocols with slight modifications. Frozen kidney sections were immersed in 100% ethanol for 30 s and rinsed 10 times in double-distilled H2O. Slides were transferred to hematoxylin for 5 min and washed five times in 0.1% HCl. After washing in tap water for 5 s, the slides were washed five times in 0.1% NH4OH and five times in tap water. The slides were stained with eosin for 3 min and immersed in 100% ethanol with 0.1% acetic acid, 100% ethanol I, 100% ethanol II, acetone I, acetone II, xylene I, and xylene II five times each. After dehydration, the slides were mounted, covered with a cover glass, and visualized by microscopy (Olympus IX73/DP72, cellsens Standard software).

**Microbial DNA extraction and 16S rRNA gene sequencing.** Microbial DNA was extracted from fecal pellets using the DNA Stool Mini Kit (Qiagen) according to the manufacturer's instructions. The quality and quantity of DNA extracts were determined by agarose gel electrophoresis (0.8% w/v agarose) and the Nanodrop II, acetone I, acetone II, xylene I, and xylene II five times each. After dehydration, the slides were mounted, covered with a cover glass, and visualized by microscopy (Olympus IX73/DP72, cellsens Standard software).

**Data analysis and bioinformatics.** Data analysis was performed using a modified protocol of Chi-Cheng Huang. Amplicon sequencing was performed using 300-bp paired-end raw reads and assembled using FLASH v. 1.2.7. De-multiplexing was conducted based on barcode identification. For quality control, Q < 20 reads were discarded from the pipeline of QIIME v. 1.9.1. If three consecutive bases had Q ≤ 10, the read was classified as truncated. The read was preserved in the data set if it contained >75% of the original length using split_libraries_fastq.py script in QIIME. Using UCHIME algorithms, sequences were checked for chimeras to acquire the effective tags and were filtered from the data set using the UPARSE function in the peptide of USEARCH v. 7. Operational taxonomic unit (OTU) abundance was normalized to the variation and rarefied to the minimum sequence depth using the QIIME script (single_rarefaction.py). Subsequently, analysis of alpha and beta diversities was performed using the normalized data. Alpha diversity indicated the species richness and the relative abundance and evenness accounting for diversity were assessed using the Shannon and Simpson indices. Beta diversity analysis was conducted to evaluate the differences in species complexity across samples. PCoA was performed using the distance matrix to acquire principal coordinates for visualization of multi-dimensional data. The PCoA of the Bray–Curtis distance was analyzed according to OTU level for microbiota beta diversity.

For statistical analysis, the significance of all microbial species within groups at various levels of taxonomy was detected by differential abundance analysis using a zero-inflated Gaussian log-normal model executed in the “fitFeatureModel” function of the Bioconductor metagenomeSeq package. Welch’s t-test was performed using Statistical Analysis of Metagenomic Profiles (STAMP) software (v. 2.1.3). Anosim and MRPP analyses were used to determine whether the community structures significantly differed among and within groups.
Significant biomarkers were evaluated by LDA of effect size using the non-parametric factorial Kruskal–Wallis rank sum test and LDA to assess differences in taxon abundance between two groups. For functional analysis, functional abundances from 16S rRNA sequencing data were analyzed to predict functional genes using PICRUSt (v. 1.1.1)26. Two-tailed and unpaired Fisher’s tests and paired Student’s t-test with graphs depicting the mean ± the standard errors of the mean (SEM) were used for comparisons between two groups. All statistical tests were conducted with a two-sided 5% level of significance (* P ≤ 0.05; ** P ≤ 0.01; *** P ≤ 0.001) using SAS statistical software (v. 9.4, SAS Institute).

Data availability
The datasets presented in this study can be found in online repositories (Sequence Read Archive data, accession: PRJNA693398).

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Author contributions

A.H.H., L.C.W., W.Y.T., and C.F.K. designed the project. A.H.H., L.C.W., Y.F.C., Y.J.H., and Y.C.T. performed the experiments, acquired the data, and interpreted the results. A.H.H., Y.J.H., and Y.C.T. prepared the figures. A.H.H., K.H.Y., C.F.K., and S.F.L. participated in the interpretation of the results and drafted and revised the manuscript. All authors reviewed the manuscript, approved the final version to be published, and accepted responsibility for all aspects of the work.
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Competing interests
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