



Gut Microbial Stability is Associated with Greater Endurance Performance in Athletes Undertaking Dietary Periodization

Matthew J. W. Furber, ^a Gregory R. Young, ^{b,c} Giles S. Holt, ^d Simone Pyle, ^g Glen Davison, ^f Michael G. Roberts, ^a Justin D. Roberts, ^h Glyn Howatson, ^{b,e} Darren L. Smith ^{b,c}

a School of Life and Medical Sciences, Department of Psychology and Sport Sciences, University of Hertfordshire, Hertfordshire, United Kingdom

bFaculty of Health & Life Sciences, Northumbria University, Newcastle upon Tyne, United Kingdom

cHub for Biotechnology in the Built Environment, Northumbria University, Newcastle upon Tyne, United Kingdom

^dBiomedicine West Wing, Newcastle University, Newcastle upon Tyne, United Kingdom

eWater Research Group, School of Environmental Sciences and Development, Northwest University, Potchefstroom, South Africa

fUniversity of Kent, Canterbury, United Kingdom

⁹University of Reading Whiteknights, Reading, United Kingdom

^hCambridge Centre for Sport and Exercise Sciences, School of Psychology and Sport Science, Anglia Ruskin University, Cambridge, United Kingdom

Matthew J.W. Furber and Gregory R. Young contributed equally to this work. Author order for equal contribution authors was decided by alphabetical order of surnames.

ABSTRACT Dietary manipulation with high-protein or high-carbohydrate content are frequently employed during elite athletic training, aiming to enhance athletic performance. Such interventions are likely to impact upon gut microbial content. This study explored the impact of acute high-protein or high-carbohydrate diets on measured endurance performance and associated gut microbial community changes. In a cohort of well-matched, highly trained endurance runners, we measured performance outcomes, as well as gut bacterial, viral (FVP), and bacteriophage (IV) communities in a double-blind, repeated-measures design randomized control trial (RCT) to explore the impact of dietary intervention with either high-protein or high-carbohydrate content. High-dietary carbohydrate improved time-trial performance by +6.5% (P < 0.03) and was associated with expansion of Ruminococcus and Collinsella bacterial spp. Conversely, high dietary protein led to a reduction in performance by -23.3% (P = 0.001). This impact was accompanied by significantly reduced diversity (IV: P = 0.04) and altered composition (IV and FVP: P = 0.02) of the gut phageome as well as enrichment of both free and inducible Sk1virus and Leuconostoc bacterial populations. Greatest performance during dietary modification was observed in participants with less substantial shifts in community composition. Gut microbial stability during acute dietary periodization was associated with greater athletic performance in this highly trained, well-matched cohort. Athletes, and those supporting them, should be mindful of the potential consequences of dietary manipulation on gut flora and implications for performance, and periodize appropriately.

IMPORTANCE Dietary periodization is employed to improve endurance exercise performance but may impact on gut microbial communities. Bacteriophage are implicated in bacterial cell homeostasis and have been identified as biomarkers of disequilibrium in the gut ecosystem possibly brought about through dietary periodization. We find high-carbohydrate and high-protein diets to have opposing impacts on endurance performance in highly trained athlete populations. Reduced performance is linked with disturbance of microbial stasis in the gut. We demonstrate bacteriophage communities are the most sensitive component of the gut microbiota to increased gut stress following dietary manipulation. Athletes undertaking dietary periodization should be aware of potential negative impacts of drastic changes to dietary composition on gut microbial stasis and, in turn, endurance performance.

Editor Seth Bordenstein, Vanderbilt University
Copyright © 2022 Furber et al. This is an openaccess article distributed under the terms of
the Creative Commons Attribution 4.0
International license.

Address correspondence to Darren L. Smith, Darren.smith@northumbria.ac.uk.

The authors declare a conflict of interest. MF and SP have previously worked for Glaxo Smith Klyne. GH and DS have previously received funding from the same institution for investigation of the gut microbiota. Previous employers or funders had no influence on the study design, participant recruitment, results presented or the conclusions drawn here.

Received 10 February 2022 **Accepted** 15 April 2022

KEYWORDS human microbiome, recovery, endurance running, high-protein diet, carbohydrate loading diet, randomized controlled trial, bacteria, bacteriophages, gut microbiome, endurance, physical

Despite decades of research, the ongoing debate around optimal nutrition persists. It is well established that carbohydrate utilization and exercise performance are intrinsically linked (1, 2). For example, carbohydrate is the preferred substrate during high intensity endurance exercise and athletes who consume a high carbohydrate diet (HCD) can maximize endurance performance (2, 3). However, in some circumstances reducing carbohydrate intake during periods of low intensity endurance training can enhance performance via metabolic and cellular adaptations associated with oxidative phosphorylation (4, 5). Dietary periodization is a concept that uses different regimes to optimize the balance between training adaptation and performance. Athletes employ low carbohydrate diets during physical training with the aim of enhancing adaptations (6) before switching to HCD prior to competition to maximize energy stores, thereby improving potential for athletic performance (7).

The implementation of high-protein diets (HPD) during training have become increasingly popular regimes among athletes (8, 9), in part to offset carbohydrate-depleted states. Body mass and body composition are additional factors to consider in endurance performance and HPDs are used during training to help manage these (10, 11) and stimulate muscle growth (12, 13). Despite the potential advantages of HPD, the impact on gut homeostasis is an area of contention (14). In particular, the effects of HPDs on metabolism and gene expression of both gut microbes and host epithelial cells raise questions about longer-term health.

HPDs, especially those rich in red meat, contain large amounts of sulfated amino acids such as cysteine and methionine (15, 16). Proteolytic fermentation of these amino acids in the distal colon is associated with microbial metabolites such as ammonia, phenols, and hydrogen sulfide (17) which can cause damage to the gut microstructure (18) or mucosa (19). Some phenolic compound products of protein metabolism have also been linked with increased gut permeability (20–22). Over prolonged periods, these physiological changes may lead to negative health outcomes. The gut microbiota plays an important role in colonic protein fermentation (23), and high dietary protein consumption has been associated with enrichment of anaerobic bacteria with proteolytic capabilities (24).

Importantly for well-trained athletes, following strict diets and rigorous training regimes, the gut microbiota is implicated in host physiological functions. Endurance exercise is linked to an increased bacterial diversity in the gut due to elevated stress levels (25). Similarly, there are differences in gut bacterial communities in physically active compared to sedentary humans (26, 27). However, these studies do not account for different diets between the populations. This issue has been previously highlighted (26) given that athletes consumed more total energy, fat, carbohydrate, and protein in comparison to less active controls. To date, there are no studies that have explored the microbial communities within the gut across microbial kingdoms in relation to dietary regimens in well-trained athletes. This is of importance given previous work has illustrated the influence of bacteriophage (the viruses of bacteria) as biomarkers of disequilibrium in gut communities (28). Phages are intrinsically linked to the metabolism, stress, and activity of active bacterial communities in the gut. Furthermore, Cronin et al. (32) identified the most descriptive part of the gut microbiota composition of sedentary participants taking on the "couch to 5 km" running challenge was the inducible viral fraction. Despite the potential influence of the viral component, this remains an understudied facet of gut microbial communities, particularly in relation to diet manipulation in highly trained athletes and the influence on human performance.

Previous work has numerous confounding factors of age, lifestyle, diet, physical activity, and other baseline characteristics. This study focuses on a homogenous group of well-trained runners who share baseline dietary and physical activity habits. The aim of this study was to assess the impact of a high-protein versus high-carbohydrate diet on performance in highly trained endurance runners while measuring associated gut

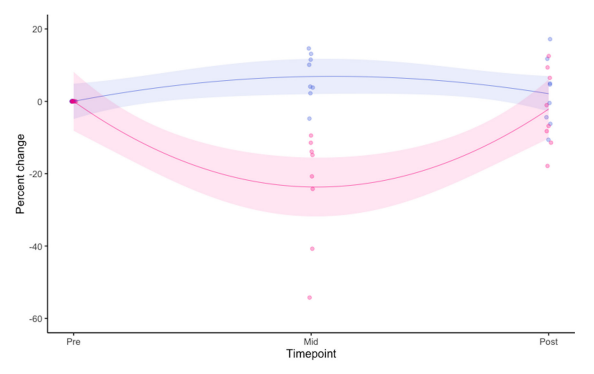


FIG 1 Illustrates longitudinal subject performance during the Max SE trial. Each point represents an individual subject through preintervention (pre), midintervention (mid), and postintervention (post), sampling time points, colored by intervention group (HCD = blue; HPD = pink). The line represents best fit of the linear regression model based on all subjects at that time point. The shaded area describes the 95% confidence intervals of the linear model.

bacterial, viral, and fungal communities. Recruiting such a well-controlled cohort enables us to highlight changes to exercise capacity and associated microbial features associated specifically with dietary interventions. Critically to the aim of this study, diet was rigorously controlled which enabled data to be attributable to the manipulation of HCD and HPD.

RESULTS

High carbohydrate and high-protein diets have opposing effects on time trial to exhaustion performance. The results of the 95% MaxSE trial are shown in Fig. 1, expressed as relative change (%) from preintervention baseline. Further detail on the impact of the dietary intervention on performance has been previously reported (29). Following HPD, time trial to exhaustion (TTE) was reduced within group by -23.3% midintervention, with all participants exhibiting decreased ability to perform (P < 0.001). Upon subsequent return to habitual dietary intake, TTE for HPD postintervention time points was comparable with preintervention performance (pre:128.3 \pm 29.3 s, mid: 98.4 \pm 31.8 s, post: 125.3 \pm 32.39 s). In contrast, a 6.5% improvement in TTE was observed midintervention in the HCD group (P = 0.05) with seven of the eight participants improving relative to preintervention performance. TTE performance for HCD also returned to preintervention levels by trial closure (pre: 182.2 \pm 44.4 s, mid: 194 \pm 45.5 s, post: 184 \pm 38.9 s).

Poor performance in HPD is associated with changes to microbiota observable in stool. A total of 48 stool samples from 16 participants (eight HPD, eight HCD intervention), were sequenced to investigate the bacterial, viral, and fungal microbiota. Each participant provided three samples capturing pre-, mid-, and postintervention gut microbiota (see supplementary materials for details of sequencing parameters).

Information on sample sizes and comparisons of samples and controls is available in the supplementary materials (Fig. S1). Significant dissimilarity was observed between controls and samples (ANOSIM: $R^2 = 0.99$; P = 0.001; Fig. S1).

Before comparing impact of dietary interventions on microbial communities we sought to define the baseline of bacterial diversity within the participants enrolled in this study. We

Month YYYY Volume XX Issue XX

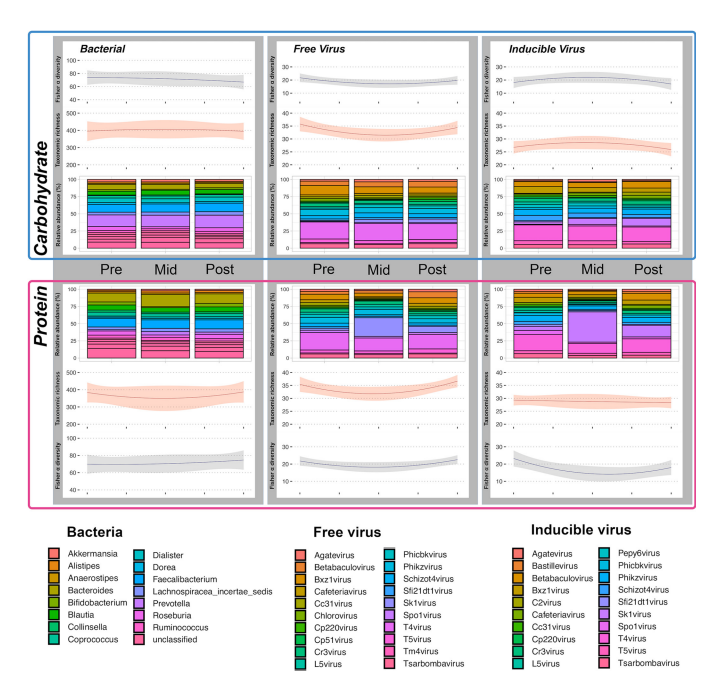


FIG 2 Illustrates overall bacterial, FVP and IV communities measured at pre-, mid-, and postintervention time points for participants enrolled on both the HPD and HPC diets. Longitudinal progression of both Fisher-alpha diversity (navy line), and observed taxonomic richness (red line), are plotted for each community. Shaded area represents 95% confidence interval based on standard error. Microbial taxa included in the bar charts represent the 16 and 20 most abundant bacteria or viruses observed within the data sets, respectively.

compared beta-diversity between participant stool samples in this cohort to age-matched males enrolled in the human microbiome project (30). Significantly lower inter-individual variability was observed between participants enrolled within this study than those enrolled in the HMP cohort (Adonis PERMANOVA P=0.001, Fig. S2). These data suggest that the highly trained athletes enrolled here harbor significantly distinct gastrointestinal bacterial communities to that of a typical western population.

Dietary intervention had a significant impact on microbial community composition. Significantly reduced Fisher-alpha diversity of inducible viruses was observed during HPD intervention (KW: P = 0.04), with levels failing to return to preintervention levels following cessation (Fig. 2) (Table S1).

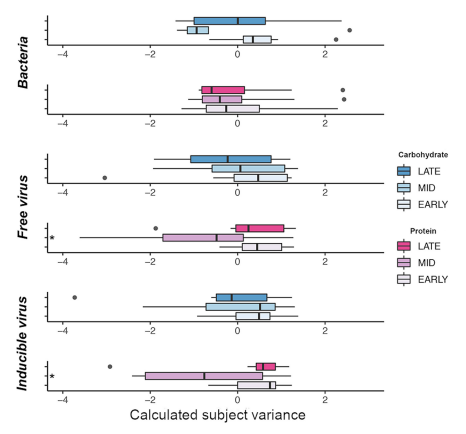


FIG 3 Illustrates shifts in beta-diversity of microbial communities following both HPD (pink), and HCD (blue), interventions. Plotted are the loadings values of the primary component of principle coordinates analysis, time point is depicted by color intensity (post-/late = dark; mid- = light; pre-/early = lightest). The center of each box represents the median, edges extend to upper and lower quantiles and whiskers describe the full range of data, excluding outliers, which are plotted as individual points. Significant shifts in beta-diversity within communities are highlighted by asterisk.

HPDs diet had the greatest impact on viral community composition while HCD diet had the greatest impact on bacterial community composition (Fig. 3). Both free viral particle (FVP) ($R^2 = 0.15$; P = 0.023), and inducible virus (IV) ($R^2 = 0.16$; P = 0.016) communities were significantly altered between during HPD intervention (Fig. 3, Table S2). Following cessation of HPD both the FVP and IV communities recovered toward preintervention community compositions (Fig. 3, Fig. S3).

Taxonomic features of the microbiota are associated with dietary interventions. Changes in viral community diversity and composition during HPD intervention is largely contributed to by an expansion in proportional abundance of Sk1virus. This expansion of Sk1virus is visible in both FVP and IV communities (Fig. 2).

Sparse partial least-squares regression (sPLS-DA), analysis was utilized to identify discriminative features between habitual and HCD or HPD interventions. We combined bacterial, FVP, and IV microbial communities to identify discriminate features between dietary interventions. sPLS-DA highlighted strong positive correlations between proportional abundances of Sk1virus and *Leuconostoc* bacteria, indicating both features are associated with HPD intervention (Fig. 4a). Furthermore, *Leuconostoc* were negatively correlated with IV Schizot4virus, which was reduced during HPD intervention in both viral communities. *Bifidobacterium* spp. proportional abundance was also substantially reduced during HPD intervention.

Despite nonsignificant shifts in the overall community during HCD intervention, proportional abundances of *Leuconostoc* were strongly associated with HCD, as in HPD. *Lactococcus* and *Collinsella* were also both strongly associated with HCD intervention (Fig. 4b). In contrast, *Streptococcus* exhibited substantially reduced proportional abundance

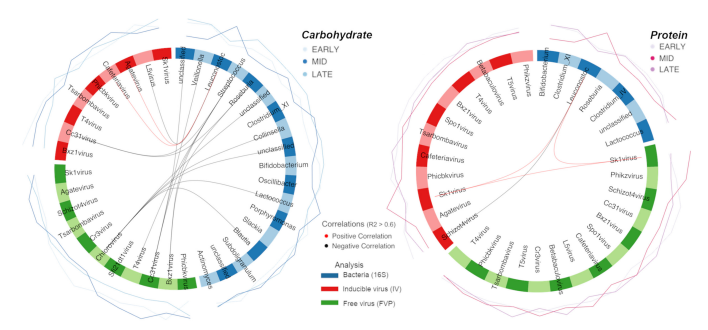


FIG 4 Shows potential biomarkers of both HPD (a) and HCD (b) dietary intervention as identified by sPLS-DA. Significant positive (red), and negative (black), correlations with R^2 values > 0.6 between features of each community are highlighted by lines joining nodes of bacterial (blue), FVP (green), and IV (red) nodes, forming the outside of the circle. Proportional abundance of each feature is illustrated by external lines colored by time point as in Fig. 2. Effect sizes of each individual feature per community are highlighted in the loadings plot and are colored by the same logic as proportional abundance.

following HCD intervention. Viral features proportionally upregulated in HCD intervention included Cc31virus which was negatively correlated with *Streptococcus* in both FVP and IV communities, as well as IV *Cafeteria virus*. Conversely, FVP Chlorovirus and IV L5virus were both negatively correlated with *Leuconostoc* and were proportionally reduced during HCD intervention.

Combination of viral and bacterial communities enabled greater discrimination between dietary intervention time points than using any community in isolation as evidenced by sPLS-DA (Fig. S4).

Gut microbial taxonomic stability is associated with greater performance. Participants were stratified in to "responders" and "nonresponders" based on percentage change in TTE performance recorded during mid-intervention tests for both HCD and HPD. Based on the results of the linear model (Fig. 1), "responders" were classified as those subjects for which TTE increased more than expected during HCD (n=4) or reduced more than expected during HPD (n=4) intervention. Likewise, four participants were classified as "nonresponders" in both the HPD and HCD interventions.

Combined microbial communities were employed to identify relationships between gut microbial communities and athletic performance during both dietary interventions. Subject variance was used as a measure of microbial stability within individual participants. Microbial communities of improving performers in the HCD intervention were far more stable over time exhibiting lower within-subject community dissimilarity than subjects who did not improve. These observations were coupled with maintenance of a subject-specific gut microbiota in HCD improvers. Nonimproving subjects on the HCD were less distinguishable from one another (Fig. 4, Fig. S3). In contrast, subjects in the HPD with reducing performance exhibited greater microbial turbulence with bigger shifts in gut communities across the study and lower subject-specificity (Fig. 5, Fig. S3). These results suggest that athletic performance may be linked with gut microbial stasis, where athletes harboring stable microbial communities consistently performed best in each dietary intervention compared to those with a more turbulent gut microbiota.

Assessing gene function in gut microbial communities and inferring impact on athletic performance. Better performance during dietary periodization was observed in participants with greater gut microbiota taxonomic stability. We explored whether

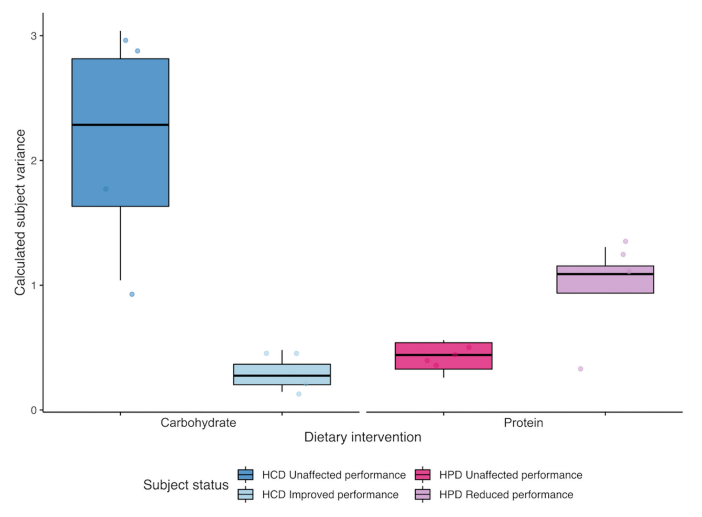


FIG 5 Illustrates within and between subject variance based on Bray-Curtis of combined bacterial, FVP and IV community dissimilarity for participants enrolled in either the HCD (blue), or HPD (pink), interventions. Within subject variance explains the dissimilarity between samples from one individual across all three sampling time points. Between subject variance explains the dissimilarity between samples from different individuals enrolled on the same dietary intervention. Subjects are stratified in to "unaffected" (dark colors), or "improved" and "reduced" groups (light colors), dependent on TTE performance changes within each dietary group. Each point represents variance calculated for an individual subject. Box edges describe the upper and lower quartile ranges while the median is depicted by the central line. Whiskers extend to the full range of data points, excluding outliers.

this was related to functional capacity of the microbiota by interrogating metagenomic data. A total of 12,581 viral orthologous gene (VOG) functional genes were identified across all 48 samples (median 287984, IQR 233105 to 372302). VOG functional genes were grouped into high level classifications as explained in Text S1. Large percentages (up to 75%) of the viral genes identified during this analysis were either hypothetical, putative or unclassifiable by alignment to the VOG database (Fig. 6a). Minor expansions in abundance of temperate viral genes associated with host metabolism were observed during HPD intervention. Genes associated with viral structure were marginally more proportionally abundant during the HCD (Fig. 6a). Percentage change from baseline of functional gene richness and dominance was not different (P > 0.05) across either temperate or lytic viral fractions between responders and nonresponders. HCD had a general, though nonsignificant, reducing effect on IV richness (Fig. 6b). Participants who performed best at TTE on the HCD diet, however, recovered from this loss of functional richness whereas nonimprovers continued to suffer reductions in IV functional richness even after intervention cessation (Fig. 6b). Up to a 17% reduction in functional gene dominance was observed in IV-HPD responders and up to 53% reduction in FVP-HCD responders (Fig. 6c), though these differences were not significant. Several significantly differentially abundant pathways (q < 0.25) of bacterial metabolism were observed between responders and

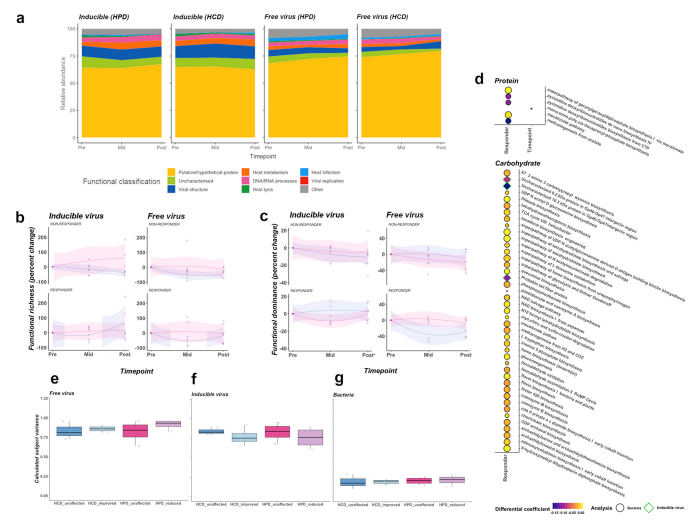


FIG 6 illustrates functional gene compositions of samples. Functional composition across each time point is represented as relative abundance of functional gene classes (a). Changes in functional gene richness (b) and dominance (c) is also presented across sampling time points, stratified by responder status. Each point represents an individual sample and is colored by diet (pink = HPD; blue = HCD). Significantly differential gene features as identified by Maaslin2 are represented by individual bubbles, colored by differential coefficient and stratified by diet (d). Significance (q value) is represented by size of the bubble. Circles represent bacterial gene pathways while diamonds represent viral genes. Associations between microbial functional composition and performance during dietary intervention in FVP (e), IV (f), and bacterial (g) compartments are represented by box and whisker plots, as in Fig. 5.

nonresponders to both dietary interventions by Maaslin2 analysis (Fig. 6d). Many of these, including inositol degradation, thiazole biosynthesis and NAD salvage/biosynthetic pathways are directly related to carbohydrate utilization. Similarly, to the results of taxonomic analyses, we observed greater change in overall FVP community composition in participants whose performance at TTE was reduced during HPD intervention (Fig. 6e). Additionally, HCD improvers had greater functional community stability in the IV compartment than their nonimproving counterparts (Fig. 6f). Results of functional and taxonomic analyses did not match for FVP HCD or IV HPD cohorts.

DISCUSSION

This study investigated the influence of two dietary strategies commonly used by athletes on gut microbiome populations and the subsequent impact on running performance. This study is the first to illustrate the impact of short-term, highly controlled HCDs and HPDs on the performance of well-trained endurance athletes. This study showed new data that isocaloric, high-protein diets in highly trained athletes resulted in reduced running performance that was correlated with alterations in gut viral communities. Importantly, changes in the viral communities represent a more sensitive

marker of gut stress during dietary intervention than bacterial community analysis alone.

Previous studies have explored changes in the gut microbiota of active versus sedentary individuals finding greater taxonomic (25, 27) and metabolic (26) diversity in active individuals. Additionally, other investigations highlighted the impact of differing levels of dietary protein (24), and carbohydrate (31) on the gut microbiota. These studies focus specifically on gut bacterial communities. However, as highlighted by, Cronin et al., bacteriophage communities can also be implicated in physical performance (32). Here, we utilized a modified approach, stratifying both free and chemically induced lysogenic phages to assess the impact of dietary periodization on each component individually. This current work has identified alterations in viral communities because of dietary change in well-trained athletes. Importantly, this study employed a blinded dietary intervention that was very tightly controlled, and in a highly trained cohort. Consequently, we identified association between the shifts in both bacterial and viral communities in the gut and athletic performance. We were able to discount any variability arising from confounding factors of diet and exercise that exist in previous work by having a well-controlled dietary regimen and homogenously active volunteers. For example, this study cohort harbored distinct gut bacterial communities to that of average human western populations; this was determined by comparison of beta-diversity between stool samples from this cohort and those collected from similar aged males participating in the human microbiome project (Fig. S2) (30). This in turn amplifies the impact of dietary manipulations for even relatively short (7 day) periods.

Of interest to well-trained endurance populations, these data showed that a high-protein diet for 7 days, alongside endurance training, significantly reduced running performance. This manifested as a dramatic -23.3% reduction in mean TTE across participants in the HPD trial arm. Conversely, improved TTE was observed (+6.5%) in athletes on the HCD, which is understandable given the large body of evidence showing high carbohydrate availability is conducive to endurance performance (2, 3). In support of this expired gas analysis performed on this cohort during the 10 km TTE showed there was a greater reliance on fat oxidation following HPD (data previously reported [33]), likely due to lower carbohydrate substrate availability as opposed to the greater use of carbohydrate oxidation. Collectively, these data further illustrate the benefit of HCD for endurance performance in well-trained athletes.

In this study the implementation of a HPD had a greater impact on the viral compartment while HCD had a greater impact on the bacterial microbiota. Increased dietary protein content was associated with reduced IV taxonomic alpha diversity and functional gene dominance, reduced FVP richness, and marked shifts in community compositions of both inducible and free viruses which we hypothesize relates to phage induction, through selective pressure on the bacterial population. Community shifts in both IV and FVP populations were attributable to an expansion in the relative abundance of *Sk1virus* observed in stools. *Sk1viruses* are members of the *Siphoviridae*, which were found to be significantly enriched during HPDs previously (34). Additionally, we found the abundance of *Leuconostoc* bacteria to be positively correlated with *Sk1virus* during HPD intervention. We propose this is a function of a predator-prey expansion relationship as *Sk1virus* are members of the 936-type phage that show broad-range infectivity within the Lactobacillales, of which *Leuconostoc* are a member (35, 36). The co-occurrence of these microbes in individuals consuming HPD may be due to supplementation with proteins derived from whey as these microbes are highly abundant in and associated with fermentation of dairy products (37).

Elevated carbohydrate intake in this study was not associated with the same changes in gut viral communities observed in HPD, but instead had an impact on bacterial community compositions through expansions of *Collinsella & Ruminococcus* spp. *Collinsella* have previously been correlated with high levels of circulating insulin (38) and have broad dietary carbohydrate metabolizing potential (39). *Ruminococcus* are well-known degraders of resistant starch in the human gut (31, 40). We propose that the enriched bacterial populations observed during HCD in this study occurred as a direct impact of the increased substrate availability associated with enhanced carbohydrate volumes in the diet. Large shifts in

bacterial taxonomy observed in this study correlated with large numbers of differential functional pathways identified in inferred metagenomes. HCD nonresponders failed to recover from loss of IV functional richness following cessation of dietary intervention which may point to links between a lack of functional plasticity and poor performance during HCD intervention. Conversely, the expansion in both inducible and free viral taxonomic communities identified in the HPD may be a result of greater bacterial cell stress in the gut environment due to a lack of fermentable substrate (carbohydrate) in the HPD and highlight the potential negative impacts of HPD.

Analysis of responders and nonresponders to the dietary interventions provide evidence of these proposed differences. For example, responders to the HCD exhibited greater improvements in TTE performance than nonresponders. This improvement was associated with markedly greater taxonomic microbial community stability and marginally greater functional stability of inducible viruses. In contrast, participants whose performance was affected by the HPD performed significantly worse than their peers. This is reflected in the greater change in overall taxonomic communities and FVP functional communities because of the intervention. Maintenance of a subject-specific community and reduced longitudinal variation was observed in athletes who performed better during dietary periodization. Changes in gut communities were associated with lower TTE performance in both dietary interventions. Athletes undergoing dietary periodization with the aim of improving performance would likely benefit from greater gut microbial stability. This is the first time such a phenomenon has been described, and in practical terms may have significant implications for nutrition strategies for training and performance. Constant shifts in dietary intake result in instability of the microbial communities. Athletes that alternate between strict dietary regimes may be more likely to cause microbial instability which may impact performance.

Limitations of this study include the relatively small sample size, lack of mechanistic analysis to link changes in microbial communities more definitively to performance, and the inability to guarantee participant compliance to dietary intervention. However, diets were rigorously controlled and training stimuli were maintained between cohorts, enabling high confidence of trial protocol compliance. This work provides a rare insight into the potential flux experienced in athletes that might engage in dietary periodization and the implications it might have on performance. We performed metagenomic sequencing of viral communities which enabled us to characterize functional genes encoded in the FVP and IV compartments. Unfortunately, the high number of genes reported with unknown function in virus genomes means that databases searches have limited utility. Up to 75% of genes identified in this study aligned to gene targets of unknown or hypothetical function in the VOG database. Future studies should attempt to explore functional characteristics of the viral microbiota, specifically that of bacteriophage, which can have a large impact on bacterial communities in the gut. Furthermore, the functional annotation of bacterial genes performed here was based on inferred metagenomes, predicted from 16S sequencing data. Links between functional plasticity within the gut microbial community, resilience to dietary change and athletic performance should be further explored in larger, metagenomic community analyses. Isolation of gut microbes to characterize carbohydrate and protein substrate utilization of microbial community members, particularly lysogens, would further confirm the hypotheses formed during this study.

Conclusion. This study provides new insights into dietary manipulation in athletic populations. Gut microbial stability was associated with greater athletic performance when highly trained individuals underwent dietary periodization. Implementation of an acute high-protein diet resulted in compromised athletic performance in individuals with less stable gut microbiota. We propose this observation is attributable to increased stress within the gut environment. Viral populations provide high sensitivity to determine gut stress and could be used as a marker of microbial instability in future studies. Short-term high carbohydrate diet improved athletic performance and was characterized by subtle alterations to the gut microbiota. Stable microbial communities were associated with better performance. Subjects consistently performed better during physical tests when microbial communities remained relatively

TABLE 1 Baseline physical characteristics of study participants

Group	Age (yrs)	Height (cm)	Body mass (kg)	Absolute V^2O_2 max (L · min ⁻¹)	Relative V^2O_2 max (mL·kg ⁻¹ ·min ⁻¹)
HPD (n = 8)	25 ± 3.6	179.4 ± 6.4	69.5 ± 3.3	4.38 ± 0.35	63.1 ± 4.8
HCD (n = 8)	27 ± 5.0	181.6 ± 3.5	67.6 ± 6.1	4.39 ± 0.28	65.3 ± 6.4

HPD, high-protein diet; HCD, high-carbohydrate diet. No differences were observed in baseline participant characteristics according to allocated starting group (mean \pm SD) (p > 0.05).

unchanged throughout dietary periodization. Athletes, and those supporting them, should be mindful of the potential consequences of altering diet on performance and gut flora and periodize dietary intake (particularly macronutrient distribution) appropriately.

MATERIALS AND METHODS

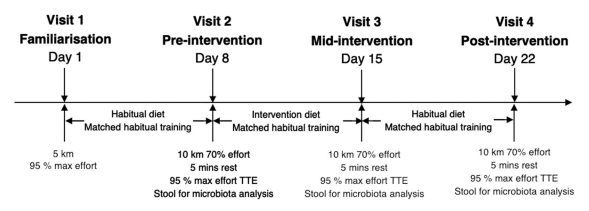
Study design and cohort characteristics. This study was conducted in accordance with the Declaration of Helsinki (2013) and was approved by the University of Hertfordshire Life and Medical Sciences Ethics Committee (protocol number LMS/PGR/UH/02227). To be considered, participants were required to be "highly trained" endurance athletes and not taking any dietary supplements before or during the trial period (see supplementary materials for detailed inclusion criteria). A minimum of six participants were required per trial arm according to power calculation assessment for sample size (G *power3, Dusseldorf) using $\alpha = 0.05$; $1 - \beta = 0.80$ based on previous exercise data. Twenty suitable participants were recruited, providing written informed consent (Table 1), with four not completing the study for reasons unrelated to the trial.

In a double-blind, parallel group, repeated-measures design, participants were randomly assigned to either isocaloric HPD (40% protein, 30% carbohydrate, and 30% fat macronutrient distribution) or energy-matched HCD (10% protein, 60% carbohydrate, and 30% fat macronutrient distribution). While participants remained blinded to which trial arm they were assigned, due to the nature of dietary intervention, we could not blind food items. Further information about diet composition is available in supplementary materials. Participants attended the laboratory on four separate occasions to undertake performance testing and provide samples for microbiota analysis. After a rest day from exercise, no morning physical exertion, and following a minimum 4-h fast, participants arrived at the laboratory via the same mode of transport at the same time each visit. Visits were preceded by 7 days of either habitual or prescribed intervention diet (Fig. 6, and Fig. 7). Participants were requested to maintain training regimes throughout the duration of the study as previously reported (29).

Participants collected their first stool of the day which was transferred to -80°C storage within a maximum of 2 h of passing and remained there until analysis. Participants then completed a 10 km steady state run at $70\% \text{ V}^2\text{O}_2\text{max}$, had 5 min rest then completed a maximum effort at $95\% \text{ V}^2\text{O}_2\text{max}$ effort until volitional exhaustion (also known as time trial to exhaustion [TTE]). Further details of sample collection and TTE are available in supplementary materials.

Control of dietary intake. A 3-day food diary (2 weekdays and 1 weekend day) was used to assess habitual dietary intake prior to visits 2 and 4. During the study briefing participants were instructed how to complete the food dairy and provided with a comprehensive example. The importance of accuracy and detail were emphasized, as was the importance of maintaining current dietary habits. All food diaries were analyzed by the same researcher using dietary analysis software (Nutritics, Dublin, Ireland).

Individual energy intake for the intervention diet was prescribed to match energy expenditure and was calculated using basal metabolic rate (BMR) multiplied by a physical activity factor (PAF) (described in detail in supplementary materials). A total of 34 diet plans (17 HCD, 17 HPD) were formulated by the investigative team, ranging from 1,800 kcal·d⁻¹ to 4200 kcal·d⁻¹. Participants were assigned the diet closest to their calculated interventional energy intake. An additional 500 kcal·day⁻¹ macronutrient-



 $\textbf{FIG 7} \ \ \textbf{Illustrates the method design employed in this study}.$

TABLE 2 Calculated, prescribed dietary intake (mean \pm S.E.) and macro-nutrient breakdown for each group during the 7-day intervention based on estimated energy requirements

	Energy intake		Carbohydrate		Protein		Fat	
Group	Kcal⋅d ⁻¹	Kcal⋅kg ⁻¹ ⋅d ⁻¹	$g \cdot d^{-1}$	g⋅kg ⁻¹ ⋅d ⁻¹	$g \cdot d^{-1}$	g⋅kg ⁻¹ ⋅d ⁻¹	$g \cdot d^{-1}$	$g \cdot kg^{-1} \cdot d^{-1}$
HPD	3185 ± 84	48 ± 1.2	239 ± 6.3	3.4 ± 0.9	319 ± 8.4^{a}	4.6 ± 0.1^{a}	106 ± 2.8	1.5 ± 0.04
HCD	3281 ± 69	49 ± 1.0	492 ± 10.4^a	7.3 ± 0.1^{a}	82 ± 1.7	1.2 ± 0.03	109 ± 2.3	1.5 ± 0.03

HPD, high-protein/reduced carbohydrate diet (40/30/30 – protein, carbohydrate, fat ratio); HCD, high carbohydrate/typical protein diet (60/10/30 – carbohydrate, protein, fat ratio).

matched meal was provided to ensure participants remained in a positive energy balance during the intervention week. Water was available *ad libitum*, but participants were requested to abstain from drinks containing caffeine or additional energy. The mean dietary intake for both prescribed diets is shown in Table 2.

Assessment of training volume. To ensure any observed changes were due to the intervention diet and not a due to change in training volume, the same weekly training program was followed by each participant throughout the experimental period. Training sessions were logged and submitted online with GPS smartwatches provided to each participant (Garmin Connect, Garmin Ltd., Schaffhausen, Switzerland).

Isolation of gut microbial DNA. Stool samples collected during each lab visit were defrosted and homogenized in 3 mL of ice cold, sterile 1x Phosphate buffered saline (PBS) then allowed to settle for 5 min. Also, 1 mL supernatant was sub-sampled for viral community analysis. The viral sub-sample was processed as per modified protocols previously described (41). Briefly, the sub-sample was centrifuged at 4,000 rpm and 4°C for 10 min. The supernatant was used for free viral particle (FVP) analysis and the pellet used for inducible virus (IV) community analysis. The IV pellet was resuspended in 1 mL sterile 1x PBS and incubated with norfloxacin at 1 μ g/mL for 1 h at 37°C. The incubated pellet was then centrifuged at 4,000 rpm and 4°C for a further 10 min and the supernatant taken for DNA isolation.

Prior to isolation of FVP, IV, and bacterial and fungal DNA, all samples were treated with of 1 μ L TURBO DNase and 1 μ L RNase Cocktail (Life Technologies, CA, USA) to deplete any free DNA in solution. Treatment consisted of 30 min at 37°C before inactivation at 65°C for 1 min in 15 mM EDTA. Viral DNA was extracted as per manufacturer's instructions using NORGEN Phage DNA Isolation Kits (Geneflow Limited, Lichfield, UK). Bacterial and fungal DNA was isolated from the remaining homogenized stool following removal and processing of the 1 mL viral supernatant using DNEasy PowerLyzer PowerSoil DNA Isolation kits (Qiagen, DE), as per manufacturer's instructions.

Sequencing and processing of gut microbial DNA. All sequencing was performed by NUOmics (Newcastle, UK). Viral DNA metagenome sequencing libraries were processed via the Illumina Nextera XT prep (Illumina, Saffron Waldon, UK). Libraries were sequenced on the Illumina MiSeq using V3 2 × 300 bp chemistry. Viral sequence reads were trimmed and quality filtered using *Trimmomatic* (42) and *Sickle* (43). Sequences homologous to the human genome were culled using *KneadData* in the *bioBakery* environment (44). Taxonomic assignment was achieved by aligning sequences against the NCBI nr database using a kmer size of 20% and 70% identity threshold with *blastn* (45). Lowest common ancestor and relative abundances were calculated in *MEGAN6* (46). Functional annotation of viral sequences was performed by aligning sequences to those in the VOG database (https://vogdb.org, accessed 6/12/21) using default *VIBRANT* (47) parameters. Normalized abundance of identified VOGs per sample was calculated using *bwa mem* (48), *samtools* (49), *bamtools* (50), and *bedtools* (51). To discriminate between multiple target alignments of a single sequence against the VOG database, a hierarchical preference criterion was applied. Firstly, actual classifications were preferred over putative ones. If multiple actual classifications were present for a single sequence, greater *VIBRANT* outputted *V.score* was preferred, followed by lower *E.score*. Custom R scripts merged feature counts per sample to create feature tables for statistical analysis.

Bacterial and fungal DNA amplicon sequencing libraries were prepared as per the Schloss protocol (52). Bacterial amplicons targeted the V4 region of the 16S rRNA gene while fungal amplicons targeted the ITS1 and ITS2 regions of rRNA spacer genes. Both bacterial and fungal libraries were sequenced on the Illumina MiSeq using V2 2 × 250 bp chemistry. Taxonomic assignment of bacterial and fungal sequence reads was performed in Mothur (53). Briefly, paired end reads were trimmed, merged, quality filtered, and clustered into *de novo* operational taxonomic units (OTUs) using the OptiClust method (54). OTUs were aligned to SILVA (bacterial [55]), and UNITE (fungal [56]), databases to determine taxonomy. *PiCrust2* (57) was used to infer functional metagenomes from, chimera filtered, 97% homology bacterial OTU reference sequences classified using "ready-to-wear" classifiers specific for human stool communities (58, 59) by aligning to the Greengenes (13_8) database (60) in *QIIME2* (61).

All raw sequencing data is freely available at European Nucleotide Archive (ENA) under study accession PRJEB45703.

Statistical analysis. Statistical analyses were performed using R (62). Normality of data was verified by the Shapiro–Wilk test. Kruskal-Wallis rank-sum test (KW) was used to compare means of continuous data. Where appropriate, a Bonferroni *post hoc* test was used.

Alpha diversity of microbial communities was assessed by richness, diversity (Fisher-alpha), and dominance (Simpson) metrics while beta diversity was assessed by Bray-Curtis dissimilarity for taxonomic analyses and Canberra distance for functional gene composition analyses in the *vegan* package (63). All taxonomic diversity metrics were calculated at the genus level. Adonis PERMANOVA was used

^aDenotes significantly more intake (p < 0.05).

to identify metadata variables significantly impacting community compositions utilizing the "strata" function to account for repeated measures. Microbial stability/change was assessed using calculated subject variance, being the average range of loadings values from the primary principal component of dissimilarity between microbial communities from a single trial participant over time. Discriminatory features of combined bacterial and viral microbiota between intervention time points were identified by sparse partial least-squares regression based on combined bacterial, FVP and IV communities. The model was tuned to prune noninformative variables using the DIABLO method (64). Significantly differential features within functional gene analyses were identified using Maaslin2 (65). Phyloseq (66), mixOmics (67) and ggplot2 (33) packages were used to visualize data. STORM checklist: https://zenodo.org/record/6530945#.YnjV0FzMLAg.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

TEXT S1, PDF file, 0.1 MB.

FIG S1, PDF file, 0.1 MB.

FIG S2, PDF file, 0.04 MB.

FIG S3, PDF file, 0.4 MB.

FIG S4, PDF file, 0.1 MB.

TABLE S1, PDF file, 0.03 MB.

TABLE S2, PDF file, 0.05 MB.

ACKNOWLEDGMENTS

Matt Furber and Simone Pyle have previously worked for Glaxo Smith Kline. Glyn Howatson and Darren Smith have previously received funding from the same institution for investigation of the gut microbiota. Previous employers or funders had no influence on the study design, participant recruitment, results presented, or the conclusions drawn here.

REFERENCES

- Bergström J, Hermansen L, Hultman E, Saltin B. 1967. Diet, muscle glycogen and physical performance. Acta Physiol Scand 71:140–150. https://doi.org/10.1111/j.1748-1716.1967.tb03720.x.
- Burke LM, Hawley JA, Wong SHS, Jeukendrup AE. 2011. Carbohydrates for training and competition. J Sports Sci 29:S17–S27. https://doi.org/10.1080/ 02640414.2011.585473.
- Hearris MA, Hammond KM, Fell JM, Morton JP. 2018. Regulation of muscle glycogen metabolism during exercise: implications for endurance performance and training adaptations. Nutr 10:298.
- Burke LM, Hawley JA, Jeukendrup A, Morton JP, Stellingwerff T, Maughan RJ. 2018. Toward a common understanding of diet–exercise strategies to manipulate fuel availability for training and competition preparation in endurance sport. Int J Sport Nutr Exerc Metab 28:451–463. https://doi.org/10.1123/ijsnem.2018-0289.
- Hulston CJ, Venables MC, Mann CH, Martin C, Philp A, Baar K, Jeukendrup AE. 2010. Training with low muscle glycogen enhances fat metabolism in well-trained cyclists. Med Sci Sport Exerc 42:2046–2055. https://doi.org/ 10.1249/MSS.0b013e3181dd5070.
- Impey SG, Hearris MA, Hammond KM, Bartlett JD, Louis J, Close GL, Morton JP. 2018. Fuel for the work required: a theoretical framework for carbohydrate periodization and the glycogen threshold hypothesis. Sports Med 48:1031–1048. https://doi.org/10.1007/s40279-018-0867-7.
- Bartlett JD, Hawley JA, Morton JP. 2015. Carbohydrate availability and exercise training adaptation: too much of a good thing? Eur J Sport Sci 15:3–12. https://doi.org/10.1080/17461391.2014.920926.
- Campbell B, Kreider RB, Ziegenfuss T, La Bounty P, Roberts M, Burke D, Landis J, Lopez H, Antonio J. 2007. International Society of Sports Nutrition position stand: protein and exercise. J Int Soc Sports Nutr 4:8. https:// doi.org/10.1186/1550-2783-4-8.
- Huecker M, Sarav M, Pearlman M, Laster J. 2019. Protein supplementation in sport: source, timing, and intended benefits. Curr Nutr Rep 8:382–396. https://doi.org/10.1007/s13668-019-00293-1.
- Wycherley TP, Buckley JD, Noakes M, Clifton PM, Brinkworth GD. 2013. Comparison of the effects of weight loss from a high-protein versus standard-protein energy-restricted diet on strength and aerobic capacity in overweight and obese men. Eur J Nutr 52:317–325. https://doi.org/10 .1007/s00394-012-0338-0.

- Leidy HJ, Clifton PM, Astrup A, Wycherley TP, Westerterp-Plantenga MS, Luscombe-Marsh ND, Woods SC, Mattes RD. 2015. The role of protein in weight loss and maintenance. Am J Clin Nutr 101:1320–1329.
- Burd NA, Beals JW, Martinez IG, Salvador AF, Skinner SK. 2019. Food-first approach to enhance the regulation of post-exercise skeletal muscle protein synthesis and remodeling. Sport Med 49:59–68. https://doi.org/10 .1007/s40279-018-1009-y.
- Moore DR, Robinson MJ, Fry JL, Tang JE, Glover EI, Wilkinson SB, Prior T, Tarnopolsky MA, Phillips SM. 2009. Ingested protein dose response of muscle and albumin protein synthesis after resistance exercise in young men. Am J Clin Nutr 89:161–168. https://doi.org/10.3945/ajcn.2008.26401.
- 14. Blachier F, Beaumont M, Portune KJ, Steuer N, Lan A, Audebert M, Khodorova N, Andriamihaja M, Airinei G, Benamouzig R, Davila AM, Armand L, Rampelli S, Brigidi P, Tomé D, Claus SP, Sanz Y. 2019. High-protein diets for weight management: Interactions with the intestinal microbiota and consequences for gut health. A position paper by the my new gut study group. Clin Nutr 38:1012–1022. https://doi.org/10.1016/j.clnu.2018.09.016.
- Yao CK, Muir JG, Gibson PR. 2016. Review article: Insights into colonic protein fermentation, its modulation and potential health implications. Aliment Pharmacol Ther 43:181–196. https://doi.org/10.1111/apt.13456.
- USDA. 2016. USDA national nutrient database for standard reference, release 28. USDA.
- Macfarlane GT, Macfarlane S. 2012. Bacteria, colonic fermentation, and gastrointestinal health. J AOAC Int 95:50–60. https://doi.org/10.5740/ jaoacint.sge_macfarlane.
- Dai Z-L, Wu G, Zhu W-Y. 2011. Amino acid metabolism in intestinal bacteria: links between gut ecology and host health. Front Biosci (Landmark Ed) 16:1768–1786. https://doi.org/10.2741/3820.
- Christl SU, Eisner HD, Dusel G, Kasper H, Scheppach W. 1996. Antagonistic effects of sulfide and butyrate on proliferation of colonic mucosa: a potential role for these agents in the pathogenesis of ulcerative colitis. Digest Dis Sci 41:2477–2481. https://doi.org/10.1007/BF02100146.
- 20. Lewis S, Cochrane S. 2007. Alteration of sulfate and hydrogen metabolism in the human colon by changing intestinal transit rate. Am J Gastroenterol 102:624–633. https://doi.org/10.1111/j.1572-0241.2006.01020.x.
- 21. Magee EA, Richardson CJ, Hughes R, Cummings JH. 2000. Contribution of dietary protein to sulfide production in the large intestine: an in vitro and

- a controlled feeding study in humans. Am J Clin Nutr 72:1488–1494. https://doi.org/10.1093/ajcn/72.6.1488.
- McCall IC, Betanzos A, Weber DA, Nava P, Miller GW, Parkos CA. 2009. Effects of phenol on barrier function of a human intestinal epithelial cell line correlate with altered tight junction protein localization. Toxicol Appl Pharmacol 241:61–70. https://doi.org/10.1016/j.taap.2009.08.002.
- Korpela K. 2018. Diet, microbiota, and metabolic health: trade-off between saccharolytic and proteolytic fermentation. Annu Rev Food Sci Technol 9: 65–84. https://doi.org/10.1146/annurev-food-030117-012830.
- Moreno-Pérez D, Bressa C, Bailén M, Hamed-Bousdar S, Naclerio F, Carmona M, Pérez M, González-Soltero R, Montalvo-Lominchar MG, Carabaña C, Larrosa M. 2018. Effect of a protein supplement on the gut microbiota of endurance athletes: a randomized, controlled, double-blind pilot study. Nutr 10:337.
- Karl JP, Margolis LM, Madslien EH, Murphy NE, Castellani JW, Gundersen Y, Hoke AV, Levangie MW, Kumar R, Chakraborty N, Gautam A, Hammamieh R, Martini S, Montain SJ, Pasiakos SM. 2017. Changes in intestinal microbiota composition and metabolism coincide with increased intestinal permeability in young adults under prolonged physiological stress. Am J Physiol - Gastrointest Liver Physiol 312:G559–G571. https://doi.org/10.1152/ajpgi.00066.2017.
- Barton W, Penney NC, Cronin O, Garcia-Perez I, Molloy MG, Holmes E, Shanahan F, Cotter PD, O'Sullivan O. 2018. The microbiome of professional athletes differs from that of more sedentary subjects in composition and particularly at the functional metabolic level. Gut 67:625–633. https://doi.org/10.1136/gutjnl-2016-313627.
- Clarke SF, Murphy EF, O'Sullivan O, Lucey AJ, Humphreys M, Hogan A, Hayes P, O'Reilly M, Jeffery IB, Wood-Martin R, Kerins DM, Quigley E, Ross RP, O'Toole PW, Molloy MG, Falvey E, Shanahan F, Cotter PD. 2014. Exercise and associated dietary extremes impact on gut microbial diversity. Gut 63:1913–1920. https://doi.org/10.1136/gutjnl-2013-306541.
- Lepage P, Colombet J, Marteau P, Sime-Ngando T, Doré J, Leclerc M. 2008. Dysbiosis in inflammatory bowel disease: a role for bacteriophages? Gut 57:424–425. https://doi.org/10.1136/gut.2007.134668.
- Furber M, Pyle S, Roberts M, Roberts J. 2021. Comparing acute, high dietary protein and carbohydrate intake on transcriptional biomarkers, fuel utilisation and exercise performance in trained male runners. Nutr 13 https://www.mdpi.com/2072-6643/13/12/4391/htm#.
- 30. HMPC THMPC. 2012. Structure, function and diversity of the healthy human microbiome. Nature 486:207–214. https://doi.org/10.1038/nature11234.
- Walker AW, Ince J, Duncan SH, Webster LM, Holtrop G, Ze X, Brown D, Stares MD, Scott P, Bergerat A, Louis P, McIntosh F, Johnstone AM, Lobley GE, Parkhill J, Flint HJ. 2011. Dominant and diet-responsive groups of bacteria within the human colonic microbiota. ISME J 5:220–230. https://doi.org/10.1038/ismej.2010.118.
- 32. Cronin O, Barton W, Skuse P, Penney NC, Garcia-Perez I, Murphy EF, Woods T, Nugent H, Fanning A, Melgar S, Falvey EC, Holmes E, Cotter PD, O'Sullivan O, Molloy MG, Shanahan F. 2018. A prospective metagenomic and metabolomic analysis of the impact of exercise and/or whey protein supplementation on the gut microbiome of sedentary adults. mSystems 3. https://doi.org/10.1128/mSystems.00044-18.
- 33. Whickham H. 2009. ggplot2: elegant graphics for data analysis. Springer-Verlag, New York.
- Minot S, Sinha R, Chen J, Li H, Keilbaugh SA, Wu GD, Lewis JD, Bushman FD. 2011. The human gut virome: inter-individual variation and dynamic response to diet. Genome Res. https://doi.org/10.1101/gr.122705.111.
- Mahony J, Murphy J, van Sinderen D. 2012. Lactococcal 936-type phages and dairy fermentation problems: from detection to evolution and prevention. Front Microbiol 3:335. https://doi.org/10.3389/fmicb.2012.00335.
- 36. Ackermann HW. 2001. Frequency of morphological phage descriptions in the year 2000. Arch Virol 146:843–857. https://doi.org/10.1007/s007050170120.
- 37. Frantzen CA, Kot W, Pedersen TB, Ardö YM, Broadbent JR, Neve H, Hansen LH, Dal Bello F, Østlie HM, Kleppen HP, Vogensen FK, Holo H. 2017. Genomic characterization of dairy associated leuconostoc species and diversity of leuconostocs in undefined mixed mesophilic starter cultures. Front Microbiol 8:132. https://doi.org/10.3389/fmicb.2017.00132.
- Gomez-Arango LF, Barrett HL, Wilkinson SA, Callaway LK, McIntyre HD, Morrison M, Dekker Nitert M. 2018. Low dietary fiber intake increases Collinsella abundance in the gut microbiota of overweight and obese pregnant women. Gut Microbes 9:189–201. https://doi.org/10.1080/19490976 .2017.1406584.
- Kageyama A, Benno Y, Nakase T. 1999. Phylogenetic and phenotypic evidence for the transfer of Eubacterium aerofaciens to the genus Collinsella as Collinsella aerofaciens gen. nov., comb. nov. Int J Syst Evol Microbiol 49:557–565. https://doi.org/10.1099/00207713-49-2-557.

- 40. Abell GCJ, Cooke CM, Bennett CN, Conlon MA, McOrist AL. 2008. Phylotypes related to Ruminococcus bromii are abundant in the large bowel of humans and increase in response to a diet high in resistant starch. FEMS Microbiol Ecol 66:505–515. https://doi.org/10.1111/j.1574-6941.2008.00527.x.
- 41. Tariq MA, Everest FLC, Cowley LA, De Soyza AD, Holt GS, Bridge SH, Perry A, Perry JD, Bourke SJ, Cummings SP, Lanyon CV, Barr JJ, Smith DL. 2015. A metagenomic approach to characterize temperate bacteriophage populations from cystic fibrosis and non-cystic fibrosis bronchiectasis patients. Front Microbiol 6. https://doi.org/10.3389/fmicb.2015.00097.
- 42. Bolger AM, Lohse M, Usadel B. 2014. Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 30:2114–2120. https://doi.org/10.1093/bioinformatics/btu170.
- Joshi N, Fass J. 2011. Sickle: a sliding-window, adaptive, quality-based trimming tool for FastQ files (Version 1.33) [Software]. https://github.com/najoshi/sickle.
- McIver LJ, Abu-Ali G, Franzosa EA, Schwager R, Morgan XC, Waldron L, Segata N, Huttenhower C. 2018. bioBakery: a meta'omic analysis environment. Bioinformatics 34:1235–1237. https://doi.org/10.1093/bioinformatics/ btx754.
- Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search tool. J Mol Biol 215:403–410. https://doi.org/10.1016/ S0022-2836(05)80360-2.
- Huson DH, Auch AF, Qi J, Schuster SC. 2007. MEGAN analysis of metagenomic data. Genome Res https://doi.org/10.1101/gr.5969107.
- Kieft K, Zhou Z, Anantharaman K. 2020. VIBRANT: automated recovery, annotation and curation of microbial viruses, and evaluation of viral community function from genomic sequences. Microbiome 8:90. https://doi .org/10.1186/s40168-020-00867-0.
- 48. Li H. 2013. Aligning sequence reads, clone sequences and assembly contigs with BWA-MEM.
- Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G, Durbin R, Subgroup 1000 Genome Project Data Processing. 2009. The Sequence Alignment/Map format and SAMtools. Bioinformatics 25:2078–2079. https://doi.org/10.1093/bioinformatics/btp352.
- Barnett DW, Garrison EK, Quinlan AR, Strömberg MP, Marth GT. 2011. Bam-Tools: a C++ API and toolkit for analyzing and managing BAM files. Bioinformatics 27:1691–1692. https://doi.org/10.1093/bioinformatics/btr174.
- Quinlan AR, Hall IM. 2010. BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26:841–842. https://doi.org/10.1093/bioinformatics/btq033.
- Kozich JJ, Westcott SL, Baxter NT, Highlander SK, Schloss PD. 2013. Development of a dual-index sequencing strategy and curation pipeline for analyzing amplicon sequence data on the MiSeq illumina sequencing platform. Appl Environ Microbiol 79:5112–5120. https://doi.org/10.1128/AEM.01043-13.
- Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, Hollister EB, Lesniewski RA, Oakley BB, Parks DH, Robinson CJ, Sahl JW, Stres B, Thallinger GG, Van Horn DJ, Weber CF. 2009. Introducing mothur: open-source, platform-independent, community-supported software for describing and comparing microbial communities. Appl Environ Microbiol 75:7537–7541. https://doi.org/10.1128/AEM.01541-09.
- Westcott SL, Schloss PD. 2017. OptiClust, an improved method for assigning amplicon-based sequence data to operational taxonomic units. mSphere 2. https://doi.org/10.1128/mSphereDirect.00073-17.
- Quast C, Pruesse E, Yilmaz P, Gerken J, Schweer T, Yarza P, Peplies J, Glöckner FO. 2013. The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. Nucleic Acids Res 41:D590–D596. https:// doi.org/10.1093/nar/gks1219.
- Nilsson RH, Abarenkov K, Larsson K-H, Köljalg U. 2011. Molecular identification of fungi: rationale, philosophical concerns, and the UNITE database. TOAINFOJ 5:81–86. https://doi.org/10.2174/1874136301105010081.
- Douglas GM, Maffei VJ, Zaneveld JR, Yurgel SN, Brown JR, Taylor CM, Huttenhower C, Langille MGI. 2020. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol 38:685–688. https://doi.org/10.1038/ s41587-020-0548-6.
- Bokulich NA, Kaehler BD, Rideout JR, Dillon M, Bolyen E, Knight R, Huttley GA, Gregory Caporaso J. 2018. Optimizing taxonomic classification of marker-gene amplicon sequences with QIIME 2's q2-feature-classifier plugin. Microbiome 6:90. https://doi.org/10.1186/s40168-018-0470-z.
- Kaehler BD, Bokulich NA, McDonald D, Knight R, Caporaso JG, Huttley GA. 2019. Species abundance information improves sequence taxonomy classification accuracy. Nat Commun 10:4643. https://doi.org/10.1038/s41467-019-12669-6.

- McDonald D, Price MN, Goodrich J, Nawrocki EP, DeSantis TZ, Probst A, Andersen GL, Knight R, Hugenholtz P. 2012. An improved Greengenes taxonomy with explicit ranks for ecological and evolutionary analyses of bacteria and archaea. ISME J 6:610–618. https://doi.org/10.1038/ismej .2011.139.
- 61. Bolyen E, Rideout JR, Dillon MR, Bokulich NA, Abnet CC, Al-Ghalith GA, Alexander H, Alm EJ, Arumugam M, Asnicar F, Bai Y, Bisanz JE, Bittinger K, Brejnrod A, Brislawn CJ, Brown CT, Callahan BJ, Caraballo-Rodríguez AM, Chase J, Cope EK, Da Silva R, Diener C, Dorrestein PC, Douglas GM, Durall DM, Duvallet C, Edwardson CF, Ernst M, Estaki M, Fouquier J, Gauglitz JM, Gibbons SM, Gibson DL, Gonzalez A, Gorlick K, Guo J, Hillmann B, Holmes S, Holste H, Huttenhower C, Huttley GA, Janssen S, Jarmusch AK, Jiang L, Kaehler BD, Kang K, Bin Keefe CR, Keim P, Kelley ST, Knights D, Koester I, Kosciolek T, et al. 2019. Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2. Nat Biotechnol 37:852–857. https://doi.org/10.1038/s41587-019-0209-9.
- 62. R Core Team. 2014. R: a language and environment for statistical computing. R Foundation for Statistical Computing.

- 63. Oksanen J, Guillaume Blanchet F, Kindt R, Legendre P, Minchin PR, O'Hara RB, Simpson GL, Solymos P, Henry M, Stevens H, Wagner H. 2015. vegan: community ecology package. http://cran.r-project.org/.
- 64. Singh A, Shannon CP, Gautier B, Rohart F, Vacher M, Tebbutt SJ, Lê Cao K-A. 2019. DIABLO: an integrative approach for identifying key molecular drivers from multi-omics assays. Bioinformatics 35:3055–3062. https://doi.org/10.1093/bioinformatics/bty1054.
- Mallick H, Rahnavard A, McIver LJ, Ma S, Zhang Y, Nguyen LH, Tickle TL, Weingart G, Ren B, Schwager EH, Chatterjee S, Thompson KN, Wilkinson JE, Subramanian A, Lu Y, Waldron L, Paulson JN, Franzosa EA, Bravo HC, Huttenhower C. 2021. Multivariable association discovery in populationscale meta-omics studies. PLoS Comput Biol 17:e1009442. https://doi .org/10.1371/journal.pcbi.1009442.
- McMurdie PJ, Holmes S. 2013. phyloseq: An R package for reproducible interactive analysis and graphics of microbiome census data. PLoS One 8: e61217. https://doi.org/10.1371/journal.pone.0061217.
- Rohart F, Gautier B, Singh A, Lê Cao KA. 2017. mixOmics: An R package for 'omics feature selection and multiple data integration. PLoS Comput Biol 13:e1005752. https://doi.org/10.1371/journal.pcbi.1005752.