

# Galactoglucomannan fibres promote a beneficial porcine gut microbiome

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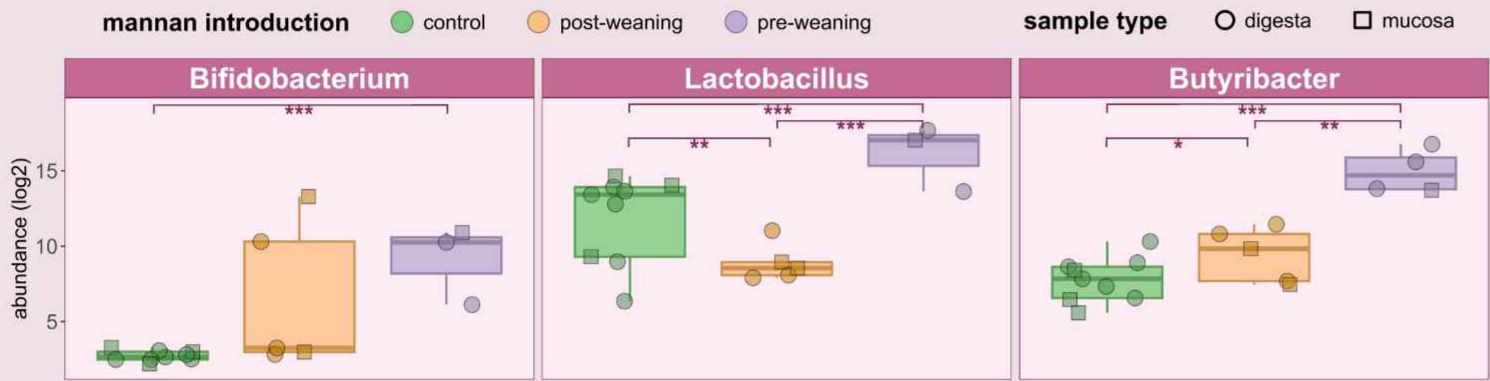
Most mammals and their microbiomes are codependent, forming a functional unit known as a holobiont. Exchanging metabolites, regulating gene expression, and combating pathogens are vital to the health and performance of the holobiont 1-2. By understanding the interactions occurring within these complex systems, we can more effectively improve animal and feed production, favouring both animal welfare, production efficiency, and the growing needs of the increasing human population<sup>34</sup>.

Mannan fibres derived from spruce can be metabolised into hostaccessible compounds by microbes with carbohydrate-

active enzymes 56. Microbes like these also ease piglets' transition from milk to solid feed 7. Hence we ask: can we jump-start the young porcine microbiome by adding mannan fibres to their preweaning diet?

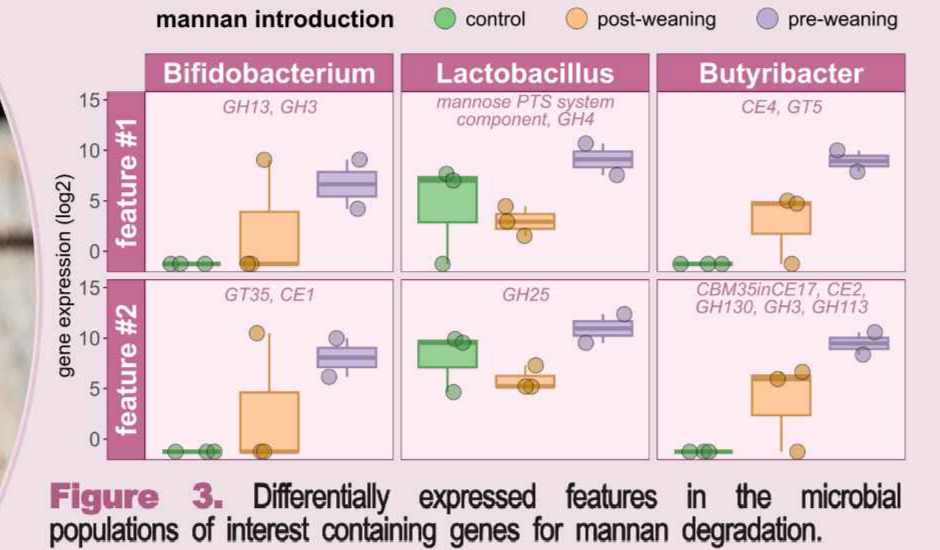
## SELECTIVE PROMOTION

Among the differentially abundant 9 populations between control and mannan-fed piglets are bacteria often associated with host health benefits: Bifidobacterium longum (log<sub>2</sub> fold change 8.3), Lactobacillus johnsonii (LFC 5.9), and Butyribacter sp. (LFC 5.7) (Fig. 2).

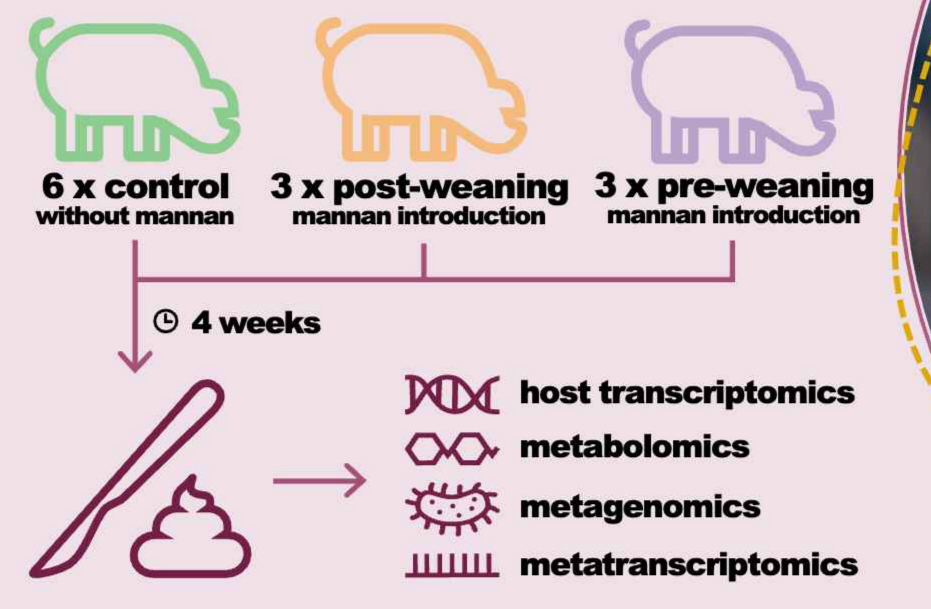


Three microbial populations with differential abundance between piglet groups. Significance by false discovery rate-adjusted p-values are indicated by asterisks: \* < 0.05, \*\* < 0.01, \*\*\* < 0.001.

Many of these populations' differentially expressed genes yield enzymes for mannan degradation, like glycoside hydrolases and transferases, and carbohydrate esterases (Fig. 3).



Thus it seems yes, we can jump-start the porcine microbiome by mannan supplementation.



The four generated datasets were analysed both as individual omic layers (Sec. 3 and 4), and jointly through a holo-omic approach (Sec. 5).

# HOLO-OMIC MODEL

A multiset correlation and factor analysis (MCFA)<sup>10</sup> model reconstructs the full dataset in three feature spaces: one shared by all omic layers; one private to each layer; and an omic-specific residual. Inferred shared components are similar to PCs, but comprises features from all omic layers. The model (Fig. 4) fits well with mannan exposure, hence it will be used to learn more about the interactions across the porcine holo-omic boundary.

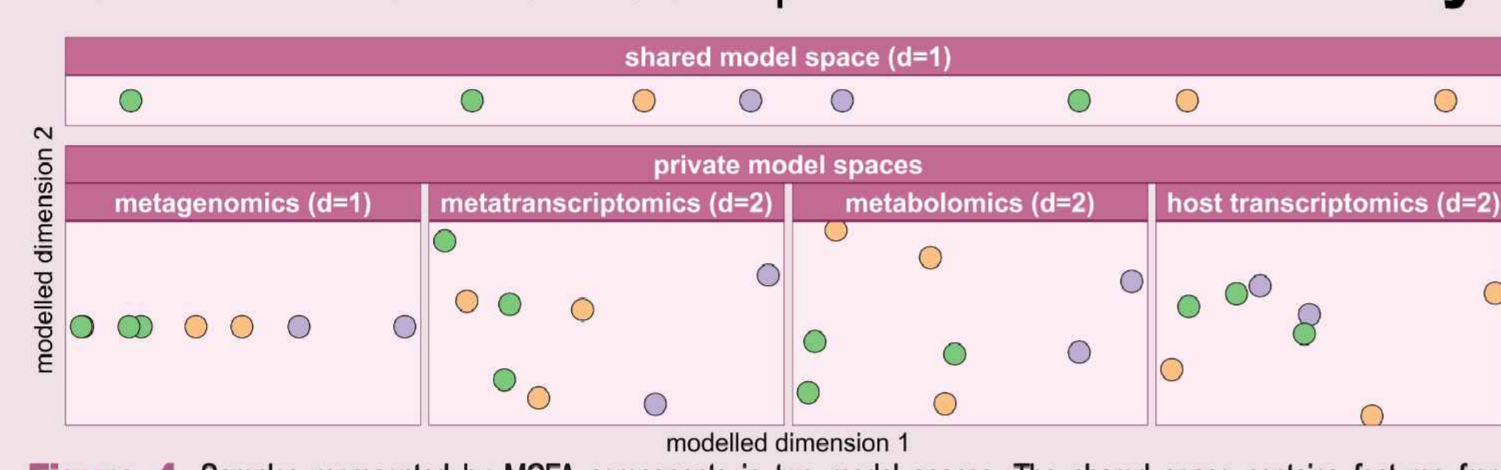


Figure 4. Samples represented by MCFA components in two model spaces. The shared space contains features from all layers. Aspects private to each omic layer add to this shared space, together recontructing the full holo-omic dataset.

MICROBIAL GRADIENTS

Principal component (PC) analyses<sup>8</sup> of individual omic layers in Fig. 1 show gradients that correspond with mannan exposure duration.

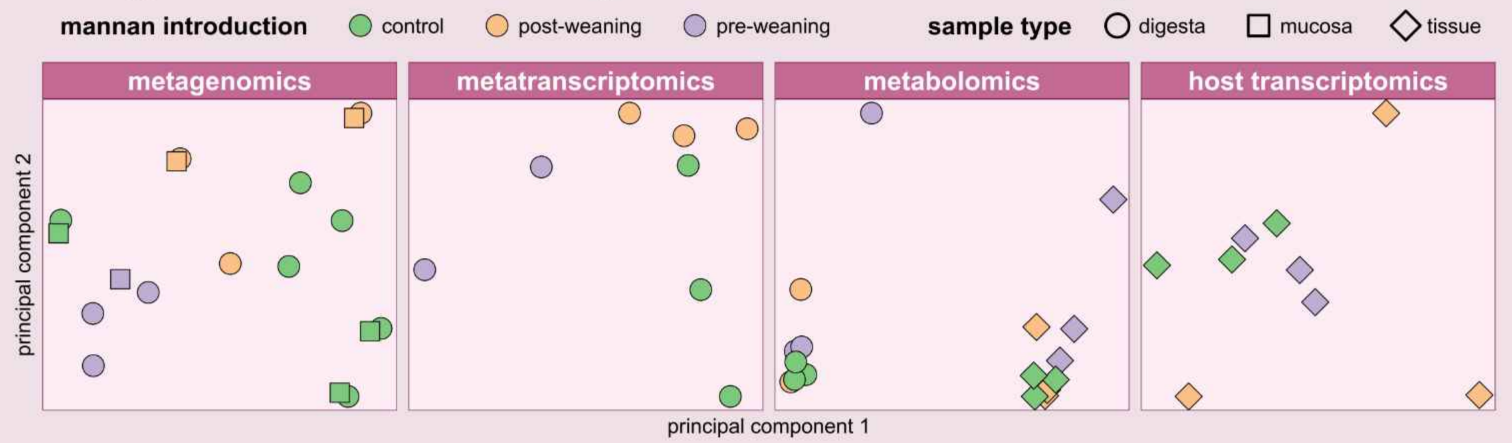


Figure 1. Principal component analyses of each omic layer. Points represent samples from piglets introduced to dietary mannan fibres at different developmental stages. Relative placements indicate similarities between samples' omic profiles.

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