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1	SUPPLEMENTARY FIGURES AND TABLES
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3	Drinking pattern and sex modulate the impact of ethanol consumption on the mouse
4	gut microbiome
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10	LEGENDS FOR SUPPLEMENTARY FIGURES
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12	Figure S1. Experimental protocol used in the current study. Male and female C57BL/6J
13	mice were randomly divided into four groups: control, chronic ethanol, binge, and chronic
14	plus binge ethanol. Mice from the chronic ethanol and chronic plus binge groups were
15	submitted to one-week-adaptation period with 5% (v/v) ethanol solution. Mice then had
16	free access to 10% (v/v) ethanol in the drinking water for 10 days. Mice from the control
17	and binge groups had free access to water during the treatment period. Mice from the
18	binge and chronic plus binge groups, received an oral gavage with a high dose of ethanol
19	(5 g.kg ⁻¹) on day 11. Mice from control and chronic ethanol groups received an oral
20	gavage with water on day 11. Blood samples and fecal pellets were collected 2 or 9 hours
21	after the oral gavage, respectively.
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23	Figure S2. Heatmap showing shifts in the gut microbiota induced by maltodextrin in male
24	and female mice. Fecal samples were collected before and 9 hours after intragastric
25	gavage with maltodextrin (9 g.kg ⁻¹). Gut microbiota composition (phyla category) was
26	evaluated by RT-qPCR. *Compared to fecal samples collected before oral gavage
27	(p<0.05, Student's t test).
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29	Figure S3. Rarefaction curves plotted using alpha diversity metric, Shannon index
30	against number of sequencing depth. (A) C57BL/6J mice were randomly divided into

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against number of sequencing depth. (A) C57BL/6J mice were randomly divided into four groups: control, chronic ethanol, binge, and chronic plus binge ethanol. Fecal pellets were collected 9 h after the oral gavage. Rarefaction curves for all gut microbiota samples sequenced from male mice groups. (B) Rarefaction curves for all gut microbiota samples sequenced from female mice groups. Each colored line represents one biological sample

35 of male or female mice groups (n=5 per group). FB: female from binge groups; FC: 36 female from control group; FE: female from chronic ethanol group; FEB: female from 37 chronic plus binge ethanol group; MB: male from binge groups; MC: male from control 38 group; ME: male from chronic ethanol group; MEB: male from chronic plus binge ethanol 39 group. 40 41 Figure S4. Ethanol-induced changes in the relative abundance of major gut microbiota 42 phyla in male and female mice. C57BL/6J mice were randomly divided into four groups: 43 control, chronic ethanol, binge, and chronic plus binge ethanol. Fecal pellets were 44 collected 9 h after the oral gavage. Bar plots illustrate the relative abundance of major 45 bacterial phyla in male (A) and female (B) groups. Results are expressed as the mean ± 46 S.E.M. (n=5 per group). 47 48 Figure S5. Top 30 detected pathways across the samples. C57BL/6J mice were randomly 49 divided into four groups: control, chronic ethanol, binge, and chronic plus binge ethanol. 50 Fecal pellets were collected 9 h after the oral gavage. FB: female from binge group; FC: 51 female from control group; FE: female from chronic ethanol group; FEB: female from 52 chronic ethanol plus binge group; MB: male from binge group; MC: male from control 53 group; ME: male from chronic ethanol group; MEB: male from chronic ethanol plus binge 54 group. 55 56 Figure S6. Predicted functional profile of male mice microbiota. C57BL/6J mice were 57 randomly divided into four groups: control, chronic ethanol, binge, and chronic plus binge 58 ethanol. Fecal pellets were collected 9 h after the oral gavage. Functional profiling of the 59 gut microbiome was performed using PICRUSt2. The Kruskal-Wallis non-parametric test 60 was used to determine statistically significant differences among the groups (p<0.05). 61 PICRUSt2: Phylogenetic Investigation of Communities by Reconstruction of 62 Unobserved States 2. 63 64 Figure S7. Predicted functional profile of female mice microbiota. C57BL/6J mice were randomly divided into four groups: control, chronic ethanol, binge, and chronic plus binge 65 ethanol. Fecal pellets were collected 9 h after the oral gavage. Functional profiling of the 66 67 gut microbiome was performed using PICRUSt2. The Kruskal-Wallis non-parametric test 68 was used to determine statistically significant differences among the groups (p<0.05).

69 PICRUSt2: Phylogenetic Investigation of Communities by Reconstruction of

70 Unobserved States 2.

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12	LEGENUS FOR SUPPLEMENTARY TABLES
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74	Table S1. Pathways identified in the mouse fecal microbiome
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76	Table S2. Primers sequences used in real-time PCR.
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