



29 **Abstract**

30 Although gut microbiota and lipid metabolites have been suggested to be closely  
31 associated with type 2 diabetes mellitus (T2DM), the interactions between gut  
32 microbiota, lipid metabolites and the host in T2DM development remains unclear.  
33 Rhesus macaques may be the best animal model to investigate these relationships  
34 given their spontaneous development of T2DM. We identified eight spontaneous  
35 T2DM macaques and conducted a comprehensive study investigating the  
36 relationships using multi-omics sequencing technology. Our results from 16S rRNA,  
37 metagenome, metabolome and transcriptome analyses identified that gut microbiota  
38 imbalance, tryptophan metabolism and fatty acid  $\beta$  oxidation disorders, long-chain  
39 fatty acid (LCFA) accumulation, and inflammation occurred in T2DM macaques. We  
40 verified the accumulation of palmitic acid (PA) and activation of inflammation in  
41 T2DM macaques. Importantly, mice transplanted with spontaneous T2DM macaque  
42 fecal microbiota and fed a high PA diet developed prediabetes within 120 days. We  
43 determined that gut microbiota mediated the absorption of excess PA in the ileum,  
44 resulting in the accumulation of PA in the serum consequently leading to T2DM in  
45 mice. In particular, we demonstrated that the specific microbiota composition was  
46 probably involved in the process. This study provides new insight into interactions  
47 between microbiota and metabolites and confirms causative effect of gut microbiota  
48 on T2DM development.

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50 **Key words:** spontaneous T2DM macaques, multi-omics, gut microbiota, palmitic  
51 acid, fecal microbiota transplantation

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## 59 **Introduction**

60 Diabetes mellitus is considered to be a refractory disease causing a significant  
61 socio-economic burden. Type 2 diabetes mellitus (T2DM) is the dominant type of  
62 diabetes mellitus characterized by metabolic disorders, insulin resistance, and  
63 deficiency of insulin secretion (1, 2). The pathogenesis of this chronic disease is  
64 complex and genetic and environmental factors, such as sugar and lipid intake, gut  
65 microbiota, many metabolites, and even air pollutants, contribute to its increase in  
66 prevalence (3, 4). Accumulating evidence has linked gut microbiota with T2DM  
67 development by variety ways. The gut microbiota can impact the integrity of the  
68 intestinal epithelial barrier, mediate insulin resistance, as well as regulate the function  
69 of mitochondria (5, 6, 7). They can regulate local or systemic immunity and  
70 inflammation, which also contributes to the development of T2DM (8). Moreover,  
71 various gut microbial metabolites, such as short-chain fatty acids, bile acid, and  
72 tryptophan-derived metabolites, have been reported to be closely related to the  
73 pathogenesis of T2DM (5, 6, 9, 10). However, the interactions between gut microbiota  
74 and its host with T2DM have not yet been fully characterized. In pathological  
75 conditions, the dysregulation of the host can lead to changes in gut microbiota  
76 composition. In turn, the microbiota plays a regulatory role to participate in the  
77 development of T2DM. Despite the complicate interactions between host and  
78 microbiota in the context of T2DM, some studies suggest antidiabetic interventions  
79 targeting the gut microbiota such as fecal microbiota transplanting (FMT) can be  
80 applied as a clinical treatment of T2DM (11, 12, 13).

81       Meanwhile, dysfunction of lipid metabolism contributes to T2DM development  
82 by inducing lipotoxicity in humans and animal models (14, 15). The long-chain fatty  
83 acids (LCFAs) are the principal lipid components naturally occurring in animal fats  
84 and vegetable oil, as well as the main metabolites of fat. LCFAs such as palmitic acid  
85 (PA, C16:0), palmitoleic acid (C16:1N7), and oleic acid (C18:1N9), are reported to  
86 have strong association with T2DM. In the last few decades, there is increasing

87 evidence that frequent consumption of LCFAs contributes to metabolic diseases such  
88 as obesity and T2DM due to the high PA content (16, 17). PA is a saturated fatty acid  
89 and its increase in serum is a significant contributing factor in T2DM development  
90 (18, 19, 20). The suggested mechanisms by which PA mediates T2DM include  
91 increasing diacylglycerol and ceramide synthesis (21), mitochondrial and  
92 endoplasmic reticulum stress (22, 23), and activation of pro-inflammatory pathways  
93 (24). Nevertheless, whether gut microbiota involving in PA mediated T2DM and the  
94 interactions between gut microbiota and LCFA metabolites in T2DM development are  
95 still unclear.

96 Spontaneous development of T2DM in non-human primates (e.g. macaques) is  
97 highly similar to human T2DM, such as insulin resistance in early stages and later  
98 abnormal glucose tolerance and T2DM development follows the same pathological  
99 changes of pancreatic islets and complications. In fact, T2DM macaques and humans  
100 have amyloidosis of pancreatic islets but it is not observed in mouse models (25, 26),  
101 suggesting that T2DM macaques are the best animal model that can simulate and  
102 reproduce human T2DM and its complications (27). However, previous studies  
103 indicated that naturally occurring spontaneous T2DM macaques in captive  
104 populations were rare even if individuals were given a high lipid and high sugar diet  
105 (28, 29). After nasally fed cynomolgus macaques with a high-fat dietary emulsion for  
106 12 months, the macaques did not experience significant increases in fasting blood  
107 glucose and glycosylated hemoglobin (29). The controversial effects of high fat on the  
108 T2DM development are worthy of further investigations to better understand this  
109 complex disease.

110 Here, this study identified spontaneous development of T2DM in individuals  
111 (hereafter spontaneous T2DM macaques) from a large group of rhesus macaques.  
112 These spontaneous T2DM macaques have never been treated with anti-diabetic drugs  
113 and therefore provide valuable models for pathogenesis investigation of T2DM.  
114 Based on the macaque model, we used multi-omics techniques to address the  
115 interactions between gut microbiota, host gene expression, and fecal metabolites and

116 the development of T2DM. Our results demonstrated that gut microbiota and LCFA  
117 metabolites played important roles in the pathogenesis of T2DM. We validated the  
118 increased content of plasma PA and activation of inflammation in the T2DM  
119 macaques. In addition, we successfully induced prediabetes in mice by transplanting  
120 fecal microbiota from T2DM macaques into mice in conjunction with a diet high in  
121 PA. We also revealed the specific structure of gut microbiota that promoted T2DM  
122 development by regulating the absorption of excess PA, which provides key evidence  
123 of the causative effect of gut microbiota on the pathogenesis of T2DM.

## 124 **Results**

### 125 **Metagenome and 16S sequencing demonstrate alterations on gut** 126 **microbiota in spontaneous T2DM macaques**

127 We identified eight spontaneous T2DM macaques out of 1698 individuals from  
128 long-term glucose monitoring in a captive population (Table S1). The fasting plasma  
129 glucose (FPG), fasting plasma insulin (FPI), and HOMA-IR levels in the T2DM  
130 macaques were significantly higher than in the control group ( $p < 0.01$ ), suggesting  
131 insulin resistance in T2DM macaques. However, glycosylated hemoglobin A1c  
132 (HbA1c), triglycerides (TG), total cholesterol (TC), high-density lipoprotein  
133 cholesterol (HDL), low-density lipoprotein cholesterol (LDL), and BMI did not  
134 significantly differ from the controls ( $p > 0.05$ , Table 1). Comparison of gut microbiota  
135 between the two groups based on 16S rRNA amplicon found that both Shannon and  
136 Simpson index decreased in T2DM macaques but it was not significantly different  
137 from controls ( $p > 0.05$ ; Figures 1A and B). PCoA indicated that the microbiota  
138 composition of spontaneous T2DM macaques was different from the control group  
139 (Figure 1C). As one of the most dominant families in both groups, Lachnospiraceae  
140 showed significantly higher abundance in the T2DM group (14.48%) than in the  
141 control group (6.66%). While the abundance of the Lactobacillaceae family was  
142 significantly greater in the control group (25.95%) compared to spontaneous T2DM  
143 macaques (20.88%) (Figure 1D). A total of 21 microbes were identified as

144 differentially microbes between the T2DM group and control group, where ten  
145 microbes including Erysipelotrichaceae and five members in Lachnospiraceae family  
146 (*Ruminococcus gnavus*, *Lachnospira*, *Coprococcus* sp. *Dorea longicatena*, and  
147 *Roseburia*) were significantly greater in the T2DM group (Figure 1E).

148 The metagenome results showed that the T2DM group had a higher abundance  
149 of Erysipelotrichaceae, *Eubacterium rectale*, Lachnospiraceae, Negativicutes, *Blautia*,  
150 and Coriobacteriia than the control group (Figure S1A). Functional enrichment  
151 demonstrated a total of 74 KEGG pathways with significant differences between the  
152 two groups. These pathways were mainly associated with T2DM and inflammation,  
153 including type II diabetes mellitus, glucagon signaling pathway, starch and sucrose  
154 metabolism, lipid metabolism, and toll-like receptor signaling pathways (Figure 1F).  
155 Among the six CAZy enzyme families, the Carbohydrate-binding module (CBMs),  
156 Glycosyl Transferases (GTs), Carbohydrate esterases (CEs), and Polysaccharide  
157 Lyases (PLs) families were significantly upregulated in the T2DM group, indicating  
158 significant changes in carbohydrate metabolism (Figure 1G). The results of  
159 metagenome and 16S sequencing demonstrated significant alterations in the  
160 composition and function of gut microbiota in spontaneous T2DM macaques, with a  
161 greater abundance of microbes associated with T2DM and fewer beneficial microbes.

## 162 **Fecal metabolome and blood transcriptome reveals dysfunction of** 163 **fatty acid $\beta$ oxidation and tryptophan metabolism in T2DM** 164 **macaques**

165 The UHPLC-MS based metabolome analysis on fecal samples of T2DM and control  
166 macaques identified 1564 metabolites belonging to various types of secondary  
167 metabolites, with lipids and lipid-like molecules being most abundant (31.1%) (Figure  
168 S1B). We found 64 significantly differential metabolites between the two groups  
169 using a combined multidimensional statistical analysis (OPLS-DA) and univariate  
170 statistical analysis (T-test) ( $VIP > 1$ ,  $p < 0.05$ ) (Figures 2A and B; Table S2). Among  
171 them, muscon, indole-3-acetaldehyde, and serotonin were significantly lower in the

172 T2DM group (Figure 2C) and are associated with anti-inflammatory activity (30, 31,  
173 32). Notably, 64 differential metabolites were significantly enriched in one pathway:  
174 tryptophan metabolism (Figure 2D), and we identified two significantly different  
175 metabolites, indole-3-acetaldehyde and serotonin in this pathway. Both metabolites  
176 are microbiota-derived tryptophan metabolites and the ligands for AhR (31, 32). The  
177 lower concentration of AhR ligands may lead to the development of inflammation and  
178 metabolic syndromes in humans (33). In addition, we found the contents of many  
179 acylcarnitine metabolites were significantly higher in the T2DM group (VIP>1,  
180  $p<0.1$ ), including l-propionylcarnitine, hexanoyl-l-carnitine, (r)-butyrylcarnitine, and  
181 isovaleryl-l-carnitine (Table S2). Among them, l-propionylcarnitine, a kind of  
182 acylcarnitines, was the most upregulated metabolite in the T2DM macaques (FC:  
183 16.19) (Figure 2C). Acylcarnitines are products of incomplete oxidation of LCFAs,  
184 which can activate pro-inflammatory signaling pathways and ultimately inhibit insulin  
185 activity (34). Our results indicated incomplete LCFAs  $\beta$  oxidation in spontaneous  
186 T2DM macaques, while similar characteristics were also observed in insulin resistant  
187 and T2DM humans (35, 36).

188 Blood transcriptome analysis was consistent with metabolome results, indicating  
189 dysfunction of fatty acid  $\beta$  oxidation and inflammation in T2DM macaques. Gene  
190 expression in T2DM macaques exhibited significant differences from the controls  
191 (Figure 2E), and a total of 161 differentially expressed genes (DEGs) (26 upregulated  
192 and 135 downregulated) were identified in T2DM macaques at a FDR level of 0.05  
193 (Figure 2F). Enrichment analysis of the DEGs was linked to diabetes, fatty acid  
194 metabolism, and inflammation, such as diabetic cardiomyopathy, glucose homeostasis,  
195 fatty acid metabolic process, and chemokine production (Figure 2G). We also  
196 identified 26 differential enrichment pathways between the two groups by aggregate  
197 fold change (AFC), and most were associated with insulin resistance and  
198 inflammation, including insulin resistance, PI3K-Akt signaling pathway, bacterial  
199 invasion of epithelial cells, and NOD-like receptor signaling pathway (Figure S2A).  
200 As shown in the insulin resistance pathway, expression of *IL6* and *IRS1* were

201 upregulated, while *INSR* was downregulated in T2DM macaques (Figure S2B). The  
202 WGCNA analysis identified four modules that were related to T2DM. Darkgreen  
203 module and brown module were significantly positively correlated with T2DM, while  
204 green module and darkred module were negatively correlated with T2DM (Figure 2H).  
205 Genes in the darkgreen module and brown module related to lipolysis and  
206 inflammation were significantly upregulated in T2DM macaques, and genes in the  
207 green module and darkred module related to fatty acid metabolism and insulin  
208 secretion were significantly downregulated (Figure S2C). With the cut-off  
209 ( $|kME| > 0.8$ ), a total of 546 genes in the four modules were identified as hub genes  
210 (Table S3). Of these 546, 102 hub genes were also DEGs (Figure 2I). Several of these  
211 genes have been reported as correlated with T2DM in humans, including *IGF2BP2*,  
212 *LEPR*, *RAP1A*, *SESTRIN 3*, and *ITLN1* (37, 38, 39, 40, 41). In addition, *ACSM3*,  
213 *HADHB*, and *EFHB* are involved in fatty acid  $\beta$  oxidation and inflammation (42, 43,  
214 44).

## 215 **Validation of LCFAs accumulation and inflammation in T2DM** 216 **macaques**

217 Collectively, metabolome and transcriptome results indicated dysfunction in fatty acid  
218  $\beta$  oxidation and tryptophan metabolism in T2DM macaques, which may lead to LCFA  
219 accumulation and inflammation. To support this conclusion, we performed targeted  
220 medium-and long-chain fatty acid mass spectrometry of plasma and examined serum  
221 inflammatory cytokines in the macaques. A total of 34 fatty acids were detected, and  
222 among the five types of fatty acids, the concentration of saturated fatty acid (SFA)  
223 was significantly greater in the T2DM group ( $p < 0.05$ , Figure 3A), while other types  
224 of fatty acids were not significantly different between the two groups ( $p > 0.05$ , Figures  
225 3B-E). In particular, concentrations of PA, palmitoleic acid, and oleic acid were  
226 significantly higher in the T2DM group than control group ( $p < 0.05$  and  $VIP > 1$ ,  
227 Figures 3F and G). Increased content of PA, palmitoleic acid, and oleic acid was also  
228 found in human T2DM (45, 46). PA was an important metabolite mediating insulin

229 resistance through three main mechanisms, being increased diacylglycerol and  
230 ceramide synthesis, mitochondrial and endoplasmic reticulum stress, and activation of  
231 pro-inflammatory pathways through membrane receptors (21, 22, 23, 24). Analysis on  
232 the serum inflammatory cytokines found that IL-1 $\beta$  was significantly higher in the  
233 T2DM group ( $p < 0.05$ , Figure 3H), but TNF- $\alpha$  and IL-6 levels showed no significant  
234 difference between the two groups ( $p > 0.05$ , Figures 3I and J). IL-1 $\beta$  is a major player  
235 in a variety of autoinflammatory diseases and a key promoter of T2DM systemic and  
236 tissue inflammation (47). Moreover, blood routine examination showed an increase of  
237 WBC number, NEU percentage and NEU number and a decrease of LYM percentage  
238 and LYM number, also indicating the inflammation in the T2DM macaques ( $p < 0.05$ ,  
239 Table 2).

240 To investigate effect of gut microbiota on the LCFAs accumulation and  
241 inflammation in T2DM macaques, we performed a correlation analysis among the  
242 DEGs, differential metabolites, and differential microbes using spearman rank  
243 correlation. Four differential microbes in Lachnospiraceae family (*Coprococcus*,  
244 *Lachnospira*, *Roseburia* and *Dorea longicatena*) were significantly associated with  
245 three differential metabolites of PA, palmitoleic acid and oleic acid ( $|r| > 0.5$ , adj  
246  $p < 0.05$ ), suggesting the participation of Lachnospiraceae microbes in LCFAs  
247 accumulation in T2DM macaques (Figure 3K). In addition, the bacteria in class  
248 Coriobacteriia was also associated with the three LCFAs ( $|r| > 0.5$ , adj  $p < 0.05$ , Figures  
249 S1C and D).

## 250 **Fecal microbiota transplantation (FMT) with high content PA food** 251 **induce prediabetes in mice**

252 To determine the causative effect of gut microbiota and PA on T2DM development,  
253 we collected feces from the spontaneous T2DM macaques and performed fecal  
254 microbiota transplantation (FMT) in antibiotic-pretreated mice. Mice were either  
255 administrated by FMT (FT), fed with high concentration PA diet (PA), or were  
256 combined FMT and PA diet (FTPA). A control group was used and they were fed with

257 normal commercial food and lacked FMT (Figure 4A). FPG monitoring found that  
258 FPG levels in the FTPA group and FT group increased continuously from day 60  
259 (Figure 4B), while the control group maintained stable FPG levels throughout the 120  
260 days. The FTPA group showed the highest FPG of 6.7 mmol/L at day 120, which was  
261 significantly higher than the control group and indicated FTPA-treated mice had  
262 developed prediabetes (Figure 4C). The oral glucose tolerance test (OGTT) showed  
263 obvious glucose intolerance in the FTPA group (Figures 4D and E), and the FPI  
264 values and insulin tolerance test (ITT) were both significantly elevated in the FTPA  
265 group, indicating insulin resistance (Figures 4F and G). Meanwhile, body weight  
266 (BW), TC, and TG significantly increased in the FTPA group (Figures 4H-J). While  
267 mice in the FT group and PA group had significant weight gain and mild insulin  
268 resistance, they did not exhibit significant glucose intolerance and no significant  
269 elevation in TC and TG levels compared to controls. Histopathological changes in the  
270 pancreas and liver of mice were investigated using H-E staining. Hepatocytes focal  
271 necrosis with inflammatory cell infiltration was commonly observed in the FTPA  
272 mice, but not frequent in hepatocytes in FT and PA groups (Figure 4K). Furthermore,  
273 decreased volume and area in pancreatic islets and inflammatory cell infiltration were  
274 detected in the FTPA mice (Figure 4L), while such pathological changes were not  
275 found in the control group.

## 276 **Specific structure of gut microbiota mediates the absorption of excess**

### 277 **PA in the ileum**

278 Given that the content of plasma PA significantly increased in the T2DM macaques  
279 (Figures 3F and G), we examined PA content in the feces, ileum, and serum in mice to  
280 compare the prediabetes mice and the spontaneous T2DM macaques. PA content in  
281 the serum and ileum of the FTPA group was significantly higher than the control  
282 group ( $p < 0.05$ , Figures 5A and B), but its content in feces was significantly lower  
283 than the control group ( $p < 0.05$ , Figure 5C). This suggested that the absorption of PA  
284 was significantly enhanced in the ileum leading to the increase of PA in serum. To

285 verify this conclusion, expression of the *Cd36* gene, a gene involved in the uptake and  
286 oxidation of LCFAs (48), was examined in the ileum of mice. Ileums showed a  
287 significantly upregulated expression of *Cd36* in the FTPA group compared to the  
288 control group ( $p < 0.05$ , Figure 5D). In contrast, the level of IL-17A, a protein  
289 inhibiting the expression of *Cd36* (49), was significantly reduced in the ileum of the  
290 FTPA group ( $p < 0.05$ , Figure 5E). Interestingly, the PA group's content of PA, *Cd36*  
291 expression and IL-17A was not significantly different from the control group. This  
292 indicated that without FMT, the ileum could not absorb PA effectively even when fed  
293 in high concentrations. Consequently, gut microbiota mediated the absorption of  
294 excess PA in the ileum.

295 We then compared diversity and composition of microbiota communities  
296 between the FTPA, FT, PA and control groups, and a fifth group consisting of the  
297 microbiota transplants (TP) from T2DM macaques. Shannon and Simpson indices  
298 were lower in the FTPA, FT and PA groups than the control group (Figures S3A and  
299 B). NMDS analysis also indicated a distinct microbiota composition from the control  
300 group compared to other groups (NMDS1). In addition, microbiota composition of TP  
301 from T2DM macaques was distinct from other groups but a little closer to the FTPA  
302 and FT groups (NMDS2, Figure 5F). In particular, the Lachnospiraceae family  
303 showed the higher abundance in the TP, FTPA and FT groups than in the PA and  
304 control groups (Figure S3C). The abundances of three members of microbes in  
305 Lachnospiraceae (*R. gnavus* and *Coprococcus* sp. and *Clostridium*) in the FTPA and  
306 FT groups gradually increased over time after FMT (Figure S3D). The FTPA and FT  
307 groups shared many differential microbes compared to the control group, such as the  
308 significantly upregulated *R. gnavus* and *Coprococcus* sp., and the significantly  
309 downregulated Christensenellaceae, F16, *Treponema* sp. and *Fibrobacter*  
310 *succinogenes* (Figures 5G and H). It is noteworthy that these microbes were also  
311 differential microbes between T2DM macaques and controls, and their abundances  
312 changed in the same way as in macaques (Figure 1E). However, mice in the PA group  
313 did not share differential microbes with the spontaneous T2DM macaques.

314 Correspondingly, the change of serum PA content in the PA group was not  
315 significantly higher than the control group (Figure S3E). Our results suggested that  
316 the transplanted microbiota from spontaneous T2DM macaques, especially the  
317 increased abundance of *R. gnavus* and *Coprococcus* sp. and decreased abundance of  
318 *Treponema*, *F. succinogenes*, Christensenellaceae and F16, promoted the absorption of  
319 excess PA by regulating the expression of IL-17A and Cd36, leading to the LCFAs  
320 accumulation and insulin resistance (Figures 5I).

## 321 Discussion

322 With a multi-omics technology, this study comprehensively characterizes the gut  
323 microbiota, metabolites and gene expression of spontaneous T2DM macaques. The  
324 gut microbiota diversity in T2DM macaques decreased. In particular, the abundance  
325 of bacteria *R. gnavus* and Erysipelotrichaceae were upregulated while the abundance  
326 of Christensenellaceae was downregulated in the T2DM macaques. Metabolome  
327 results demonstrated a decrease of microbiota-derived tryptophan and  
328 anti-inflammatory metabolites, indicating that T2DM macaques were prone to  
329 inflammation. Notably, the accumulation of acylcarnitine metabolites, the suggested  
330 biomarkers for human T2DM (46), indicated incomplete mitochondrial LCFA  $\beta$   
331 oxidation in T2DM macaques. Transcriptome results identified many DEGs linked to  
332 insulin resistance, fatty acid  $\beta$  oxidation and inflammation, including *IGF2BP2*, *LEPR*,  
333 *RAP1A*, *SESTRIN 3*, and *ITLN1* that have also been reported in human T2DM (37, 38,  
334 39, 40, 41). Combining the multi-omics results we revealed the complex pathological  
335 mechanisms in the spontaneous T2DM macaques (Figure 6), which is comparable to  
336 T2DM humans. Firstly, expression of genes related to lipolysis, fatty acid oxidation,  
337 LCFA accumulation, inflammation, and insulin secretion are dysregulated, promoting  
338 the development of T2DM (Figure 6A). This is particularly related to lipid  
339 metabolism, where the increase of lipolysis and the downregulated expression of fatty  
340 acid metabolism-related genes *HADHB* and *ACSM3* cause the accumulation of  
341 acylcarnitine and LCFAs, which results in incomplete LCFA oxidation in T2DM  
342 macaques (Figure 6B). We also suggest that the decrease of *Lactobacillus* sp. cause

343 the reduction of Aryl hydrocarbon receptor (AhR) ligands (serotonin and  
344 indole-3-acetaldehyde) given that *Lactobacillus* is producer of AhR ligands (50). This  
345 then promotes the expansion of the PA producer Erysipelotrichacea (51, 52) and  
346 ultimately leads to the increase of PA levels. The increase in the abundance of  
347 Erysipelotrichacea and *R. gnavus* also promotes the development of T2DM by  
348 activation of inflammation (53, 54, 55) (Figure 6C). The accumulation of PA was  
349 reported to lead to insulin resistance by affecting genes in the insulin signaling  
350 pathway (16). Our study demonstrates that the significantly changed expression of  
351 *RAP1A*, *SESTRIN3*, and *IRS1* in PA-mTORC1-Akt pathway causes insulin resistance  
352 in T2DM macaques. Moreover, the increase of PA can promote the development of  
353 T2DM by upregulating the NF- $\kappa$ B signaling pathway (Figure 6D).

354 We validated the observations from the multi-omics analysis finding the  
355 significantly higher inflammatory cytokines IL-1 $\beta$  and LCFA accumulation,  
356 especially significant PA accumulation in the T2DM macaques. And found LCFA  
357 metabolites were significantly correlated with bacteria in the Lachnospiraceae family.  
358 Numerous studies imply an association of gut microbiota with T2DM development.  
359 By transplanting gut microbiota from healthy individuals to T2DM individuals,  
360 symptoms such as insulin resistance or inflammation were improved (6, 56). However,  
361 there is no evidence to date suggesting gut microbiota have directly causative effects  
362 on T2DM development. Our study confirmed the causative effect of gut microbiota  
363 and PA on T2DM development by transplanting fecal microbiota from spontaneous  
364 T2DM macaques to antibiotic pretreated mice. We successfully induced prediabetes in  
365 mice after combining FMT administration and high PA ingestion. However, when the  
366 treatments were administered on their own, the mice did not develop prediabetes. We  
367 determined, for the first time, that gut microbiota mediated the absorption of excess  
368 PA in the ileum by quantitative examining PA contents in feces, ileum, and serum, and  
369 analysis of the expression of *Cd36* and IL-17A level in the ileum. Most notably, this  
370 then resulted in accumulation of PA in the serum and finally led to T2DM  
371 development. Without the transplanting gut microbiota, the ileum could not absorb the

372 PA effectively even at a high concentration of ingested PA. Our study highlights the  
373 essential roles of gut microbiota in T2DM development, and this is probable also the  
374 reason that previous studies have failed to induce T2DM in macaques because they  
375 have only used a high-fat diet (28, 29).

376 We then determined the specific gut microbiota structure that related to T2DM  
377 development in the prediabetes mice and spontaneous T2DM macaques. We found  
378 that the increased abundance of *R. gnavus* and *Coprococcus* sp., and the decreased  
379 abundance of *Treponema*, *F. succinogenes*, Christensenellaceae, and F16, were  
380 involved in the T2DM development. *R. gnavus* can promote insulin resistance by  
381 regulating the content of tryptamine/phenethylamine (57). Moreover, *R. gnavus* is a  
382 mucin-degrading microbe that leads to an increase of inflammation (58, 59, 60). The  
383 intestinal mucous layer is an important barrier separating intestinal tissue from  
384 microbiota, and microbiota composition plays a major role in affecting the integrity of  
385 intestinal mucous layer (58). The increase of *R. gnavus* suggested a higher risk of  
386 damage to the integrity of the intestinal mucous layer. This is further supported by our  
387 results of the lower abundance of *Lactobacillus* sp., which are AhR ligand producers,  
388 and the decreased content of AhR ligands (tryptophan microbial metabolites) in  
389 spontaneous T2DM macaques. The deficiency of AhR ligand reduced the production  
390 of intestinal mucus and increased the risk of microbial invasion, which in turn  
391 affected the immune cell differentiation and cytokine production (61, 62). The  
392 cytokine IL-17A is a regulator of the fatty acid transporter *Cd36* and lipid absorption  
393 can be promoted by reducing the inhibition of *Cd36* expression by IL-17A (54). The  
394 decreased IL-17A level and increased *Cd36* expression in prediabetes mice indicated  
395 that the specific gut microbiota promoted the absorption of excess PA by disrupting  
396 the integrity of the intestinal mucous layer and regulating the expression of IL-17A  
397 and *Cd36*. In addition, the beneficial bacteria decreased in abundance in T2DM  
398 macaques and prediabetes mice. These beneficial bacteria, such as *F. succinogenes*,  
399 Christensenellaceae, and F16, protect the mucosal barrier and improved insulin  
400 resistance (63, 64, 65). We inferred the collectively effects of these gut microbes

401 determined the absorption of excess PA from ileum to serum, which might contribute  
402 to the development of T2DM.

403 In conclusion, spontaneous T2DM macaques that have never been treated with  
404 diabetes-related drugs provide a valuable model for our understanding of the  
405 pathological characteristics and pathogenesis of T2DM. This study characterized  
406 changes in gene expression, metabolites, and gut microbiota levels of spontaneous  
407 T2DM macaques using multi-omics techniques. We found gut abnormal microbiota,  
408 tryptophan metabolism and fatty acid  $\beta$  oxidation disorders, inflammation, and PA  
409 accumulation. We also successfully induced prediabetes in mice by transplanting fecal  
410 microbiota from T2DM macaques into antibiotic pretreated mice fed a high PA diet.  
411 Our study confirms the causative effect of gut microbiota composition change and PA  
412 in T2DM development. The microbiota composition, specifically higher abundance of  
413 *R. gnavus* and *Coprococcus* sp. and lower abundance of *Treponema*, *F. succinogenes*,  
414 Christensenellaceae and F16, promoted the absorption of excess PA thus led to the  
415 development of T2DM. This study provides new insights into the interaction of gut  
416 microbiota and metabolites in the development of T2DM, which expands our  
417 understanding of the pathogenesis of this metabolic disease and may provide novel  
418 insights for the treatment of T2DM in the future.

## 419 **Materials and methods**

### 420 **The screening of spontaneous T2DM macaques**

421 The experimental macaques used in this study were all from Greenhouse  
422 Biotechnology Co., LTD (Meishan, China). We obtained eight spontaneous T2DM  
423 macaques with  $FPG \geq 7$  mmol/L and eight healthy control macaques with  $FPG \leq 6.1$   
424 mmol/L (three consecutive detections, each detection interval of one month) from a  
425 population of 1698 captive macaques. None of these 16 screened macaques received  
426 any medical treatment for diabetes.

### 427 **Sample collection**

428 Each experimental macaque was kept in a single cage, and was fasted for 12 hours but  
429 had free access to drinking water. We obtained serum, plasma and whole blood for the

430 detection of physiological and biochemical parameters, and metabolome and  
431 transcriptome analysis. Fecal samples were collected within ten minutes after  
432 deposition. During the sampling process, fecal samples were loaded into a 50 mL  
433 sterile centrifuge tube were stored at -80°C. We followed animal welfare guidelines  
434 throughout the sample collection process, and all observations and samplings were  
435 approved by the Sichuan University's Animal Care Committee.

#### 436 **Physiological and biochemical parameters**

437 In this study, FGP, HbA1c and FPI were detected by hexokinase method, high  
438 performance liquid chromatography and electro-chemiluminescence method,  
439 respectively. TC, TG, HDL and LDL were detected by automatic biochemical  
440 analyzer. HOMA-IR is one of the criteria for T2DM, calculated as:  $(FPI \times FPG) / 22.5$ .

#### 441 **Feces 16S rRNA amplicon sequencing and analysis**

442 Total DNA from fecal samples was extracted using the QIAamp Fast DNA Stool Mini  
443 Kit. The V3-V4 region of the 16S rRNA gene was amplified using the 341F/806R  
444 primer (341F: 5'-CCTAYGGGRBGCASCAG -3', 806R:  
445 5'-GGACTACNNGGGTATCTAAT -3'). The purification and quantification of the  
446 amplified products were performed and followed by the sequencing library  
447 preparation with TruSeq Nano DNA LT Library Prep Kit (Illumina, USA). The library  
448 sequencing was performed on Illumina MiSeq platform and 250 bp paired-end reads  
449 were generated. Raw sequencing reads were merged using FLASH (V1.2.7,  
450 <http://ccb.jhu.edu/software/FLASH/>) (66) and analyzed by QIIME2 (version  
451 2020.11.1) pipeline with default parameters. Reads after denoising by DADA2 were  
452 clustered into OTUs at 99% similarity threshold. Taxonomy of OTUs was assigned  
453 based on Greengenes reference database (67). The QIIME2 diversity plugin was used  
454 to calculate alpha diversity (68). Principal Coordinate Analysis (PCoA) was  
455 determined by using the R package vegan. Differentially microbes were determined  
456 using linear discriminant analysis effect size (LEfSe) (69).

#### 457 **Feces shotgun metagenome sequencing and analysis**

458 Total DNA from each sample was extracted using Tiangen DNA Stool Mini Kit

459 (Tiangen Biotech Co., Ltd., China). Metagenome sequencing was performed using the  
460 Illumina NovaSeq 6000 platform with a paired-end sequencing length of 150 bp.  
461 Trimmomatic was used for removing the adapters and low-quality raw reads based on  
462 a four-base wide sliding window, with average quality per base >20 and minimum  
463 length 90 bp (70). The rhesus macaque potential sequences were removed using  
464 Bowtie2 (71) with the reference genome (assembly Mmul\_10). Taxonomy of  
465 remaining reads was assigned using Kraken2 (72) with the option “--use-mpa-style”.  
466 *De novo* assembly of remaining reads was performed using MEGAHIT (73) with the  
467 option “-m 0.90 --min-contig-len 300”. Prodigal (74) was used for gene prediction.  
468 The construction of non-redundant gene catalogue was performed with CD-HIT (75)  
469 with the option “-c 0.95 -aS 0.90”. Quantification of the non-redundant genes in each  
470 sample was performed using Salmon (76). The amino acid sequences translated from  
471 non-redundant genes were aligned (--id 80% --query-cover 70% --evaluate 1e-5) by  
472 DIAMOND (77) in the Carbohydrate-Active enZymes (CAZy) database (78). The  
473 annotation metabolic pathway was performed using HUMANN3 (79).

#### 474 **RNA sequencing and DEG analysis**

475 Total RNA was extracted using PAXgene Blood RNA kit. The cDNA Library was  
476 constructed following the NEBNext® Ultra™ RNA Library Prep Kit for Illumina®  
477 (NEB, USA) manual, and index was added to each sample for sample differentiation.  
478 cBot Cluster Generation System was used to cluster the sequences with the same  
479 index. Illumina HiSeq 2500 sequencing platform was used to obtain the paired-end  
480 sequencing reads (PE150). NGS QC Toolkit v2.3.3 (80) was used to obtain high  
481 quality reads (high-quality paired reads with more than 90% of bases with  
482 Q-value ≥ 20 were retained). Processed reads of each sample were mapped to the  
483 macaque reference genome using HISAT2 v2.1.0 (Kim, et al. 2015). Each alignment  
484 output file was assembled into a separate transcriptome using StringTie v1.3.6 (81),  
485 resulting in a transcript GTF file. To obtain the expression value of TPM (transcripts  
486 per million) and raw read counts of each gene, transcript GTF file was used as the  
487 reference annotation file. Differential expression analysis was performed using

488 DESeq2 R package (82). DEGs were screened according to Foldchange value (FC)  
489 and p value corrected by FDR (Benjamini-Hochberg method was selected) (adj  
490  $p < 0.05$ ,  $|\log_2FC| > 1$ ).

491 Weighted Gene Co-Expression Network Analysis (WGCNA) (83) was used to  
492 analyse the correlation between genes and phenotypes. We selected an appropriate  
493 “soft thresholding power” using the “picksoftthreshold” function in the WGCNA  
494 package (v1.61). Next, “blockwiseModules” function was used to construct the co-  
495 expression matrix with the option “checkMissingData=TRUE, power=16,  
496 TOMType='unsigned', minModuleSize=30, maxBlockSize=6000,  
497 mergeCutHeight=0.25”. The modules with high correlation to T2DM phenotype and  
498  $p < 0.05$  were selected for downstream analysis, and genes in these modules were used  
499 for GO and KEGG functional enrichment analysis. Hub genes were identified based  
500 on the module eigengene-based connectivity (kME),  $|kME| > 0.8$  as the cut-off criteria.

501 Instead of identifying the differentially expressed genes within a pathway  
502 between the two groups, Aggregate Fold Change (AFC) calculated the average  
503 multiple change for each gene and defined the pathway score as the average  
504 difference multiple for all genes in that pathway. A null hypothesis test was performed  
505 using the pathway scores of the gene expression dataset, and the significance of each  
506 pathway was estimated by p-value (84). STRING (v11.0) provides a tool for  
507 functional enrichment analysis based on AFC. GO and KEGG enrichment analysis  
508 was performed using g: Profiler (85), AFC enrichment analysis was performed using  
509 STRING  
510 ([https://www.string-db.org/cgi/input?sessionId=bfRWW6asD8S4&input\\_page\\_show\\_](https://www.string-db.org/cgi/input?sessionId=bfRWW6asD8S4&input_page_show_search=on)  
511 [search=on](https://www.string-db.org/cgi/input?sessionId=bfRWW6asD8S4&input_page_show_search=on)).

### 512 **Untargeted metabolomics processing**

513 The 200  $\mu$ L homogenized fecal sample was mixed with 800  $\mu$ L cold  
514 methanol/acetonitrile (1:1, v/v) to remove the protein. The mixture was centrifuged  
515 for 15 min (14000g, 4°C) followed by the drying of the supernatant in a vacuum  
516 centrifuge. For LC-MS analysis, the samples were re-dissolved in 100  $\mu$ L

517 acetonitrile/water (1:1, v/v) solvent. LC-MS/MS analysis was performed using an  
518 UHPLC (1290 Infinity LC, Agilent Technologies) coupled to a quadrupole  
519 time-of-flight (AB Sciex TripleTOF 6600) in Shanghai Applied Protein Technology  
520 Co., Ltd. The samples were separated by Agilent 1290 Infinity LC ultra-high  
521 performance liquid chromatography (UHPLC) HILIC column. In both ESI positive  
522 and negative modes, the mobile phase contained A=25 mM ammonium acetate and 25  
523 mM ammonium hydroxide in water and B=acetonitrile. The gradient was 85% B for 1  
524 min and was linearly reduced to 65% in 11 min, and then was reduced to 40% in 0.1  
525 min and kept for 4 min, and then increased to 85% in 0.1 min, with a 5 min  
526 re-equilibration period employed. AB Triple TOF 6600 mass spectrometer was used  
527 to collect the primary and secondary spectra of per sample. After separation by  
528 UHPLC, the samples were analyzed by Triple TOF 6600 mass spectrometer (AB  
529 SCIEX). Positive and negative modes of electrospray ionization (ESI) were  
530 respectively detected.

### 531 **Targeted medium-and long-chain fatty acid metabolomics processing**

532 A total amount of 100  $\mu$ L plasma per sample was taken in 2 mL glass centrifuge tubes  
533 and 1mL chloroform methanol solution was added. After 30min of ultrasound, the  
534 supernatant was taken and 2mL of 1% sulfuric acid-methanol solution was added. The  
535 mixed solution was placed in a water bath at 80°C, for 30min, and then 1mL of  
536 N-hexane and 5 mL of pure water were added in turn. Next, 500  $\mu$ L of supernatant  
537 was absorbed, 25  $\mu$ L of methyl N-nonaconate was added and mixed. The final sample  
538 was detected by GC-MS. All samples were separated by Agilent DB-WAX capillary  
539 column (30 m $\times$ 0.25 mm ID $\times$ 0.25  $\mu$ m) gas chromatography. Agilent 7890/5975C  
540 gas-mass spectrometer was used for mass spectrometry. The chromatographic peak  
541 area and retention time were extracted by MSD ChemStation software. The content of  
542 medium-and long-chain fatty acids was calculated by drawing a standard curve.

### 543 **Metabolomics statistical processing**

544 The screening of significant changed metabolites was performed using univariate and  
545 multidimensional analysis. Student's t test was applied to determine the significance

546 of differences between two groups. The variable importance in the projection (VIP)  
547 value of each variable was obtained from the Orthogonal Partial Least Squares  
548 Discriminant Analysis (OPLS-DA) model, which was used to indicate its contribution  
549 to the classification. The screening criteria were  $p < 0.05$  and  $VIP > 1$ .

#### 550 **Animal treatments**

551 Specific-pathogen-free C57BL/6J male mice at 6 weeks of age (vendor: Chengdu  
552 dossey experimental animals Co., LTD) were randomly assigned to four groups (FTPA,  
553 FT, PA and Control). All the mice lived in cages with the same conditions, including  
554 12h light and 12h dark cycles, temperature 22-25 °C and humidity 40-60%.

#### 555 **Diets**

556 HPAD was prepared by adding 12% PA to conventional forage. Both the conventional  
557 forage and the HPAD were sterile and the fresh forage was renewed three times a  
558 week. On day 0, the diets of FTPA-treated mice and PA-treated mice were switched to  
559 HPAD, FT-treated mice and control mice were still fed with conventional forage.  
560 Drinking water was sterile and renewed twice a day.

#### 561 **Transplant preparation and use**

562 After single cage feeding, FPG detection and fecal collection were performed, and  
563 fecal samples of seven T2DM macaques were mixed for the preparation of transplants.  
564 The appropriate volume of diluent was added to the fecal sample (i.e. add 4 ml diluent  
565 per gram of feces) and the preparation of diluent can be found in (86). Sodium  
566 L-ascorbic acid and L-cysteine hydrochloride monohydrate were added to all  
567 suspensions at final concentrations of 5% (w/v) and 0.1% (w/v), respectively (The  
568 sterile diluent of control group was added with the same amount of reagent). The  
569 mixture was homogenized and filtered with a 200-mesh sterile mesh screen to remove  
570 large particles from the feces, and the filtrate was passed through 400 and 800 sterile  
571 mesh screens to remove undigested food and smaller particulate matter. The filtrate  
572 was divided into 50 ml centrifuge tubes, centrifuged at 600×g for 5 min, and the  
573 precipitation was discarded. Finally, the fecal supernatant was divided into new  
574 centrifuge tubes in equal parts (400 μL per tube) and frozen at -80°C. For use, the

575 transplant was quickly thawed in a 37°C water bath.

#### 576 **Fecal microbiota transplantation (FMT)**

577 After one week of changed feeding regime, the FTPA and FT mice groups were  
578 pre-treated with 1 g/L neomycin sulfate, 1 g/L ampicillin and 1 g/L metronidazole in  
579 the drinking water for 14 days, and the control group was not treated. For FMT  
580 treatment, the gavage with 400 µL transplant, which were thawed ahead of time, was  
581 performed for 14 days in FTPA and FT-treated mice. At the same time, the gavage in  
582 control and PA-treated mice were performed with sterile diluent.

#### 583 **Metabolic measurements**

584 Throughout the experiment, body weight and feces were collected every month, FPG  
585 detection was every half month. OGTT was performed on day 110 and ITT was  
586 performed on day 115. For OGTT, after a 12h overnight fast, oral glucose gavage (1.2  
587 g/kg of 12% dextrose solution) was performed and followed by blood sample  
588 collection from the tail vein at 0, 15, 30, 60, 90 and 120 min. For ITT, after a 6h fast,  
589 intraperitoneal insulin injection was performed (0.75 U/kg, human regular insulin),  
590 followed by the blood samples collection from the tail vein at 0, 20, 40, 60, 90 and  
591 120 min. On day 120, after FPG detection, blood collection and tissues were collected  
592 for quantitative RT-PCR, H-E staining and ELISA.

#### 593 **Isolation of tissue for quantitative RT-PCR and ELISA**

594 Tissue was homogenized for RNA extraction, after adjusting the final concentration of  
595 RNA, and DNA reverse transcription was performed. Quantitative RT-PCR was  
596 performed on CFX Connect Real-Time PCR Detection System (Bio-rad, USA) using  
597 SYBR Green (Table 3-5). PA, IL-1 $\beta$ , IL-6, TNF- $\alpha$  and IL-17A were detected using  
598 Jiangsu Meimian ELISA kit and followed operator instructions.

#### 599 **H-E staining**

600 The isolated livers and pancreas were dehydrated and embedded after fixation with  
601 formalin for 24 h. Paraffin-embedded livers and pancreas specimens were cut at a  
602 thickness of 3 µm. All sections were stained with hematoxylin then eosin, and finally  
603 microscopy and image acquisition were performed.

## 604 **Statistical analysis**

605 In this study, one-way ANOVA was used to determine statistical significance for  
606 comparisons of more than three groups, and for comparisons of two groups,  
607 two-tailed t-test was used. p values are represented on figures as follows: ns, not  
608 significant, \* $p < 0.05$ , \*\* $p < 0.01$ .

609

## 610 **Data availability**

611 The raw data of transcriptomes, metagenomes, and 16S rRNA have been submitted to  
612 the China National Center for Bioinformation/Beijing Institute of Genomics, Chinese  
613 Academy of Sciences with BioProject accession no. PRJCA021499. We provide four  
614 links that reviewers can use to access the raw data. Transcriptomes data:  
615 <https://ngdc.cncb.ac.cn/gsa/s/4xAVJ3Lk> (GSA: CRA013604). 16S rRNA data  
616 (macaque): <https://ngdc.cncb.ac.cn/gsa/s/bsDz9qrN> (GSA: CRA013637). 16S rRNA  
617 data (mouse): <https://ngdc.cncb.ac.cn/gsa/s/1YcuC464> (GSA: CRA013638).  
618 metagenomes data: <https://ngdc.cncb.ac.cn/gsa/s/3k38T849> (GSA: CRA013607). The  
619 identified untargeted metabolites were listed in the Table S4. The identified targeted  
620 medium-and long-chain fatty acid metabolites were listed in the Table S5.

621

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627 manuscript.

## 628 **Author contributions**

629 X.L., G.H., Q.H.L., S.Z.Y. and C.J. collected the samples; X.L. and S.Z.Y. performed  
630 the bioinformatics analyses; X.L., Y.C.X., K.S., C.J., J.X.L. and L.Z. performed the  
631 experiments; X.L. wrote the manuscript; J.L., Z.L.H., Z.X.F. and B.S.Y. revised the

632 manuscript; J.L. conceived and designed the experiments.

### 633 **Declaration of interests**

634 The authors declare no competing interests.

### 635 **Figure legends**

#### 636 **Figure 1. The changes in gut microbiota in spontaneous T2DM macaques.**

637 (A) Alpha diversity estimates (Shannon index) between T2DM and control groups (ns,  
638 not significant, two-tailed t-test, n=8).

639 (B) Alpha diversity estimates (Simpson index) between T2DM and control groups (ns,  
640 not significant, two-tailed t-test, n=8).

641 (C) Principal Coordinate Analysis (PCoA) (n=8).

642 (D) Differential analysis of gut microbial composition in T2DM and control groups  
643 (n=8).

644 (E) LEfSe analysis between T2DM and control groups (n=8).

645 (F) Differential analysis of gut microbial function in T2DM and control groups (n=5).

646 The pathways with red color were associated with T2DM and inflammation. Error bar  
647 is mean with  $\pm$  standard deviation (s.d.).

648 (G) Differential analysis of gut microbial CAZy enzyme in T2DM and control groups  
649 (n=5). CBMs: carbohydrate-binding module ( $p=0.022$ , two-tailed t-test); GTs:  
650 Glycosyl Transferases ( $p=0.013$ , two-tailed t-test); PLs: Polysaccharide Lyases  
651 ( $p=0.017$ , two-tailed t-test); AA: Auxiliary activity enzymes (ns, not significant,  
652 two-tailed t-test); GH: Glycoside hydrolases (ns, not significant, two-tailed t-test); CE:  
653 Carbohydrate esterases ( $p=0.039$ , two-tailed t-test). For all boxplots: centre lines, upper  
654 and lower bounds show median values, 25th and 75th quantiles; upper and lower  
655 whiskers show the largest and smallest non-outlier values. In c, ellipses represent the  
656 95% confidence intervals.

657 **Figure 2. The alterations of fecal metabolites and gene expression in spontaneous**  
658 **T2DM macaques.**

659 (A) Orthogonal partial least squares discriminant analysis (OPLS-DA) score plots  
660 based on the metabolic profiles.

661 (B) Volcano plots of metabolomics ( $p < 0.05$ , two-tailed t-test).

662 (C) Fecal metabolites with significant differences between T2DM and control groups  
663 ( $VIP > 1$ ,  $p < 0.05$ , two-tailed t-test).

664 (D) Enrichment analysis of the differentially abundant pathways between T2DM and  
665 control groups ( $p < 0.05$ , two-tailed t-test).

666 (E) Non-metric multidimensional scaling (NMDS) analysis between T2DM and  
667 control groups ( $p = 0.019$ , two-tailed t-test).

668 (F) Volcano plots of DEGs ( $\log \text{fold change} \geq 1$ ,  $p < 0.05$ , two-tailed t-test).

669 (G) The GO and KEGG pathway enrichment analyses ( $p < 0.05$ , two-tailed t-test).

670 (H) Weighted Gene Co-Expression Network Analysis (WGCNA).

671 (I) Venn analysis between hub genes and DEGs. In A and E, ellipses represent the 95%  
672 confidence intervals. Data shown are from 8 individuals per group.

673

674 **Figure 3. LCFAs accumulation and inflammation occurred in spontaneous T2DM**  
675 **macaques.**

676 (A-E) The contents of SFA (A,  $p = 0.038$ ), MUFA (B), PUFA (C), N3 (D), and N6 (E) in  
677 plasma (ns, not significant, two-tailed t-test).

678 (F) The univariate analysis by two-tailed t-test, error bar is mean with  $\pm$  s.d.

679 (G) The multidimensional analysis by VIP value ( $VIP > 1$ ).

680 (H-J) The contents of serum inflammatory cytokines, including IL-1 $\beta$  (H,  $p = 0.032$ ),

681 TNF- $\alpha$  (I) and IL-6 (J) (ns, not significant, two-tailed t-test).

682 (K) Correlation analysis between DEGs, differential metabolites, and differential  
683 microbes using Spearman rank correlation ( $|r|>0.5$ ,  $\text{adj } p<0.05$ ). For all boxplots: centre  
684 lines, upper and lower bounds show median values, 25th and 75th quantiles; upper and  
685 lower whiskers show the largest and smallest non-outlier values. Data shown are from 7  
686 individuals per group.

687 **Figure 4. The FMT and high PA diet mice developed pre-T2DM characteristics.**

688 (A) Experimental scheme of FMT and high PA diet treatment.

689 (B-H) Metabolic analysis, including the trend of FPG within 120 days (B), FPG (C,  
690  $p=0.0003$ ), OGTT (D), AUC of OGTT (E,  $p=0.028$ ), FPI (F,  $p=0.007$ ), ITT (G), and  
691 body weight change (H) on day 120.

692 (I-J) The contents of TC (I,  $p=0.005$ ) and TG (J,  $p=0.041$ ) in serum on day 120.

693 (K and L) Representative H-E staining images of liver (K) and pancreas (L). For all  
694 boxplots: centre lines, upper and lower bounds show median values, 25th and 75th  
695 quantiles; upper and lower whiskers show the largest and smallest non-outlier values.  
696 Significance was determined using one-way ANOVA. In d, g, and h:  $*p<0.05$ ,  
697  $**p<0.01$ . Data shown are from 4-6 individuals per group.

698

699

700 **Figure 5. The PA accumulation required the specific gut microbiota.**

701 (A-C) Total PA contents in serum (A,  $p=0.013$ ), ileum (B,  $p=0.016$ ) and feces (C,  
702  $p=0.014$ ) on day 120.

703 (D) Quantitative RT-PCR for Cd36 transcripts in ileum on day 120 ( $p=0.049$ ).

704 (E) The content of IL-17A in ileum on day 120 ( $p=0.027$ ). For all boxplots: centre lines,  
705 upper and lower bounds show median values, 25th and 75th quantiles; upper and lower  
706 whiskers show the largest and smallest non-outlier values. Significance was determined  
707 using one-way ANOVA. Data shown are from 3-4 individual macaques per group.

708 (F) NMDS analysis ( $p=0.001$ , one-way ANOVA), ellipses represent the 95%  
709 confidence intervals.

710 (G and H) LEfSe analysis between FTPA and control groups (G), FT and control  
711 groups (H). Data shown are from 4 individuals per group.

712 (I) Specific gut microbiota structure promoted the absorption of excess PA by  
713 regulating the expression of IL-17A and *Cd36*, leading to the LCFAs accumulation  
714 and insulin resistance.

715 **Figure 6. Integration of multi-omics results.**

716 (A) Insulin resistance, fatty acid oxidation disorders, LCFAs accumulation and  
717 inflammation occurred in spontaneous T2DM macaques.

718 (B) Incomplete mitochondrial LCFAs  $\beta$  oxidation. The expression levels of fatty acid  
719 metabolism-related genes HADHB and ACSM3 were downregulated in spontaneous  
720 T2DM macaques, which could lead to accumulation of acylcarnitine, including  
721 l-propionylcarnitine, hexanoyl-l-carnitine, (r)-butyrylcarnitine, and  
722 isovaleryl-l-carnitine.

723 (C) Gut inflammation. The decrease of *Lactobacillus* sp. likely caused the reduction of  
724 serotonin and indole-3-acetaldehyde, which promotes the expansion of PA producer  
725 *Erysipelotrichacea* and ultimately led to PA accumulation. Both *Erysipelotrichacea* and  
726 *Ruminococcus gnavus* promote the development of inflammation. Accumulation of PA  
727 and inflammation are important factors in the development of T2DM.

728 (D) Accumulation of PA promoted the development of insulin resistance. In the  
729 PA-mTORC1-Akt pathway, the changes of RAP1A, SESTRIN3, and IRS1 expression  
730 promoted the development of insulin resistance in spontaneous T2DM macaques. The  
731 increase of PA promoted the development of T2DM by up-regulating the NF- $\kappa$ B  
732 signaling pathway.

733 **Tables**

734 **Table 1 Physiological and biochemical parameters of Control and T2DM group**

Index	T2DM (n=8)	Control (n=8)
FPG (mmol/L)	7.75±1	4.05±1.02 <sup>**</sup>
FPI (μU/mL)	17.97±8.50	6.45±2.56 <sup>**</sup>
HOMA-IR	6.24±3.06	1.20±0.65 <sup>**</sup>
BMI	14.03±7.71	14.30±1.42
HbA1c (%)	3.78±0.70	3.26±0.62
TG (mmol/L)	0.97±0.53	0.68±0.42
TC (mmol/L)	3.30±0.98	3.52±0.86
HDL (mmol/L)	1.23±0.40	1.42±0.38
LDL (mmol/L)	1.36±0.44	1.47±0.57

735 \*  $p < 0.05$ , \*\*  $p < 0.01$

736 FPG: fasting plasma glucose; FPI: fasting plasma insulin; HOMA-IR:  
 737 homeostasis model assessment- insulin resistance; BMI: body mass index; HbA1c:  
 738 glycosylated hemoglobin A1c; TG: triglycerides; TC: total cholesterol; HDL:  
 739 high-density lipoprotein cholesterol; LDL: low-density lipoprotein cholesterol.

740

741 **Table 2 blood routine examination of Control and T2DM group**

Index	T2DM (n=7)	Control (n=7)
WBC (10e9/L)	15.63±4.66	11.32±2.19*
RBC (10e12/L)	5.47±0.51	5.77±0.43
HGB (g/L)	129.57±15.48	136.71±9.60
HCT (%)	41.31±3.64	44.22±3.12
MCV (fl)	74.13±2.49	76.74±2.00
MCH (pg)	23.17±1.00	23.71±0.60
MCHC (g/L)	312.86±10.16	309±8.14

RDW (%)	15.23±2.12	14.69±1.74
PLT (10e9/L)	402±86.66	371.57±86.42
MPV (fl)	10.54±1.62	10.24±1.12
PCT (%)	0.42±0.08	0.38±0.09
PDW (%)	14.94±0.58	14.83±1.64
LYM% (%)	23.67±10.26	47.71±8.13**
LYM# (10e9/L)	3.46±1.66	5.39±1.52*
MON% (%)	4.18±3.34	6.18±2.29
MON# (10e9/L)	0.65±0.62	0.71±0.38
NEU% (%)	71.13±13.23	44.28±8.96**
NEU# (10e9/L)	11.36±4.91	5.00±1.38**
EOS% (%)	0.91±0.69	1.50±1.94
EOS# (10e9/L)	0.14±0.12	0.17±0.22
BAS% (%)	0.12±0.13	0.33±0.47
BAS# (10e9/L)	0.02±0.02	0.04±0.07
NRBC# (10e9/L)	0	0
NRBC% (%)	0	0

742 \*  $p < 0.05$ , \*\*  $p < 0.01$

743 WBC: white blood cell; RBC: red blood cell; HGB: hemoglobin; HCT: hematocrit;  
 744 MCV: mean corpuscular volume; MCH: mean corpuscular hemoglobin; MCHC:  
 745 mean corpuscular hemoglobin concentration; RDW: red blood cell distribution width;  
 746 PLT: platelet; MPV: mean platelet volume; PCT: procalcitonin; PDW: platelet volume  
 747 distribution width; LYM%: lymphocyte percentage; LYM#: lymphocyte value;  
 748 MON%: monocytes percentage; MON#: monocytes value; NEU%: neutrophil  
 749 percentage; NEU#: neutrophil value; EOS%: eosinophil percentage; EOS#:  
 750 eosinophil value; BAS%: basophil percentage; BAS#: basophil value; NRBC%:  
 751 nucleated red blood cell percentage; NRBC#: nucleated red blood cell value.

752

753

### Table 3 Primers of RT-PCR

Gene name	Sequences of primers
<i>CD36/F</i>	5'-ATGGGCTGTGATCGGAACTG-3'
<i>CD36/R</i>	5'-GTCTTCCCAATAAGCATGTCTCC-3'
<i>GADPH/F</i>	5'-CCTCGTCCCGTAGACAAAATG-3'
<i>GADPH/R</i>	5'-TCTCCACTTTGCCACTGCAA-3'

754

**Table 4 RT-PCR reaction components**

Reagent	Volume
2×SG Fast qPCR Master Mix	10.0 μL
DNF Buffer	2.0 μL
F primer (10umol/L)	0.4 μL
R primer (10umol/L)	0.4 μL
cDNA	1.0 μL
ddH <sub>2</sub> O	6.2 μL

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**Table 5 RT-PCR cycle proctol**

Step	Temperature	Time
1	95°C	3 min
2	95°C	1-3 s
3	60°C	30 s
4		
5	72°C	1 min

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757

## 758 **Supplementary Figures**

### 759 **Figure S1 Metagenome analysis of microbiota.**

760 (A) Differential microbes screened by metagenome analysis (p<0.05, two-tailed t-test).

761 T2DM: type 2 diabetes mellitus

762 (B) The proportion of all metabolites.

763 (C) Correlation analysis between differential untargeted metabolites and differential

764 microbes (Spearman's Rho).

765 (D) Correlation analysis between differential targeted metabolites and differential  
766 microbes (Spearman's Rho), \* $p < 0.05$ . Data shown are from 5 individuals per group.

767 **Figure S2 Pathway enrichment analysis by AFC and WGCNA.**

768 (A) Total 26 differential pathways were enriched (FDR<0.05).

769 (B) The changes of gene expression in insulin resistance pathway compared to the  
770 control group, red color illustrating the up-regulation of genes and blue showed the  
771 down-regulation of genes in T2DM group, and the darker the color of the genes, the  
772 greater the  $|\log_{10}FC|$  value.

773 (C)WGCNA functional enrichment. Data shown are from 8 individuals per group.

774 **Figure S3 The changes in gut microbiota in FTPA, FT, and PA-treated mice.**

775 (A) Alpha diversity estimates (Shannon index) ( $p > 0.05$ , one-way ANOVA,  $n=4$ ).

776 (B) Alpha diversity estimates (Simpson index) between T2DM and control groups  
777 ( $p > 0.05$ , one-way ANOVA,  $n=4$ ).

778 (C) Family level taxonomy and relative abundance of five groups.

779 (D) The changes of members of Lachnospiraceae family from day -14 to day 120.

780 (E) LEfSe analysis between PA and control groups. Data shown are from 4  
781 individuals per group.

782 **Supplementary Tables: Table. S1.xlsx- Table. S5.xlsx**

783 Table S1. Information on the macaque samples.

784 Table S2. List of 64 differential metabolites.

785 Table S3. List of genes in four modules significantly correlated with T2DM.

786 Table S4. List of all identified untargeted metabolites.

787 Table S5. List of all targeted medium-and long-chain fatty acid metabolites.

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