The vagino-cervical microbiome as a woman's life history

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The gut microbiome has been the center of attention for human commensal microbiome studies. The vaginal microbiome is also densely populated with bacteria, viruses and fungi, and the presence of microorganisms beyond the cervix is increasingly reported in non-infectious conditions^{1–3}. Due to the over 90% of human sequences in female reproductive tract samples 3,4 , metagenomic information has been very limited. 16S rRNA gene amplicon sequencing studies have identified community types in the vaginal microbiota, and observed its dynamic changes due to menstrual cycles and sexual behaviors in small cohorts^{5,6}. Here we perform metagenomic shotgun sequencing on cervical samples from 516 women of reproductive age (more than 10-fold of the Human Microbiome Project (HMP)⁴), and dissect major factors, especially pregnancy and delivery histories and contraception methods on the microbiome composition. Features of other body sites, such as mood fluctuations and facial speckles could potentially be deduced from the vagino-cervical microbiome. Our results offer an unprecedented glimpse into the microbiota in the female reproductive tract and imply disease susceptibilities that may be relieved by behavioural changes.

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Dominant bacterial and non-bacterial members of the vagino-cervical metagenome

To explore the vagino-cervical microbiome, 516 healthy Chinese women aged 21-52
(median 29, 95% CI: 23–39) were recruited during a physical examination (Sup. Data 1).
Cervical samples were collected using a cytobrush and immediately preserved on
Flinders Technology Associates (FTA) cards. Metagenomic shotgun sequencing was

8 performed on the DNA samples, and high-quality non-human reads were used for

9 taxonomic profiling of the vagino-cervical microbiome (**Fig. 1**).

10

In agreement with 16S rRNA gene amplicon sequencing data from the US⁵, the vagino-11 12 cervical microbiota of this Asian cohort was mostly Lactobacilli-dominated, while 19.19% of women harboured over 50% Gardnerella vaginalis (Fig. 1a, Extended Data Fig. 1). 13 14 Some of the volunteers were dominated by *Atopobium vaginae* but not *G. vaginalis*, which were commonly believed to co-occur in bacterial vaginosis (BV)⁷. 22.67% of the 15 cohort was dominated by Lactobacillus iners (Fig. 1a, Extended Data Fig. 1), which 16 17 was far less protective against bacterial and viral infections compared to Lactobacillus 18 crispatus. Rare subtypes such as Bifidobacterium breve, Lactobacillus johnsonii, 19 Streptococcus anginosis, Lactobacillus amylovorus and Escherichia coli were also 20 detected in this cohort (Fig. 1a). Other microorganisms including *Clamydia trachomatis*, 21 Ureaplasma parvum, human papillomavirus (HPV), herpesviruses, Haemophilus 22 influenzae and Enterobacteria phage were abundant in some individuals (Fig. 1). 23 According to the metagenomic data, the vagino-cervical microbiome varied widely 24 among individuals (Fig. 1a, Extended Data Fig. 1a), and the mean proportion of non-25 bacterial sequences was 3.45%. The vaginal types of G. vaginalis, L. crispatus, and B. 26 breve showed lower proportions of human sequences than other types (Extended Data Fig. 1b). 27

28

29 Factors shaping the vagino-cervical microbiome and beyond

We computed the prediction value (5-fold cross-validated random forest model) of female life history questionnaire on the microbiome data, and found the most important factors to be pregnancy, marriage, mode of delivery, age at vaginal sexual debut, actively breast-feeding, contraceptive methods, menstrual cycle, and sexual intercourse frequency (**Fig. 2a**).

35

Weaker but nonetheless significant dependences were observed between the vaginocervical microbiome and practices such as sexual intercourse within 24 hours, HPV vaccination, vaginal douching, oral probiotics, antibiotics and herbal medicine (**Fig. 2a**, **Sup. Data 2**).

40

As a whole, these questionnaire data collected for the female reproductive tract 41 samples showed the greatest explained variances for the vagino-cervical microbial 42 43 composition, followed by other data collected on the same day, such as faecal 44 microbiome composition, psychological questionnaire, plasma metabolites, immune indices, facial skin imaging and medical test data (Fig 2b, Extended Data Fig. 2). For 45 instance, self-assessed depression was the strongest factor in the psychological 46 47 guestionnaire to predict vagino-cervical microbiome composition, spots on the cheeks constituted the strongest factor in facial skin measurements, and plasma phosphoserine 48 49 was the most important among metabolites (Extended Data Fig. 2).

50

51 Specific influences from pregnancy histories and contraception

52 Marriage was one of the most significant factors to influence the vagino-cervical 53 microbiome (Fig. 2a). It showed negative correlations with relative abundances of L. crispatus, Acinetobacter species, L. jensenii, L. vaginalis, Ureaplasma parvum and 54 55 Comamonas testosteroni, and positive correlation with Bifidobacterium breve (Fig 3a, Sup. Data 2). Women who got married had higher concentration of vitamin D, and their 56 plasma testosterone (responsible for sexual drive), dehydroepiandrosterone and 57 58 creatinine had declined (Fig 3a). The age at vaginal sexual debut showed positive 59 correlations with Gardnerella vaginalis and Scardovia, and negative associations with L. 60 crispatus and L. iners (Fig 3a, Sup. Data 2).

61

62 Similarly, the women who went through pregnancy showed decreased *Lactobacillus*,

63 including L. crispatus, L. vaginalis, and L. jensenii, but were enriched for Actinobacteria,

64 Bifidobacteriaceae, Anelloviridae and Torque teno virus (the most abundant anellovirus

in the human virome) in the vagino-cervical microbiome (**Fig. 3c, Sup. Data 2**,

66 **Extended Data Fig.3**). We observed increased concentrations of vitamin D, blood

67 glucose and direct bilirubin in women with previous pregnancy, but lower concentrations

of testosterone, androstenedione, dehydroepiandrosterone and methionine (**Fig. 3c**).

69 Vaginal deliveries were associated with decreased *L. crispatus*, *L. jensenii*, *Bartonella*

species, *Prevotella timonensis*, and increased *Lactobacillus* sp. 7_1_47FAA (**Fig. 3d**,

71 **Sup. Data 2**). We observed a lower relative abundance of *Ureaplasma parvum* in

individuals who delivered by cesarean section (Fig. 3e, Sup. Data 2), a bacterium

commonly isolated from pregnant women⁸ and recently reported in the lower respiratory

⁷⁴ tract of preterm infants⁹. Compared with vaginal deliveries, the concentrations of

testosterone, androstenedione, methionine, threonine and tryptophan were significantly

- ⁷⁶ lower in women who experienced caesarean section (**Fig 4d,3e**). In addition, these
- 77 women more often suffered from an abnormal leucorrhoea (**Fig 3e**).
- 78

79	Among our cohort, 63 volunteers happened to be actively breast-feeding. L. crispatus,
80	and L. iners was found to be less abundant in these actively breast-feeding individuals
81	(Fig. 3f). Alkaline phosphatase, vitamin D and lead (Pb) were found to be more
82	abundant in the breast-feeding individuals, while progesterone and creatine levels were
83	slightly reduced (Fig 3f). Gravida and para, especially deliveries by caesarean section,
84	appeared associated with facial skin problems such as UV spots and porphyrins, but
85	these problems may be reduced by breast-feeding (Fig 3c-3f).
86	
87	Integrated association network using a wisdom of crowds approach ¹⁰ also revealed
88	interesting patterns (Spearman's correlation, random forest and linear regression,
89	Extended Data Fig. 4). The relative abundance of G. vaginalis was positively
90	associated with a poor mental state, poor sleep and appetite (Extended Data Fig. 4),
91	possibly due to reduced GABA (γ -amino butyric acid) production from Lactobacilli ¹¹ , and
92	the results advocate for getting a partner at a younger age (Fig. 3). L. iners negatively
93	correlated with times of spontaneous abortion, days after last menses, condom usage
94	and the concentration of vitamin D and vitamin A, and positively correlated with plasma
95	concentrations of asparagine and alanine. In addition, vaccination history, such as these
96	against HPV, tetanus toxoid, and bacillus Calmette-Guerin, may be associated with
97	higher relative abundances of <i>L. iners</i> .

98

99 We found multiple significant associations between the contraceptive methods of 100 participants and their vagino-cervical microbiome. Condom usage showed negative 101 correlations with L. iners and Comamonas species, and positive associations with 102 Prevotella species (Sup. Data 2, Extended Data Fig. 4). Oral contraceptives, on the 103 other hand, were associated with increased Ureaplasma parvum, Comamonas species 104 and Mycoplasmataceae (Fig. 4g, Sup. Data 2, and Extended Data Fig. 4). 105 Comamonas has been implicated in fecundity in C. elegans and identified as a marker for infertility due to endometriosis ^{2,12}. Moreover, oral contraceptives exhibited a positive 106 107 correlation with plasma homocitrulline, and a negative correlation with good physical 108 shape (such as waist-to-hip ratio and body weight), consistent with decline in butyrate-109 producing faecal microbiota species such as Faecalibacterium prausnizi and Roseburia 110 *intestinalis* (**Fig. 3g**). Thus, none of the contraceptive methods appeared healthy for the 111 microbiome.

112

Menstrual phases are known to influence the vaginal microbiota^{2,6}. We confirmed in our 113 114 cohort that L. iners gradually decreased in proportion after menses, coinciding with the 115 increase in progesterone and the decrease of plasma threonine and arginine (Fig 3h). 116 White blood cell counts (WBC) also recovered after menses, consistent with greatest 117 susceptibility to infection at the onset of menses. L. vaginalis, L. johnsonii and Weissella 118 species showed negative correlations with self-reported regular periods (Extended 119 Data Fig. 4). Streptococcus agalactiae, Propionibacterium acnes and Dialister 120 micraerophilus were positively associated with a heavier flow (Extended Data Fig. 4).

Many women experience dysmenorrhea during menses. Dysmenorrhea enriched for Pseudomonadales, *Acinetobacter* and Moraxellaceae, while lower in plasma level of histidine (**Fig 3i**), consistent with these bacteria producing histidine decarboxylases to convert histidine into histamine¹³.

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126 We also found some significant omics factors varied among the different vaginal types 127 (Extended Data Fig. 5). The women with types of *B. breve* and *G. vaginalis* had older 128 age at vaginal sexual debut (Extended Data Fig. 5a). The types of L. crispatus and L. 129 jensenii were overrepresented in the women who were not married yet or had fewer 130 pregnancies (Extended Data Fig. 5b, c). We observed more abundant of proline and 131 creatinine in the S. anginosus-type individuals, but lower concentrations of 11-132 deoxycortisol and 17a-hydroxyprogesterone (Extended Data Fig. 5d-g). The 133 concentrations of 11-deoxycortisol, 17α -hydroxyprogesterone and testosterone were 134 higher in L. crispatus-type than L. iners and G. vaginalis types (Extended Data Fig. 5f-135 h). Vaginal types also appeared associated with facial skin features such as A. vaginae 136 and L. johnsonii types had higher rates of skin problem than other types (Extended 137 Data Fig. 5i-I). Faecal Neisseria species, Haemophilus species and Aggregatibacter 138 species showed greater abundance in *L. jensenii*-type (Extended Data Fig. 5m-o).

139

While lacking significant association with the vagino-cervical microbial composition, the immune repertoire data showed multiple associations with functional pathways in the vagino-cervical microbiome, such as purine and pyrimidine metabolism, synthesis of branched chain amino acids, histidine and arginine (**Extended Data Fig. 6**). Red blood

cell counts, vitamin A and hydroxyl vitamin D levels negatively associated with CDPdiacylglycerol biosynthesis pathways, consistent with presence of diacylglycerol kinase in Lactobacilli with anti-inflammatory functions^{14,15}. Fecal *Coprococcus comes*, a bacterium previously reported to associate with cytokine response to *Candida albicans*¹⁶, was seen here to associate with isoleucine pathways in the vagino-cervical microbiome (**Extended Data Fig. 6**).

150

151 Overall, our comprehensive data on a healthy cohort of young women could help target 152 efforts aimed at promoting a healthy reproductive tract microbiota and minimizing preterm birth, as well as preventing infections from viruses such as the human 153 immunodeficiency virus (HIV)¹⁷ and HPV. For example, whether vitamin D 154 155 supplementation should be considered in African countries to reduce vaginal L. iners, 156 and therefore HIV and BV susceptibility. While the focus in the field has always been 157 infection, our data highlight major aspects (e.g. marriage, delivery methods, breast-158 feeding, contraception) that are worth further investigations for women in the modern 159 world. As a densely populated microbiota other than the distal gut, the vagino-cervical 160 microbiota also has the potential to reflect or even influence physiology elsewhere in the 161 human body.

162

163 Methods Summary

164 Informed consent was obtained from all 516 healthy women from the Institutional 165 Review Boards (IRB) at BGI-Shenzhen before participation in the study. Cervical 166 samples and faecal samples were collected from all volunteers during physical

examination. DNA was extracted as described^{2,18}, singled-end libraries were 167 constructed and sequenced using the BGISEQ-500 platform¹⁹. Medical test data were 168 169 performed by the physical examination center. Blood was drawn from a cubital vein of 170 volunteers, then carried out a series of experiments, including plasma metabolites 171 (hormone, vitamin, amino acid and microelement) and immune indices. Facial skin characteristics were evaluation by VISIA-CR[™] imaging system. Physical fitness tests 172 173 were evaluation by a series of HK6800 testers. Three kinds of online guestionnaires 174 about female life history questionnaire, lifestyle questionnaire and psychological questionnaire were filled in by volunteers. 175

176 Accession codes

Metagenomic sequencing data for all samples have been deposited to the (CNGB)database under the accession code CNP0000287.

179

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186 Author contributions

H.J. and C.C. conceived and organized this study. J.W. initiated the overall health
project. C.C., L.H., F.L, L.S., Y.L., X.Y. and N.L. performed the sample collection and

- 189 questionnaire collection. X.T., Y.K., W.R., Y.L., D.Z., X.Q., X.C., and J.Z. provided the
- 190 omics data. Z.J., C.C., F.L, L.H., and L.T. performed the bioinformatic analyses, H.J.,
- 191 C.C., and Z.J. wrote the manuscript. All authors contributed to data and texts in this
- 192 manuscript.
- 193

194 **Competing financial interests**

- 195 The authors declare no competing financial interest.
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- 259

260 Figure Captions

261

Figure 1 Vagino-cervical microbiome of the cohort. The microbial composition in each sample at the species level according to MetaPhylAn2 is shown. The samples are ordered according to hierachical clustering. Taxa names in black and blue denote bacteria and viruses, respectively.

266

267 Figure 2 Global view of factors influencing the vagino-cervical microbiome. a. 268 Female life history questionnaire entries on the vagino-cervical microbiome, ordered 269 according to their 5-fold cross-validated random forest (RFCV) importance on the 270 microbiome composition. b, Predicting the vagino-cervical microbiome from each omics 271 and vice versa using stepwise redundancy analysis. Numbers on the straight arrows 272 indicate adjusted R-squared from omics data to vagino-cervical microbiome; numbers 273 on the curved arrows indicate adjusted R-squared from vagino-cervical microbiome to 274 omics data.

275

276 Figure 3 Specific influences from reproductive factors on omics data. Effect size 277 of the factors shown (a, Marriage; b, Age at vaginal sexual debut; c, Pregnancy history; d. Number of vaginal deliveries; e, Number of cesarean deliveries; f, Current breast-278 279 feeding; **g**, Oral contraceptives; **h**, Days since last menstrual bleeding; **i**, 280 Dysmenorrhea). Wisdom of crowds was performed to calculate the correlation of omics 281 data. The bars are colored according to the type of omics, as in **Fig. 2b**. The length of 282 the bars represents Spearman's correlation coefficient between the actual 283 measurements and the values predicted by 5-fold cross-validated random forest models.

- A plus sign denotes Q-value <0.1, an asterisk denotes Q-value <0.05, two asterisks
- 285 denote Q-value <0.01, three asterisks denote Q-value <0.001, four asterisks denote Q-
- value <0.0001, and five asterisks denote Q-value <0.00001.

288 Online Methods

289 Study Cohort

All the 516 volunteers were recruited between May 2017 and July 2017 during physical

291 examination in BGI-Shenzhen, China. Baseline characteristics of the cohort are shown

- in **Sup. Data 1**.
- 293 The study was approved by the Institutional Review Boards (IRB) at BGI-Shenzhen,
- and all participants provided written informed consent at enrolment.

295 **Demographic Data Collection**

During physical examination, the volunteers received three kinds of online questionnaire. The female life history questionnaire contained pregnancy and delivery history, menstrual phases, sexual activity, contraceptive methods and so on. The lifestyle questionnaire contained age, disease history, eating and exercise habits and so on. The sychological questionnaire contained the evaluation of irritability, dizziness, frustration, fear, appetite, self-confidence, resilience and so on (**Sup. Data 1**).

302

303 Samples Collection

304 Cervical samples were collected and smeared in the Flinders Technology Associates 305 (FTA) cards by the doctor during gynecological examination. Fecal samples were self-306 collected by volunteers. Cervical samples and faecal samples were stored for 307 metagenomic shotgun sequencing. The overnight fasting blood samples were drawn 308 from a cubital vein of volunteers by the doctors.

309

310 DNA extraction and metagenomics shotgun sequencing

311 DNA extraction of cervical samples and fecal samples was performed as described^{2,18}, 312 respectively. Metagenomic sequencing was done on the BGISEQ-500 platform (50bp of 313 singled-end reads for cervical samples and four libraries were constructed for each lane; 314 100bp of singled-end reads for fecal samples and four libraries were constructed for 315 each lane)¹⁹.

316

317 Amino Acid Measurements

318 40 µl plasma was deproteinized with 20 µl 10% (w/v) sulfosalicylic acid (Sigma) 319 containing internal standards, then 120 µl aqueous solution was added. After 320 centrifuged, the supernatant was used for analysis. The analysis was performed by ultra 321 high pressure liquid chromatography (UHPLC) coupled to an AB Sciex Qtrap 5500 322 mass spectrometry (AB Sciex, US) with the electrospray ionization (ESI) source in 323 positive ion mode. A Waters ACQUITY UPLC HSS T3 column (1.8 µm, 2.1 × 100 mm) 324 was used for amino compound separation with a flow rate at 0.5 ml/min and column 325 temperature of 55 °C. The mobile phases were (A) water containing 0.05% and 0.1% 326 formic acid (v/v), (B) acetonitrile containing 0.05% and 0.1% formic acid (v/v). The 327 gradient elution was 2% B kept for 0.5 min, then changed linearly to 10% B during 1 328 min, continued up to 35% B in 2 min, increased to 95% B in 0.1 min and maintained for 329 1.4 min. Multiple Reaction Monitoring (MRM) was used to monitor all amino compounds. The mass parameters were as follows, Curtain gas flow 35 L/min, Collision 330 331 Gas (CAD) was medium, Ion Source Gas 1 (GS 1) flow 60 l/min, Ion Source Gas 2 (GS

332 2) flow 60 l/min, IonSpray Voltage (IS) 5500V, temperature 600 °C. All amino compound
 333 standards were purchased from sigma and Toronto research chemical (TRC).

334

335 Hormone Measurements

336 250 µl plasma was diluted with 205 µl aqueous solution. For SPE experiments, HLB 337 (Waters, USA) was activated with 1.0 ml of dichloromethane, acetonitrile, methanol, 338 respectively and was equilibrated with 1.0 ml of water. The pretreated plasma sample 339 was loaded onto the cartridge and was extracted using gravity. Clean up was 340 accomplished by washing the cartridges with 1.0 ml of 25% methanol in water. After 341 drying under vacuum, samples on the cartridges were eluted with 1.0 ml of 342 dichloromethane. The eluted extract was dried under nitrogen and the residual was 343 dissolved with 25% methanol in water and was transferred to an autosampler vial prior 344 to LC-MS/MS analysis. The analysis was performed by UHPLC coupled to an AB Sciex 345 Qtrap 5500 mass spectrometry (AB Sciex, US) with the atmospheric pressure chemical 346 ionization (APCI) source in positive ion mode. A Phenomone Kinetex C18 column (2.6 347 μ um. 2.1 × 50 mm) was used for steroid hormone separation with a flow rate at 0.8 348 ml/min and column temperature of 55 °C. The mobile phases were (A) water containing 349 1mM Ammonium acetate, (B) Methanol containing 1mM Ammonium acetate. The 350 gradient elution was 25% B kept for 0.9min, then changed linearly to 40% B during 351 0.9min, continued up to 70% B in 2 min, increased to 95% B in 0.1 min and maintained 352 for 1.6 min. Multiple Reaction Monitoring (MRM) was used to monitor all steroid 353 hormone compounds. The mass parameters were as follows, Curtain gas flow 35 l/min, 354 Collision Gas (CAD) was medium. Ion Source Gas 1 (GS 1) flow 60 l/min. Ion Source

- 355 Gas 2 (GS 2) flow 60 l/min, Nebulizer Current (NC) 5, temperature 500 °C. All steroid
- 356 hormone profiling compound standards were purchased from sigma, Toronto research
- 357 chemical (TRC), Cerilliant and DR. Ehrenstorfer.
- 358

359 Microelement Measurements

- 360 200 µl of whole blood were accurately transferred into a 15 mL polyethylene tube and
- 361 were diluted 1:25 with a diluent solution consisting of 0.1% (v/v) Triton X-100, 0.1% (v/v)
- 362 HNO₃,2mg/L AU, and internal standards (20 μg/L). The mixture was sonicated for 10min
- 363 before ICP-MS analysis. Multi-element determination was performed on an Agilent
- 364 7700x ICP-MS (Agilent Technologies, Tokyo, Japan) equipped with an octupole
- 365 reaction system (ORS) collision/reaction cell technology to minimize spectral
- 366 interferences. The continuous sample introduction system consisted of an autosampler,
- a quartz torch with a 2.5-mmdiameter injector with a Shield Torch system, a Scott
- 368 double-pass spray chamber and nickel cones (Agilent Technologies, Tokyo, Japan). A
- 369 glass concentric MicroMist nebuliser (Agilent Technologies, Tokyo, Japan) was used for
- 370 the analysis of diluted samples.
- 371

372 Water-soluble Vitamins Measurements

200 µl plasma was deproteinized with 600 µl methanol (Merck), water, acetic acid
(9:1:0.1) containing internal standards, thiamine-(4-methyl-13C-thiazol-5-yl-13C3)
hydrochloride (Sigma-Aldrich), levomefolic acid-13C, d3, riboflavin-13C,15N2, 4pyridoxic acid-d3 and pantothenic acid-13C3,15N hemi calcium salt (Toronto Research
Chemicals). 500 µl supernatant were dried by nitrogen flow. 60 µl water were added to

378	the residues, vortexed, the mixture was centrifuged and the supernatant was for
379	analysis. The analysis was performed by UPLC coupled to a Waters Xevo TQ-S Triple
380	Quad mass spectrometry (Waters, USA) with the electrospray ionization (ESI) source in
381	positive ion mode. A Waters ACQUITY UPLC HSS T3 column (1.7 μ m, 2.1 × 50 mm)
382	was used for water-soluble vitamins separation with a flow rate at 0.45 ml/min and
383	column temperature of 45 °C. The mobile phases were (A) 0.1 % formic acid in water,
384	(B) 0.1% formic acid in methanol. The following elution gradient was used: 0–1
385	min,99.0%–99.0% A; 1–1.5 min, 99.0% A–97.0% A; 1.5–2 min, 97.0% A–70.0% A,2–
386	3.5 min, 70% A-30% A; 3.5-4.0 min, 30%A-10.0%A; 4.0-4.8 min, 10%A-10.0%A; 4.9-
387	6.0 min, 99.0%A–99.0%A. Multiple Reaction Monitoring (MRM) was used to monitor all
388	water-soluble vitamins. The mass parameters were as follows, the capillary voltages of
389	3000V and source temperature of 150°C were adopted. The desolvation temperature
390	was 500°C. The collision gas flow was set at 0.10 ml/min. The cone gas and
391	desolvation gas flow were 150 l/h and 1000 l/h, respectively. All water-soluble vitamins
392	standards were purchased from Sigma-Aldrich (USA).
393	

394 Fat-soluble Vitamins Measurements

250 µl plasma was deproteinized with 1000 µl methanol and acetonitrile, (v/v,1:1) 395 396 (Fisher Chemical) containing internal standards, all-trans-Retinol-d5, 25-397 HydroxyVitamin-D2-d6, 25-HydroxyVitamin-D3-d6, vitamin K1-d7, α -Tocopherol-d6 398 (Toronto Research Chemicals). 900 µl supernatant was dried by nitrogen flow. 80 µl 399 80% acetonitrile added to the residues, vortexed, the mixture was centrifuged and the 400 supernatant was for analysis. The analysis was performed by UPLC coupled to an AB 401 Sciex Qtrap 4500 mass spectrometry (AB Sciex, USA) with the atmospheric pressure 402 chemical ionization (APCI) source in positive ion mode. A Waters ACQUITY UPLC BEH 403 C18 column (1.7 μ m, 2.1 × 50 mm) was used for fat-soluble vitamins separation with a 404 flow rate at 0.50 ml/min and column temperature of 45 °C. The mobile phases were (A) 405 0.1 % formic acid in water, (B) 0.1% formic acid in acetonitrile. The following elution 406 gradient was used: 0-0.5 min,60.0%-60.0% A; 0.5-1.5 min, 60.0% A-20.0% A; 1.5-2.5 407 min, 20.0% A–0% A,2.5–4.5 min, 0% A–0% A; 4.5–4.6 min, 0%A–60.0%A; 4.6–5.0 min, 408 60.0%A-60.0%A. Multiple Reaction Monitoring (MRM) was used to monitor all fat-409 soluble vitamins. The mass parameters were as follows, Curtain gas flow 30 l/min, 410 Collision Gas (CAD) was medium, Ion Source Gas 1 (GS 1) flow 40 l/min, Ion Source 411 Gas 2 (GS 2) flow 50 l/min, Nebulizer Current (NC) 5, temperature 400 °C. All fat-412 soluble vitamins standards were purchased from Sigma-Aldrich (USA), Toronto 413 research chemical (TRC).

414

415 *Immune indices Measurements*

10 ml whole blood was centrifuged at 3,000 r/min for 10 min, then 165 µl buffy coat was
obtained to extract DNA using MagPure Buffy Coat DNA Midi KF Kit (Magen, China).
The DNA was then sequenced on BGISEQ- 500 platform in 200 bp singled-end reads.
The data processing was using Immune IMonitor²⁰.

421 *Medical Parameters*

All the volunteers were recruited during the physical examination. The medical test including blood tests, urinalysis, routine examination of cervical secretion and so on. All the medical parameters were measured by the physical examination center and shown in **Sup. Data 1**.

426

427 Facial Skin feature

The volunteers were required to clean their face without makeup after got up in the 428 429 morning. The volunteer 's frontal face was photographed by VISIA-CRTM imaging system (Canfield Scientific, Fairfield, NJ, USA) which is equipped with chin supports 430 431 and forehead clamps that fix the face during the photographing process and maintain a 432 fixed distance between the volunteers and the camera at all times. Eight indices were obtained including spots, pores, wrinkles, texture, UV spots, porphyrins, brown spots 433 434 and red area from the cheek and forehead, respectively (Sup. Data 1). The percentile of 435 index was calculated based on the index value ranked in the age-matched database (Sup. Data 1). The higher the percentile of index, the better the facial skin appears. 436

437

438 Physical fitness test

439 8 kinds of tests were performed to evaluate volunteers' physical fitness condition (Sup. 440 Data 1). Vital capacity was measured by HK6800-FH (Hengkangjiaye, China). Eye-441 closed and single-legged standing was measured by HK6800-ZL. Choice reaction time 442 was measured by HK6800-FY. Grip strength was measured by HK6800-WL. Sit and reach was measured by HK6800-TQ. Sit-ups was measured by HK6800-YW. Step 443 444 index was measured by HK6800-TJ. Vertical jump was measured by HK6800-ZT. We 445 got a measure value from each test. Then each measure value score was assigned 446 grades A through E based on its corresponding age-matched database.

447

448 STATISTICAL ANALYSIS

449 **Quality control, Taxonomic annotation and abundance calculation**

450 The sequencing reads were quality-controlled as described previously¹⁹. The more

451 stringent condition for removal of host sequences was used for cervical samples³,

452 through alignment to the GRCh38 reference.

Taxonomic assignment of the high-quality cervical metagenomic data was performed
 using MetaPhlAn2²¹.

455 Taxonomic assignment of the high-quality faecal metagenomic data was performed

using the reference gene catalog comprising 9,879,896 genes²². Taxonomy of the

457 vagino-cervical MLGs and the faecal MGSs were then determined from their constituent

458 genes, as previously described $^{18,23-25}$.

459 Random-forest on the influence of female life history factors

- 460 The factors in female life history questionnaire were fitted against the relative 461 abundances of metaphlan2 profile (found in at least 10% of the samples) in the cervical 462 samples using default parameters in the RFCV regression function from randomForest 463 package in R. All the categorical variables were converted into continue variables, and 464 nominal variables were converted into dummy variables. Spearman's correlation 465 between measured value and 5-fold cross-validation predicted value was calculated, 466 then rank the key predictable factors. 467 The global effect size between vagino-cervical microbiome and omics data. 468 469 To evaluate the combined effect size of vagino-cervical microbiome on omics data, we 470 used forward stepwise redundancy analysis of omics data lists on the relative 471 abundances of metaphlan2 profile in forward.sel function in the packfor package in R. 472 This analysis provided a global versus global association between any two omics 473 datasets that maximize the associations by use the most predict power non-redundant 474 predictors.
- 475
- 476

477 The factors in each type of omics predicted by vagino-cervical microbiome

The factors in each type of omics were fitted against the relative abundances of metaphlan2 profile (found in at least 10% of the samples) in the cervical samples using

480 default parameters in the RFCV regression function from randomForest package in R.

481 All the categorical variables were converted into continue variables, and nominal

482 variables were converted into dummy variables. Spearman's correlation between

483 measured value and 5-fold cross-validation predicted value was calculated, then rank

484 the top 8 predictable factors in each type.

485

486 Wisdom of crowds for robust network construction between vagino-cervical 487 microbial species and other omics data

A new Multi-omics analyses method¹⁰ was used to integrated coefficient of linear 488 489 regression, variance importance from randomForest and Spearman's correlation to 490 construct omics flux networks and then visualized in CytoScape. The detail as indicated: Step 1: Data processing. All categorical variables in other omics data were converted 491 492 into continuous variables, and nominal variables were converted into dummy variables. 493 Missing values were filled with median, the samples which contained more than 70% 494 missing variables were removed. The microbial species less than10% in all the samples 495 were also removed. Removed near zero variable variables. For linear models, variables 496 were normalized. Outliers were defined as outside of the 95% quartiles and removed. 497 Step 2: Method implementation. Random forest variable importance was used to identify the most important predictor variables²⁶. RFCV regression function from 498 499 randomForest package in R with default parameter was used to get the 5-fold average 500 variable importance. We calculated the Spearman's correlation with the cor.test function 501 in base R software. For linear regression, we considered penalty regression to 502 overcome the sparse and co-linear problem, cv.glmnet function from glmnet package in 503 R was first used to figure out the best lambda parameter, bootstrapping glmnet with 504 0.632 re-sampling was performed 100 times, then we obtained the best lambda.

Step 3: Construction of robust networks. We kept first 5 average ranks for each target
variable and retained edges with Spearman's correlation Q-value <0.1. Then ggplot
package in R was used to make barplot for some representative female life history
factors (Fig 3). CytoScape was also used to visualized the omics network (Extended
Data Fig. 4).

510

511 Association between microbiome pathways and other omics

- 512 Pathway profile was calculated from the vagino-cervical metagenomic data using
- 513 humann2. Spearman's correlation was calculated between the relative abundance of
- 514 each pathway and other numerical data collected. R package heatmap were used for
- 515 visualization. Q-value <0.1 was considered as significant.



Figure 1 Vagino-cervical microbiome of the cohort.

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Figure 2 Global view of factors influencing the vagino-cervical microbiome.

Pregnancy_history -0.5 0.0 0.25 Vaginal delivery Delivery by cesarean section Breast-feeding 0.0 0.2 0.0 0.2 00 00 00 00 00 Days_after_last_menses Oral contraceptives Dysmenorrhea me Faecal microbiome Medical test Immune indices Physical fitness test Psychological questionnaire. Facial Skin feature Lifestyle que

Age at vaginal sexual debut

Marriage

Figure 3 Specific influences from reproductive factors on omics data.