Spectral binning of cervicovaginal fluid metabolites improves prediction of spontaneous preterm birth and *Lactobacillus* species dominance

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**Lay summary**

Health-promoting bacteria (lactobacilli) exist in harmony with the vaginal environment. They are the predominant vaginal bacterial species during pregnancy. However, the possibility of infection and inappropriate immune response are linked with unprompted preterm delivery (PTD). Other invasive lactobacilli can alter the chemical environment of the vagina as they seek to promote their growth. This study measured the change in concentration of biochemical compounds and predominant bacterial species in vaginal fluid that are linked to PTD. The study recruited 300 healthy pregnant women who provided vaginal fluid samples during the second trimester. The women who harboured more of *Lactobacillus jensenii* over *Lactobacillus crispatus* (both reported as health-promoting bacteria) in their vaginal fluid had less lactate and glutamate and experienced more PTD. This suggests that lactate and glutamate levels in vaginal fluid may have clinical application in identifying which *Lactobacillus* species is most active. These chemical biomarkers could provide quick and accurate prediction of PTD risk in clinical settings.

**Key Words:** preterm delivery, vaginal microbiota, microbiota community state type, lactobacilli, 1H-NMR, metabolomics

The connection between vaginal microbiota-metabolite profiles and preterm delivery (PTD, birth <37 weeks' gestation) has been a topical subject. Previous research has focused on specific metabolites using proton NMR spectroscopy (1H-NMR) (Amabebe et al. 2016, Stafford et al. 2017, Ansari et al. 2020). Spectral binning, that is, partitioning the spectrum into contiguous regions (bins), does not require prior knowledge of specific metabolites and is amenable to multivariate statistical analyses (Emwas et al. 2018).

This study retrospectively applied spectral binning to 1H-NMR spectra of cervicovaginal fluid (CVF) taken from 300 consented asymptomatic high-risk pregnant women at 19th to 22nd weeks' gestation. After delivery (term = 250 and preterm = 50), we examined its potential to identify metabolic features associated with spontaneous PTD (sPTD) and microbiota community state types (CST).

CSTs were determined for 83 of these samples by sequencing the V1–V3 region of the 16S rRNA gene (Stafford et al. 2017). All 1H-NMR spectra were binned at 0.02 ppm, with differences between term/PTD or CST groups assessed by receiver operating characteristic (ROC) curve and partial least squares discriminant analysis (PLS-DA) (www.metaboanalyst.ca).
There were no significant differences in demographics between term and preterm women in the overall (n = 300) or CST-determined (n = 83) cohort (Stafford et al. 2017). sPTD rates in the CST-determined (20.5%) and total cohort (16.7%) were not significantly different (P = 0.42). However, women with CST-V (Lactobacillus jensenii-dominated) had higher sPTD rates compared to those with CST-I (Lactobacillus crispatus-dominated) (Table 1).

From 1H-NMR spectral binning, lactate (bins 1.28 and 1.30 ppm) was higher in term than preterm women both for the overall cohort (P < 0.0001) and CST-determined women (P < 0.0001). Similarly, unassigned bins at 3.61, 3.67 and 3.69 ppm were higher in preterm than term-delivered women (all P < 0.0001). Irrespective of pregnancy outcome, lactate had the highest variable importance in projection scores in differentiating the dominant CST of women. L. crispatus-dominated CST-I showed higher lactate (P < 0.0001) and glutamate (P = 0.007) bin integrals compared to other CST groups, whereas L. gasseri-dominated CST-II and L. jensenii-dominated CST-V had a higher bin integral at 3.83 ppm compared to other CST groups (P = 0.003). Summing lactate and glutamate bins improved the ROC discrimination for CST-I vs CST-V (Table 2).

Similar to previous observations, both lactate, glutamate and unidentified bins (3.61, 3.67 and 3.69 ppm) show potential as predictors of dysbiosis/infection (Ceccarani et al. 2019) and sPTD (Stafford et al. 2017), that is, high levels of lactate and glutamate indicate L. crispatus dominance over L. jensenii. Future studies using 2D NMR methods could identify other metabolites associated with these differences (3.61, 3.67 and 3.69 ppm).

CVF lactate and glutamate levels may have clinical application in providing a quick, accurate and reliable snapshot of the functional activity of the predominant vaginal Lactobacillus spp. However, the PLS-DA data

### Table 1
Summary of analysis for CST group (I, II, III, V) showing dominant Lactobacillus species and prevalence of preterm delivery in asymptomatic women studied at 19\(^{th}\)–22\(^{th}\) weeks’ gestation. Lactobacillus\(^{+}\) species and prevalence of preterm delivery at birth were compared to term deliveries. Comparisons for prevalence of preterm delivery were made between each CST group and the total cohort (P = 0.03). There were no significant differences in demographics between term and preterm women in the overall cohort (n = 300) or CST-determined (n = 83) cohort (Stafford et al. 2017). sPTD rates in the CST-determined (20.5%) and total cohort (16.7%) were not significantly different (P = 0.42). However, women with CST-V (Lactobacillus jensenii-dominated) had higher sPTD rates compared to those with CST-I (Lactobacillus crispatus-dominated) (Table 1).

| CST | Dominant species | n | Term | Preterm | Prevalence of PTD (%) |
|-----|------------------|---|------|---------|----------------------|
| I   | L. crispatus      | 26| 24   | 2       | 7.6            |
| II  | L. gasseri        | 10| 7    | 3       | 30.0           |
| III | L. iners          | 34| 28   | 6       | 17.6           |
| V   | L. jensenii       | 11| 6    | 5       | 45.5*          |
| III/V| L. iners/L. jensenii | 2 | 1    | 1       | NA              |
| Undetermined | NA | 217 | 184  | 33      | 15.2           |
| Total | NA     | 300 | 250  | 50      | 16.7           |

*Prevalence of sPTD significantly higher in the CST-V women than in CST-I women (P = 0.02), and total study cohort including women with undetermined CSTs (P = 0.03).

CST, community state type; sPTD, spontaneous preterm delivery; n, sample population in the subset; Undetermined, no CST; NA, not applicable.

### Table 2
Predictive capacity of lactate and glutamate 1H-NMR spectrum bins for asymptomatic pregnant women with L. crispatus (CST-I) vs. L. jensenii-dominated (CST-V) vaginal microbiota.

| Metabolite, bins (ppm) | AUC | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) |
|------------------------|-----|-----------------|-----------------|---------|---------|
| Lactate, 1.28          | 0.84| 96.2            | 72.7            | 89.3    | 88.9    |
| Lactate, 1.30          | 0.82| 84.6            | 81.8            | 91.7    | 69.2    |
| Glutamate, 2.38        | 0.81| 65.4            | 100             | 100     | 55.0    |
| Lactate, 4.11          | 0.84| 76.9            | 90.9            | 95.2    | 62.5    |
| All                    | 0.94| 96.2            | 81.8            | 92.6    | 90.0    |

CST, community state type; 1H-NMR, proton nuclear magnetic resonance spectroscopy; AUC, area under the ROC curve; PPV, positive predictive value; NPV, negative predictive value; ppm, parts per million.
showed a high degree of overlap for CST-II, -III and -V groups making it difficult to distinguish these groups by metabolite profile alone - only CST-I appeared to be separated.

Although the vaginal microbiota may increasingly be dominated by lactobacilli during pregnancy, the risk of infection-inflammation-associated spontaneous PTD may still be substantial depending on the predominant species.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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Ethics approval
This study was reviewed and approved by the Yorkshire & Humber (Sheffield) Committee of the UK National Research Ethics Service (REC Number 13/YH/0167).

Author contribution statement
E A and S R designed the study. E A, S R and D A all contributed to writing the manuscript. S R and E A processed and analysed the 1H-NMR metabolite data. D A collected some of the cervicovaginal fluid samples and collated patients’ clinical data. All authors read and approved the final manuscript for submission.

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