Is the phase of the menstrual cycle relevant when getting the covid-19 vaccine?

OBJECTIVE: Stability of the menstrual cycle is a key indicator of health, and its alteration can affect the physical, emotional, sexual, and social aspects of menstruating individuals’ lives.1 A recently published study showed a statistically significant increase in cycle length after vaccination against COVID-19 and no significant changes in the menses length.2 However, there is no information about the potential association between vaccination time and change in cycle length. This study aims at assessing the association between the phase of the menstrual cycle at vaccination time and change in cycle length.

STUDY DESIGN: We analyzed data collected by the menstrual cycle tracking smartphone application Lunar App.3 This application allows users to track their menstrual cycle and menses, recording the beginning and end dates, pain intensity, blood loss quantity during menses (more, equal, or less than usual), and their COVID-19 vaccination status.

The database contained 28,876 users and 162,529 cycles. The distribution of the percentages of the users’ age ranges (years) was as follows: 18 to 24, 11.85%; 25–34, 49.15%; 35 to 44, 28.56%; 45 to 54, 8.31%; other, 2.13%. We filtered the database, keeping only users who had reported their vaccination status and at least 5 consecutive cycles. We considered the first doses or monodoses of the vaccine for the analysis and removed incomplete and/or wrong data. After this filtering process, we ended up with 371 users and 1855 cycles registered between September 2020 and February 2022. The relatively small size of the final sample is caused by the imposed restrictive inclusion and exclusion criteria to ensure the maximum attainable data quality.

For analysis, we employed the self-controlled case series method.4 Each participant in our cohort was a control and a case before and after getting the COVID-19 vaccine, respectively. Our primary outcome was menstrual cycle length change in days. The secondary outcomes were menses length change in days and variations in the usual blood quantity and pain intensity during the menses. We stratified the analysis of all outcomes by the phase of the menstrual cycle of the user at vaccination time. We considered the luteal phase, ie, the period between menstruation and the 14 days before it5 owing to the relative robustness of this phase. We considered the rest of the cycle as the follicular phase. The distribution of the medians (over each user) of cycle lengths before the vaccine had a median value of 28 days, with a (5–95) interpercentile range of (22–34) days.

For calculating the menstrual cycle length change, we computed the difference between the median length of the 3 cycles before the vaccine and the length of the cycle in which the vaccine was given (4th cycle) for each user. We then computed the median over all the users and the 95% confidence intervals of the point estimate. We used medians, because the data was not normally distributed. We proceeded similarly for the menses length but employed data from the fifth cycle. For the blood loss quantity and pain intensity, we computed the differences in the percentages of cycles with abnormalities in each endpoint before and after the vaccine and the 95% confidence intervals of the point estimates. Users reported abnormalities when they had more or less blood loss quantity or pain intensity than usual during menses. We employed the Wilcoxon signed-rank and chi-square tests for statistical hypothesis testing of medians and proportions, respectively. Statistical significance was set at P<.005. The participants of this study provided their consent for the analysis of their data for menstrual or reproductive health research purposes on registration in the app, and the study obtained the approval of an ethics committee. The app does not gather information about the usage of contraception or cycle control methods, and this is a potential limitation of our study, as it could affect the outcomes.

RESULTS: We observed an increase in the median cycle length of 0.5 (0.0–1.0) days (P value <0.005) for all

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individuals, with 8.08% of the individuals having an increase of 8 or more days, which is considered clinically significant.\(^6\) We observed no variation in menses length, which is in line with results previously reported in the literature.\(^2\) In addition, we observed no significant variations in the percentages of cycles with abnormal blood loss or pain intensity.

Furthermore, the stratified analysis showed an association between the phase of the menstrual cycle of the individual at vaccination time and cycle length change. Thus, individuals vaccinated during the follicular phase showed a median cycle length increase of 1 (0.0–1.0) day (\(P\) value <.005), with 11.82% of the users having an increase of 8 or more days. Individuals vaccinated during the luteal phase showed no change (Table).

**CONCLUSION:** Our results show an association between the phase of the menstrual cycle at vaccination time and change in cycle length. Thus, vaccination during the luteal phase would have a protective effect over Covid-19 vaccine-related menstrual cycle disorders, compared to vaccination during the follicular phase. The presented results suggest considering the phase of the menstrual cycle for the design of future COVID-19 vaccination policies and recommend vaccination during the luteal phase.

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OBJECTIVE: Grand rounds are a central component of medical education across specialties, with demonstrated benefits. Obstetrics and gynecology (OB-GYN) departments across the country transitioned from in-person grand rounds (IPGR) to virtual grand rounds (VGR) during the COVID-19 pandemic. Knowledge gaps exist around OB-GYN educators’ and trainees’ perceptions of VGR. Single-center studies of faculty perspectives in other specialties have shown positive correlations with improved attendance. This multicenter observational study sought to explore the OB-GYN experience with VGR and to assess perceptions by role (educator vs trainee).

STUDY DESIGN: After institutional review board exemption, a cross-sectional survey comparing VGR and IPGR was developed de novo using themes from the literature, and then reviewed and edited by subject matter experts. Likert-style questions asked about satisfaction, engagement, learning, and multitasking (Supplemental Material). The e-survey was deployed in May 2021 to all members of 5 academic OB-GYN departments, with a 2-week reminder email.

Data were analyzed in aggregate. Trainee (resident or fellow) and educator responses were then compared, excluding research staff and emeritus faculty respondents. We used bivariate statistics and regression to control for founders that were significant in bivariate analyses.

RESULTS: Of the 591 potential participants, 306 (52%) responded. Among respondents, 69% were faculty, 21% residents, 7% fellows, and 3% others (research staff or emeritus faculty).

Compared with IPGR, 91% felt satisfied with VGR; 90% reported being more likely to attend VGR. Presentation quality was assessed as the same or better by 91%, and 93% described presenter caliber as the same or better; 48% reported learning the same amount. However, 90% were more likely to multitask; 69% felt the sense of community was worse.

The secondary analysis included 297 respondents. Response rates were 45% (86/193) for trainees and 53% (211/397) among educators. Compared with educators, trainees were more likely to be dissatisfied, less likely to attend, and reported learning less during VGR (Table). Almost all trainees were more likely to multitask (Table). After controlling for gender and institution, compared with

| Perspectives                                      | Educators N = 211 (%) | Trainees N = 86 (%) | P valuea | aOR (95% CI)b |
|--------------------------------------------------|-----------------------|---------------------|----------|---------------|
| Overall unsatisfied with virtual grand rounds    | 11 (5.3)              | 16 (18.8)           | .001     | 0.14 (0.05–0.40) |
| Less likely to attend VGR than in-person grand rounds | 14 (5.7)              | 18 (20.9)           | <.001    | 0.17 (0.07–0.43) |
| Learned less during VGR than during in-person grand rounds | 38 (18.1)              | 26 (30.6)           | .03      | 0.42 (0.22–0.82) |
| Felt a loss of sense of community with VGR compared with in-person grand rounds | 142 (67.6)              | 64 (74.4)           | .19      | —             |
| More likely to ask questions in VGR than in in-person grand rounds | 105 (50.5)              | 38 (44.2)           | .37      | —             |
| More likely to multitask during VGR than during in-person grand rounds | 185 (88.9)              | 80 (93.0)           | .29      | —             |

aOR, adjusted odds ratio; CI, confidence interval; VGR, virtual grand rounds.

a P values obtained by nonparametric bivariate statistics; Kruskal–Wallis and Wilcoxon rank-sum tests, as appropriate; b adjusted for gender and site.

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