Reproducibility of Plasma and Urinary Sex Hormone Levels in Premenopausal Women over a One-Year Period\textsuperscript{1}

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Abstract

Although endogenous sex steroid hormones in premenopausal women may be associated with the risk of breast cancer and other illnesses, direct evidence to support this hypothesis is limited in large part by methodological issues in the conduct of relevant studies. One major unresolved issue is whether a single blood sample (such as is available in most epidemiological studies), collected in a specific phase of the menstrual cycle, reflects long-term levels in that phase. To address this issue, two sets of blood and urine samples were obtained from 87 premenopausal women over a 1-year period in both the follicular and luteal phases. Plasma estradiol, estrone, and estrone sulfate were measured in the blood samples obtained in both phases, whereas progesterone and urinary 2- and 16α-hydroxyestrone were measured in luteal-phase samples only. For all of the women combined, intraclass correlation coefficients (ICCs) ranged, with one exception, from 0.52 to 0.71 for the plasma estrogens and the estrogen metabolites. The sole exception was for estradiol in the luteal phase (ICC = 0.19); inclusion of only women who were ovulatory in both cycles and who collected each sample 4–10 days before their next period resulted in a substantially higher ICC for estradiol in the luteal phase (ICC = 0.62; 95% confidence interval, 0.43–0.78). These data indicate that, for several plasma and urinary sex hormones, a single follicular- or luteal-phase measurement in premenopausal women is reasonably representative of hormone levels in that phase for at least a 1-year period.

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\textsuperscript{2}The abbreviations used are: ICC, intraclass correlation coefficient; NHSII, Nurses’ Health Study II; CV, coefficient(s) of variation; CI, confidence interval.

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(i.e., still having menstrual periods); (b) had not used oral contraceptives or other hormonal preparations (e.g., for infertility) in the previous 6 months and had no plans to begin using them; (c) had not been pregnant or lactating in the previous 6 months and were not planning to become pregnant in the next year; and (d) had completed and returned the 1989, 1991, and 1993 NHSII questionnaires. From a sample of 746 NHSII women who were invited to participate, 438 (59%) responders were willing to collect two timed samples of blood (follicular and luteal) and one urine sample. Of these, 112 women were no longer eligible to participate (e.g., had recently become pregnant); thus, sample collection kits were mailed to the 326 eligible women. Two hundred women (61% of 326) returned the kits. A second collection kit was mailed to the 121 women who had returned the first kit, remained eligible to participate, and subsequently agreed to provide a second set of samples. Eighty-seven women (72% of 121) returned the second set of samples.

Blood and Urine Collections. Each blood collection kit contained all of the supplies needed to have blood samples drawn by a local laboratory or a colleague (e.g., needle, tourniquet, and blood collection tubes with sodium heparin). The participants were asked to have their first blood sample (15 ml) drawn on the 3rd, 4th, or 5th day of their menstrual cycle (“follicular blood draw”), and to have the second blood sample (30 ml) drawn 7 to 9 days before the anticipated start of their next cycle (“luteal blood draw”). Timing of the luteal sample from the estimated first day of the next menstrual cycle is generally more accurate than counting forward from day 1 of the current cycle, because the length of the follicular phase is more variable than the length of the luteal phase (15, 16). The participants were instructed to draw the blood samples, whenever possible, in the morning while fasting; 35% of the women collected blood while fasting. Participants placed their follicular blood samples in a refrigerator for 8–24 h after it was drawn; they then aliquoted the plasma into cryotubes. The plasma was kept in the participant’s home freezer until the second (luteal) blood collection. A spot morning urine sample also was collected on the same day as the second blood draw. All of the samples were then returned to our laboratory by overnight mail, with a frozen water bottle to keep them cool. Upon arrival in the laboratory, the whole blood from the luteal collection was centrifuged and plasma aliquoted into labeled cryotubes, and all of the samples were then stored in liquid nitrogen freezers. In a pilot study of postmenopausal women, levels of estradiol and free estradiol (in addition to other hormones) were observed to be stable when processing was delayed for 24 to 48 h (17). In a small pilot study of 16 premenopausal women, we collected two blood samples from each woman to compare follicular hormone levels in samples processed immediately or processed exactly as described above (which included simulating transport in a cooler with a frozen water bottle). We observed similar means and standard deviations for estrone and estradiol when using the two different processing methods, and the ICCs were 0.93 and 0.80 for estradiol and estrone, respectively.

A questionnaire was sent to each subject on which to record the first day of the menstrual cycle during which the blood samples were drawn, the dates of both blood draws, and the date of urine sample collection. In addition, information was collected on the time of day of blood collection, the number of hours since last food intake prior to the two blood draws, and the participant’s current weight. Finally, a postcard on which to record the first day of the next menstrual cycle was provided (95% of postcards were returned). Other data (e.g., information on smoking status) were available for each participant from the NHSII questionnaires completed in 1993.

Laboratory Methods. Plasma estradiol, estrone, and progesterone (for the luteal samples only) were assayed by Quest Laboratory (San Juan Capistrano, CA), and estrone sulfate was assayed by the laboratory of Dr. Longcope (University of Massachusetts Medical Center, Worcester, MA). Because of financial constraints, samples from only 30 women were measured for estrone sulfate levels. The laboratory of one of the authors (H. L. B.) assayed the urine samples for 2- and 16α-hydroxyestra-1,4-diene and for creatinine. The ratio of 2-hydroxyestrone to 16α-hydroxyestrone was calculated.

Estradiol (18) and estrone (19) were assayed by organic extraction, celite chromatography and RIA. Reported values are corrected for procedural losses. This method is highly specific and is the ‘gold standard’ for estimating steroid levels in plasma. Estrone sulfate was assayed by RIA (of estrone) after initial extraction of estrone, enzyme hydrolysis, organic extraction, and separation by column chromatography (20). Progesterone was assayed by RIA preceded by organic extraction (21). Urine samples were assayed for 2-hydroxyestrone and 16α-hydroxyestrone by means of a new ELISA procedure, as detailed elsewhere (22). Because urine samples were not collected over a 24-h period and total urinary output was unknown, creatinine levels were measured for each sample with a Beckman manual creatinine analyzer. All of the urinary metabolite levels are thus presented as standardized values (estrogen metabolite value divided by creatinine level in each sample) to account for differences arising from variations in urine concentration.

All of the luteal and follicular samples from a single woman were assayed together in the same laboratory batch but were labeled to prevent laboratory personnel from identifying which samples were from the same woman. Intra-assay laboratory CV were obtained by sending masked replicate samples. For estradiol, estrone, and progesterone, CV were less than 10%. The CV for each of the urinary metabolites was 15%. Estrone sulfate was assayed in two batches, and the CV percentages were 15 and 7.4%, respectively.

Statistical Analysis. The natural logarithms of the plasma hormone and metabolite values were used in analyses because the transformed values were more normally distributed. Between-person and within-person variances were estimated from the two sets of hormone measurements by random effects models, as implemented by SAS (SAS Institute, Cary, NC). To assess reproducibility, ICCs were calculated by dividing the between-person variance by the sum of the between-person and within-person variances (23). CIs (95%) were calculated for the ICCs (8). Because substantial batch-to-batch variation was observed in estrone sulfate levels, we calculated the ICCs for each batch individually and then obtained a weighted average (based on the number of women in each batch) for the two batches.

Although 87 women contributed all of the requested samples of blood and urine, several provided an insufficient volume to assay all of the hormones. As a result, ICCs in this study were based on 83 women for follicular estrone, 85 women for all of the other plasma hormones, and 86 women for urinary metabolites.
Results

The characteristics of the premenopausal women in this study are summarized in Table 1. The mean number of months between the collection of the two follicular blood samples was 11.3 months (range, 5.3–22.1 months). At the time of collection of the first blood sample, the women were 31–49 years of age, with a mean age of 41 years. The median cycle length for the women in this study was 27.5 days. The ICC for the cycle length in the first compared with the second blood collection was 0.26, increasing to 0.55 (95% CI, 0.39–0.70) when five women with 1 cycle greater than 100 days in length were removed. Baseline characteristics of participants in this study (listed in Table 1) were not substantially different from those of the women who initially met the eligibility criteria (n = 633), however, the participants were less likely to smoke (5.7% versus 20.2%).

The women in this study successfully collected their follicular samples at the beginning of their menstrual cycles (0.6% on the first day, 26.8% on the second day, 42.3% on the third day, and 30.3% on the fourth day). The mean levels of follicular and luteal estrogens, progesterone, and the two urinary metabolites (measured per gram of creatinine) were similar at the two sample collections (Table 2). As expected, luteal estrogen levels were much higher than follicular estrogen levels. The mean number of months between the collection of the two follicular blood samples was 11.3 (3.1). At the time of first blood draw, the women were 31–49 years of age, with a mean age of 41 years. The median cycle length for the women in this study was 27.5 days. The ICC for the cycle length in the first compared with the second blood collection was 0.26, increasing to 0.55 (95% CI, 0.39–0.70) when five women with 1 cycle greater than 100 days in length were removed. Baseline characteristics of participants in this study (listed in Table 1) were not substantially different from those of the women who initially met the eligibility criteria (n = 633), however, the participants were less likely to smoke (5.7% versus 20.2%).

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For all of the women combined, ICCs for plasma estrogens and urinary metabolites ranged, with one exception, from 0.52 to 0.71; the sole exception was for estradiol in the luteal phase (ICC = 0.19; Table 3). The ICC for progesterone was 0.54. Correlations did not vary substantially whether the first and second blood samples were collected over a period of ≤11 months or >11 months, or if the women were ≤40 or >40 years old (data not shown). ICCs were essentially unchanged after removing women who reported being current smokers on the 1993 NHSII questionnaire (n = 6). In a separate analysis, we removed women (n = 26) who had gained or lost weight (plus or minus more than 1 body mass index kg/m2 unit) between the two blood collections. ICCs for women with a stable weight were similar for luteal estrogens but slightly stronger for follicular
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Table 4  Cross-classification of premenopausal plasma hormone and urine hormone metabolite levels: first sample by second sample

<table>
<thead>
<tr>
<th>Plasma follicular estradiol (n = 85)</th>
<th>Quartiles for sample 1</th>
<th>Quartiles for sample 2</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>9</td>
<td>4</td>
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<tr>
<td>2</td>
<td>7</td>
<td>7</td>
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</tr>
<tr>
<td>4</td>
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<td>5</td>
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<td>19</td>
<td>24</td>
<td>24</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Urinary ratio* (n = 86)</th>
<th>Quartiles for sample 1</th>
<th>Quartiles for sample 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>16</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>3</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td>20</td>
<td>23</td>
<td>24</td>
</tr>
</tbody>
</table>

*2:Hydroxyestrone:16α-hydroxyestrone.

estrogens and progesterone, compared with ICCs for all of the women combined (estradiol ICC = 0.63, 95% CI = 0.47–0.77; estrone ICC = 0.57, 95% CI = 0.39–0.72; progesterone ICC = 0.70, 95% CI = 0.55–0.81).

Twenty-two women (26%) had at least one progesterone level of <300 ng/dl (which, in the luteal phase, generally indicates an anovulatory cycle); 11 (50%) of these 22 women reported having irregular cycles (overall, 65% of women with irregular cycles had a normal progesterone level). Eight of the 22 women had progesterone levels of <300 ng/dl on both blood draws and 14 on only one blood draw. Of these 22 women, 16 had missed their luteal phase altogether in at least one cycle (i.e., had collected blood <4 or >13 days before the beginning of their next cycle); 3 were presumed to be anovulatory (i.e., had a progesterone level of <300 ng/dl in a sample that was collected within 4–13 days of the next menses); and 3 were lacking information on the date of the blood draw and thus could not be assigned a reason for the low progesterone level. After exclusion of these women, ICCs for luteal-phase estrogens improved substantially (ICC for estradiol = 0.49; Table 3). The ICC for progesterone decreased somewhat (from 0.54 to 0.45) after restricting the analysis to women with progesterone levels ≥300 ng/dl; this change could be due to a decrease in the between-person variation contributed by those women with repeated low progesterone levels.

To further investigate the reproducibility of luteal estradiol and estrone levels, we next excluded women who did not collect both blood samples during the midluteal phase of their menstrual cycle, defined here as 4 to 10 days before their next cycle. This additional exclusion substantially improved the ICCs of the plasma estrogens, compared with the estrogen ICCs for all of the women combined (r = 0.62 for estradiol and r = 0.69 for estrone; see Table 3) but did not affect the ICCs for urinary estrogen metabolites (data not shown).

In an additional analysis, we examined the reproducibility of plasma luteal estrogens among women who collected the second plasma sample at the same time during their menstrual cycle as the previous sample (within 4 days), regardless of whether that time reflected the midluteal phase. The ICCs in this analysis were similar to those obtained by the exclusion of women with progesterone levels of <300 ng/dl and those women who did not collect blood between 4 and 10 days before their next cycle (ICC = 0.71, 95% CI = 0.54–0.83 for estrone; ICC = 0.54, 95% CI = 0.33–0.73 for estradiol).

Because most studies in epidemiology will categorize continuous exposures (i.e., plasma hormones) into quartiles to examine their association with disease, we have cross-classified two samples of plasma follicular estradiol and of urinary ratio (2-hydroxyestrone:16α-hydroxyestrone) to show concordance and discordance for the same women (Table 4). For follicular estradiol, 34 (40%) of 85 were perfectly classified, and 65 (77%) of 85 were off by one category or less. Similarly, for the urinary ratio, 43 (50%) of 86 were perfectly classified, and 71 (83%) of 86 were off by one category or less. Only three women (4%) for follicular estradiol and two women (2%) for the urinary ratio were extremely misclassified. These data show that one sample can classify women into the appropriate quartile of exposure relatively well.

Discussion

ICCs for levels of plasma estrogens and urinary estrogen metabolites ranged from 0.19 to 0.71 when measured 1 year apart (on average) in a sample of premenopausal women. The exclusion of women with low plasma progesterone levels from analyses substantially improved the correlations for both estradiol and estrone in the luteal phase. For example, the ICC for luteal phase estradiol increased from 0.19 to 0.49. We observed the highest ICC for plasma estradiol and estrone in a subset of women who collected both blood samples at approximately the same time in the luteal phase and were ovulatory in each cycle.

Reproducibility was high for total estradiol, estrone, and estrone sulfate measured in the follicular phase of the menstrual cycle. To our knowledge, no other study has measured the reproducibility of estrogens in the follicular phase in premenopausal women. Similarly, the two urinary estrogen metabolites measured in this study, 2-hydroxyestrone and 16α-hydroxyestrone, as well as the ratio of the two, had good reproducibility over a 1-year period when adjusted for creatinine levels. Pasagian-Macaulay et al. (13) reported a similar ICC for...
the ratio of these two metabolites over a 6-month period in a sample of 171 premenopausal women (ICC = 0.67), although the two samples were collected at random during the menstrual cycle.

Only one previous study has assessed the reproducibility over time of total estradiol in premenopausal women (9). In this study, estradiol was measured in two blood samples obtained from each of 60 premenopausal women. The samples were collected on the same day of the luteal phase (between the 20th and 24th day, counting forward) about 1 year apart. The authors observed a very low ICC for total estradiol (ICC = 0.06). This result is consistent with our data, inasmuch as we also observed poor reliability (ICC = 0.19) for total estradiol in the luteal phase before the exclusion of women who were anovulatory or missed their luteal phase.

Studies on menstrual cycle fluctuations have shown that the luteal phase is more consistent in length than the follicular phase (15, 16). For this reason, we requested that women collect luteal-phase samples 7–9 days before the first day of their next anticipated menses rather than between the 20th and 24th days of the cycle (counting forward). By subsequently obtaining postcards from the women to confirm the first day of their next cycle, we were able to remove from the analysis samples that were not collected in the luteal phase. By measuring luteal-phase progesterone levels, we also were able to exclude women who had one or more anovulatory cycles. Because the age of the women and the interval over which samples were collected in our study are similar to these variables in the study of Muti et al. (9), the low ICC they reported was probably due to an inability to make these important exclusions.

This study was relatively small, and only a fraction of eligible women provided two sets of samples. However, when we compared the women who participated to the entire group of eligibles, with just two exceptions, the two groups were quite similar. Participants were less likely to smoke and were more likely to report a history of benign breast disease. Adjusting for smoking status and history of benign breast disease did not alter our ICCs, and, although we had insufficient power to assess whether the ICCs varied according to these attributes (e.g., ICCs in smokers versus nonsmokers), substantial differences would seem unlikely.

Because reproducibility studies require two samples from each participant, our sample size for the restricted (luteal) analysis was limited to the 39 women who obtained both samples 4–10 days before their next cycle and were ovulatory at each cycle. However, if one sample from each woman had been sufficient, the sample size would have been substantially larger; i.e., 63% of all of the samples obtained were collected between 4 and 10 days. Because of study size, cost considerations, and concerns about participation rates, only one sample per participant can be obtained in most epidemiological studies. Therefore, it would be reasonable for researchers to collect luteal samples in a manner similar to that used in this study and to match cases and controls on the number of days the samples were collected before the next menses.

The ICCs observed in this study are comparable in magnitude to those for a number of other exposure measures commonly used in epidemiology. For example, correlations have been reported for serum cholesterol of 0.65 over 1 year (24) and 0.76 over 2 years (25) and for systolic blood pressure of 0.60 over 2 years (25). Correlations in this range result in relatively modest decreases in the estimated relative risk, although the degree of attenuation will depend on the magnitude of the relative risk (8). For example, measurement error in a variable with an ICC of 0.68 will lower a true relative risk of 2.0 and 2.5 to 1.6 and 1.9, respectively (8). Besides providing important information on the reproducibility of a variable, ICCs can be used to correct relative risk estimates for random within-person measurement error in epidemiological studies (26).

Results from this study indicate that a single sample of blood or urine collected during the follicular or midluteal phase of the menstrual cycle in premenopausal women can reasonably represent estrogen levels in blood or their metabolites in urine over at least a one-year period. Studies that assess reproducibility over a longer period are needed to further delineate long-term stability in premenopausal women.

References

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