Comparison of Anogenital Distance and Correlation with Vulvo-vaginal Atrophy: A Pilot Study on Premenopausal and Postmenopausal Women

Lavinia Domenici, Angela Musella, Carlotta Bracchi, Francesca Lecce, Michele Carlo Schiavi, Vanessa Colagiovanni, Violante Di Donato, Claudia Marchetti, Federica Tomao, Innocenza Palaia, Ludovico Muzii, Pierluigi Benedetti Panici

Department of Gynecological and Obstetric Sciences, and Urological Sciences, University “Sapienza” of Rome, Rome, Italy

Objectives: Anogenital distance (AGD) represents the space between labia posterior commissure and anus. This was pilot study to investigate how menopause and so lack of oestrogens affects AGD.

Methods: A total of 109 patients were enrolled. AGD was measured in lithotomy position using sterile paper ruler. Anogenital index (AGI) was used to control 2 variables of height and weight (body mass index, kg/m²). Vaginal health index (VHI) was used to evaluate vaginal wellness. Female sexual function index (FSFI) questionnaire was administered to all women to evaluate the impact of menopause on their sexual function.

Results: AGD (30.87 ± 2.98 vs. 17.57 ± 2.18; P = 0.0001) and AGI (1.40 ± 0.21 vs. 0.70 ± 0.15; P = 0.0001) were both significantly lower in the postmenopausal group. Postmenopausal women were affected by vulvovaginal atrophy (VVA) significantly. Thus, VHI scores were dramatically worse in postmenopausal group (23.95 ± 1.28 vs. 10.75 ± 3.41; P = 0.0001) as well as FSFI results (32.68 ± 2.25 vs. 19.78 ± 5.46; P = 0.0001).

Conclusions: This study confirms that AGD in post-menopausal women was significantly shorter than AGD in premenopausal women, correlating with an increase of VVA and sexual impairment. Changes of AGD and AGI demonstrated to predict hormonal changes that may occur after menopause. (J Menopausal Med 2018;24:108-112)

Key Words: Anal canal · Atrophy · Genitalia female · Menopause · Sexuality

Introduction

The anogenital distance (AGD) is identified as the length between labia posterior commissure and the centre of the anus measured in millimetres. Its size depends on hormonal changes, particularly in estrogen and androgen level modifications.1~3 The relationship between AGD and sex steroids is still well known, considering that AGD has been used as bioindicator of fetal androgen exposure in humans and specifically to estimate the consequences of adverse in utero hormonal exposure.4~7 Menopause hormonal variations are responsible of several anatomical and physiological changes in internal and external genitalia. Decreasing estrogen levels, the production of collagen changes with an increase in collagen strength, skin elasticity reduces and genitalia water content declines severely. Generally, genitalia trophism is dramatically affected by menopause, having in women a negative effect on intimacy and even on quality.
of life. Recent studies confirm that one of the most severe menopause-related problems is vulvovaginal atrophy (VVA), involving more than 50% of women in Europe and reaching over 78% in the Italian population.8,9

Nevertheless, there is a lack of studies on anatomical changes in adult females. This is a pilot study aimed to evaluate anatomical and morphological changes and comparison between pre-menopausal and post-menopausal women, focused on AGD and vaginal health index (VHI) modifications.

**Materials and Methods**

A total of 109 patients have been identified, recruited and enrolled in the outpatient clinic at the Department of Obstetrics and Gynecology, “Sapienza” University of Rome between April 2016 and December 2016.

Inclusion criteria for premenopausal group were: age between 20 and 45 years, regular periods in the past 1-year (intervals of 20–45 days). Inclusion criteria for premenopausal group were: age between 45 to 80 years, physiologic menopause since at least 1 year.

Exclusion criteria in both groups were: gynecological surgeries in the past affecting fertility and hormone production (e.g., oophorectomy), chemotherapy or irradiation of the pelvis, hormonal treatment in the previous year (combined oral contraceptives or hormone replacement therapy), congenital anatomical abnormalities, pregnancy, breastfeeding, severe comorbidities (e.g., autoimmune diseases, diabetes, etc.), infections, bleeding, and tumors.

All patients were extensively informed of study design and hypothesis. Written informed consent was obtained by all patients enrolled. Institutional Review Board (IRB) provided exempt to ethical approval. The present study was conducted following the principles of the Declaration of Helsinki. Candidates were consecutively randomized in the 2 groups.

Patients’ data (height, weight, body mass index [BMI], AGD) were collected in the same day during outpatient examination. AGD measurement was conducted using a paper ruler in the lithotomy position to define the distance between posterior commissure of labia and anus center (Fig. 1).

Anogenital index (AGI) was used to control 2 variables of height and weight,10 by dividing AGD by BMI (kg/m²).

VHI11 was used to evaluate vaginal wellness by analysing 5 parameters by clinical inspection: elasticity, fluid volume, pH, epithelial integrity and moisture (Table 1). Each parameter is graded from 1 (worst condition) to 5 (best condition). Scores ≤15 are considered to denote vaginal atrophy.

Female sexual function index (FSFI) questionnaire was administered to all women to evaluate the impact of genitalia changes on sexual life, FSFI is a validated, multidimensional, self-reported questionnaire to assess female sexual function,12 It consists of 19 items evaluating the FSFI lubrication, sexual arousal, orgasm, sexual satisfaction, and pain. The total score (range, 2–36; severe sexual dysfunction to full sexual function respectively) is calculated by summing the 5-domain score and interpreted by comparison with age- and population-dependent reference values for normal and impaired sexual function, A FSFI score ≤ 26.55 is considered to indicate sexual dysfunction.13

Interval data were analysed using the Mann–Whitney U test. Nominal data were evaluated using χ² test or Fisher’s exact test when appropriate. Parameters are expressed as mean ± standard deviation (SD) and 95% confidence interval (CI). All statistical tests were performed using SPSS statistical software program (SPSS 20.0; SPSS Inc., Chicago, IL, USA). All P values of less than 0.05 were considered to indicate statistical significance.
Results

To compare AGD in relation with VHI and FSFI of premenopausal and postmenopausal women, a total of 109 subjects were examined, being 48 in premenopause and 61 in menopause.

Patients’ characteristics are shown in Table 2, BMI significantly varied between 2 groups (22.34 ± 2.84 vs. 25.36 ± 3.59; P = 0.0001), probably in relation to metabolic changes happening during menopause, AGD (30.87 ± 2.98 vs. 17.57 ± 2.18; P = 0.0001) and AGI (1.40 ± 0.21 vs. 0.70 ± 0.15; P = 0.0001) were both significantly lower in the postmenopausal group. Comparison between AGD in the 2 groups is evaluable in Figure 2 as well.

As highlighted in Table 2, postmenopausal women were affected by VVA significantly and negatively compared with premenopausal patients. Thus, VHI scores were dramatically worse in postmenopausal group (23.95 ± 1.28 vs. 10.75 ± 3.41; P = 0.0001) as well as FSFI results (32.68 ± 2.25 vs. 19.78 ± 5.46; P = 0.0001). Specific data on VHI and FSFI are shown in Figure 3 and 4 respectively.

Table 1. Vaginal health index

| Score | 1 | 2 | 3 | 4 | 5 |
|-------|---|---|---|---|---|
| Elasticity | None | Poor | Fair | Good | Excellent |
| Fluid volume | None | Scar amount, vault not entirely covered | Superficial amount, vault entirely covered | Moderate amount | Normal amount |
| pH | >6.1 | 5.6-6.0 | 5.1-5.5 | 4.7-5.0 | ≤4.6 |
| Epithelial integrity | Petechiae noted before contact | Bleeds with light contact | Bleeds with scraping | Not friable, thin epithelium | Normal |
| Moisture | None, surface inflamed | None, surface not inflamed | Minimal | Moderate | Normal |

Table 2. Characteristics of patients

| Variables | Premenopausal women (n = 48) | Postmenopausal women (n = 61) | P value |
|-----------|-----------------------------|-------------------------------|--------|
| Age (years) | 28.48 ± 6.54 (25.6-30.4) | 62.39 ± 7.14 (60.5-64.2) | 0.0001 |
| BMI (kg/m²) | 22.34 ± 2.84 (21.5-23.2) | 25.36 ± 3.59 (24.4-26.3) | 0.0001 |
| AGD (mm) | 30.87 ± 2.98 (30.0-31.7) | 17.57 ± 2.18 (17.0-18.1) | 0.0001 |
| AGI | 1.40 ± 0.21 (1.3-1.5) | 0.70 ± 0.15 (0.6-0.7) | 0.0001 |
| VHI score | 23.95 ± 1.28 (23.6-24.3) | 10.75 ± 3.41 (9.9-11.6) | 0.0001 |
| FSFI score | 32.68 ± 2.25 (32.0-33.3) | 19.78 ± 5.46 (18.4-21.2) | 0.0001 |

The data is presented as mean ± standard deviation (95% confidence interval)

BMI: body mass index, AGD: anogenital distance, AGI: anogenital index, VHI: vaginal health index, FSFI: female sexual function index
Discussion

AGD has been frequently investigated in newborn infants and adult males but lacking evidences are present in women. Moreover, in animal tests, AGD was usually used to evaluate drug toxicity. For instance, phthalates are synthetic chemical compound used in cosmetics and it have been demonstrated as responsible of a shortening of AGD in rodents.

On the other hand, AGD length is positively influenced by male sex hormones, becoming longer.

In fact, AGD may be considered as an alternative biomarker of foetal testicular function, reflecting androgen action during the masculinisation process in animal models. This developmental androgen exposure allows normal differentiation and subsequent growth of male reproductive organs.

To our knowledge there is just another study investigating on AGD modifications before and after menopause. Lee and colleagues analysed 50 women (25 premenopause and 25 postmenopause) suggesting AGD and AGI as possible physical marker of menopausal aging of female genitalia.

As in our study, AGD was significantly longer in premenopausal women compared to postmenopausal women ($34.8 \pm 6.4$ vs. $30.3 \pm 6.6$, $P = 0.019$), AGI was significantly higher in premenopausal women than postmenopausal women ($1.7 \pm 0.4$ vs. $1.3 \pm 0.3$, $P < 0.0001$).

Menopause is responsible of several female physical changes having an important influence on the whole body,

Skin elasticity, bone production/resorption, female genitalia trophism or function and cardiovascular system are entirely influenced by hormonal changes occurring in postmenopause.

This study confirms that AGD in postmenopausal women was significantly shorter than AGD in premenopausal women, correlating with an increase of VVA and sexual impairment.

Probably, AGD length modification can be the result of a decrease in collagen production, atrophoderma, a drop-in water content and epidermal thickness reduction caused by the decreased production of sex hormones after menopause.

Conclusion

According to the results, AGD can be used as a metric to predict the changes related to menopause. Certainly, further studies are needed to confirm and standardize its use in clinical practice.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.
References

1. Mendiola J, Roca M, Mínguez-Alarcón L, Mira-Escalonó MP, López-Espín JJ, Barrett ES, et al. Anogenital distance is related to ovarian follicular number in young Spanish women: a cross-sectional study. Environ Health 2012; 11: 90.

2. Basaran M, Kosif R, Bayar U, Civelek B. Characteristics of external genitalia in pre- and postmenopausal women. Climacteric 2008; 11: 416–21.

3. Dean A, Sharpe RM. Clinical review: anogenital distance or digit length ratio as measures of fetal androgen exposure: relationship to male reproductive development and its disorders. J Clin Endocrinol Metab 2013; 98: 2230–8.

4. Barrett ES, Parlett LE, Redmon JB, Swan SH. Evidence for sexually dimorphic associations between maternal characteristics and anogenital distance, a marker of reproductive development, Am J Epidemiol 2014; 179: 57–66.

5. Swan SH, Sathyanarayana S, Barrett ES, Janssen S, Liu F, Nguyen RH, et al. First trimester phthalate exposure and anogenital distance in newborns, Hum Reprod 2015; 30: 963–72.

6. Courant F, Antignac JP, Maume D, Monteu F, Andersson AM, Skakkebaek N, et al. Exposure assessment of prepubertal children to steroid endocrine disrupters 1. Analytical strategy for estrogens measurement in plasma at ultra-trace level. Anal Chim Acta 2007; 586: 105–14.

7. Barrett ES, Parlett LE, Sathyanarayana S, Liu F, Redmon JB, Wang C, et al. Prenatal exposure to stressful life events is associated with masculinized anogenital distance (AGD) in female infants, Physiol Behav 2013; 114–115: 14–20.

8. Nappi RE, Palacios S, Particco M, Panay N. The REVIVE (REal Women’s Views of Treatment Options for Menopausal Vaginal ChangEs) survey in Europe: country-specific comparisons of postmenopausal women’s perceptions, experiences and needs, Maturitas 2016; 91: 81–90.

9. Nappi RE, Particco M, Biglia N, Cagnacci A, Di Carlo C, Luisi S, et al. Attitudes and perceptions towards vulvar and vaginal atrophy in Italian post-menopausal women: evidence from the European REVIVE survey. Maturitas 2016; 91: 74–80.

10. Bachmann G. Urogenital ageing: an old problem newly recognized, Maturitas 1995; 22 Suppl: S1–S5.

11. Lee D, Kim TH, Lee HH, Kim JM, Jeon DS, Kim YS. A pilot study of the impacts of menopause on the anogenital distance, J Menopausal Med 2015; 21: 41–6.

12. Meston CM. Validation of the female sexual function index (FSFI) in women with female orgasmic disorder and in women with hypoactive sexual desire disorder, J Sex Marital Ther 2003; 29: 39–46.

13. Wiegel M, Meston C, Rosen R. The female sexual function index (FSFI): cross-validation and development of clinical cutoff scores, J Sex Marital Ther 2005; 31: 1–20.

14. Welsh M, Saunders PT, Fiskin M, Scott HM, Hutchison GR, Smith LB, et al. Identification in rats of a programming window for reproductive tract masculinization, disruption of which leads to hypospadias and cryptorchidism, J Clin Invest 2008; 118: 1479–90.

15. Parks LG, Ostby JS, Lambricht CR, Abbott BD, Klinefelter GR, Barlow NJ, et al. The plasticizer diethylhexyl phthalate induces malformations by decreasing fetal testosterone synthesis during sexual differentiation in the male rat, Toxicol Sci 2000; 58: 339–49.

16. Gray LE, Jr., Wilson VS, Stoker T, Lambricht C, Furr J, Noriega N, et al. Adverse effects of environmental anti-androgens and androgens on reproductive development in mammals, Int J Androl 2006; 29: 96–104; discussion 5–8.

17. Salazar-Martínez E, Romano-Ráquer P, Yanez-Marquez E, Longnecker MP, Hernandez-Avila M. Anogenital distance in human male and female newborns: a descriptive, cross-sectional study, Environ Health 2004; 3: 8.

18. Wilson VS, Blystone CR, Hotchkiss AK, Rider CV, Gray LE, Jr. Diverse mechanisms of anti-androgen action: impact on male rat reproductive tract development, Int J Androl 2008; 31: 178–87.

19. Mendiola J, Stahlhut RW, Jorgensen N, Liu F, Swan SH. Shorter anogenital distance predicts poorer semen quality in young men in Rochester, New York, Environ Health Perspect 2011; 119: 958–63.

20. Pastuski V, Acerini CL, Dunger DB, Org KK, Hughes IA, Thankamony A, et al. Postnatal penile growth concurrent with mini-puberty predicts later sex—typed play behavior: evidence for neurobehavioral effects of the postnatal androgen surge in typically developing boys, Horm Behav 2015; 69: 98–105.

21. Thankamony A, Lek N, Carroll D, Williams M, Dunger DB, Acerini CL, et al. Anogenital distance and penile length in infants with hypospadias or cryptorchidism: comparison with normative data, Environ Health Perspect 2014; 122: 207–11.