Sex-specific ranges and ratios for anogenital distance among Thai full-term newborns

Nattakarn Numsriskulrat1*, Khomsak Srilanchakon1, Chaiyat Pronprechatham2, Sopon Pornkunwilai2 and Vichit Supornsilchai1

Abstract

Introduction: Anogenital distance (AGD) is a marker of prenatal androgen exposure and a tool for assessment of differences of sex development. Data for AGD in newborns have been published, but these findings may not be applicable to Thai newborns.

Aim: To provide the sex-specific ranges for AGD in Thai full-term newborns.

Methods: A cross-sectional study was conducted in term newborns in Thailand, during 2016–2018. AGD was measured from anus to anterior base of penis (AGDAP) and to perineoscrotal junction (AGDAS) in males and from anus to clitoris (AGDAC) and to posterior fourchette (AGDAF) in females. AGD ratio is defined as AGDAS divided by AGDAP in males and AGDAF divided by AGDAC in females.

Results: A total of 364 newborns were studied (male 51.4%). The mean AGDAS, AGDAP and AGD ratio in males were 25.20 ± 4.80, 52.60 ± 6.90 and 0.48 ± 0.08 mm, respectively. The mean AGDAF, AGDAC, and AGD ratio in females were 16.50 ± 3.90, 42.60 ± 6.20 and 0.39 ± 0.08 mm, respectively. There were significant differences between AGDAS and AGDAF, AGDAP and AGDAC, and AGD ratio between males and females (p < 0.001). The AGDAC, AGDAF, AGDAS, AGDAP were correlated with birth weight and length, but AGD ratio showed no correlation.

Conclusion: The sex-specific ranges for AGD in Thai full-term newborns were determined. AGD ratio is a useful marker of prenatal androgen exposure since it differs between sexes, but constant between races and did not vary by body size.

Keywords: Anogenital distance (AGD) ratio, Anopenile distance (AGDAP), Anoscrotal distance (AGDAS), Anoclitoral distance (AGDAC), Anofourchette distance (AGDAF)

What is already known on this topic

1. Anogenital distance is a marker for prenatal androgen exposure during the critical period of external genital development (8th-16th week of gestation).

What this paper adds

1. The AGD ratio is a more reliable indicator of prenatal androgen exposure than other AGD parameters since it is consistent between races, different between sexes, and unrelated to anthropometrics.
2. Reference ranges of AGD in healthy Thai newborn.
Introduction
The distance between the anus and genitalia is referred to as the anogenital distance (AGD), which serves as a marker of prenatal androgen exposure [1]. Under-androgenization or over-androgenization can be inferred indirectly from AGD. For example, an under-androgenized newborn has a shorter AGD than normal [1]. Male offspring born to polycystic ovary syndrome (PCOS) mothers are more likely to have longer AGD due to increased androgen levels during the 8th-16th gestation weeks during pregnancy which is the critical period of external genitalia development [2]. AGD is also longer in androgenized females due to labioscrotal fusion, as demonstrated in neonates with virilized congenital adrenal hyperplasia [3].

Gender, gestational age, and anthropometric characteristics including birth weight and length are all significant correlates to AGD [4, 5]. AGD is becoming more widely used in clinical settings to assess potential reproductive dysfunction. Since AGD can also be affected by endocrine disruptors, it has been utilized in environmental toxicology to assess the health implications of chemicals such as Bisphenol A (BPA) and Polychlorinated Bisphenyls (PCBs) with endocrine-altering capabilities [6]. While data exists on Caucasian and selected Asian newborn populations, there is a paucity of data among newborns in Thailand to determine normal distance ranges and ratios by gender. In this study, the sex-specific reference ranges for AGD in healthy Thai full-term newborns were presented including comparisons to previous studies both locally and worldwide.

Materials and methods
Participants
A cross-sectional study was undertaken among newborns in Thailand from 2016 to 2018. Healthy newborns aged zero to 72 h were enrolled in the study. Exclusion factors included the presence of genital ambiguity, dysmorphic features, and known maternal ingestion of androgenic medications or substances.

The recommended minimum sample size to determine statistically significance was calculated using the formula $N = (2\sigma / E)^2$ [7].

$Z$ = the value from the standard normal distribution reflecting the confidence level ($Z = 1.96$ for 95%)

$\sigma$ = standard deviation from the reference study.

$E$ = desired margin of error.

To estimate the SD for the present study, we used data published from a reference study where the mean anoscrotal distance was 24.7 ± 4.5 mm [8]. A margin of error of 0.5 mm was considered.

$N = (1.96(4.5)/0.5)^2 = 312$

Anthropometric measurements of birth weight, supine length, head and thoracic circumferences and AGD of all newborns were performed by well-trained physicians. Weight was measured using a digital infant scale with clothing and diapers removed to the nearest 0.01 kg. With a Harpenden Infantometer, length was measured to the nearest 0.1 cm. The newborn was placed naked on the infantometer in a supine position, the head was held against the immovable head-board by an assistant, the knees were held together and held down against the board surface, and the heels were also held down to ensure the child’s body and pelvis were straight along the measuring device, the foot-board was then drawn up to meet the heels and this was measured as the length. A flexible measuring tape was used to obtain head and thoracic circumferences. A complete physical examination of each baby occurred in a warmed environment. The obstetric history of each child was obtained.

Genital distance measurements
A digital sliding caliper (SuperCaliper SERIES 500, Mitutoyo, Thailand) was used to measure AGD to the nearest 0.1 mm. Newborns were placed in a supine position. An assistant held both hips in flexion, flexed and pulled the knees back towards the shoulders. The caliper was positioned adjacent to the surface of the genitalia, digital screen turned away from the researcher, and the single AGD measurement was obtained.

In males, AGD was measured from the center of the anus to the anterior base of the penis (anopenile distance, AGDAP) and to the perineoscrotal junction (anoscrotal distance, AGDAC). In females, AGD was measured from the center of the anus to the clitoris (anoclitoral distance, AGDAc) and to the posterior fourchette (anofourchette distance, AGDAf) (Fig. 1). The anogenital distance ratio (AGD ratio) is calculated as AGDAc divided by AGDAp in males and AGDAf divided by AGDAC in females.

Statistical analysis
For continuous variables, we calculated means and standard deviations, whereas for categorical variables, we calculated frequencies and percentages. The unpaired t-test was used to test for means differences between continuous variables. Pearson’s correlation coefficient (r) assessed the correlation between continuous variables. Associations between variables were evaluated using linear regression. A $p$-value < 0.05 was considered statistically significant. All statistical data were analyzed using SPSS software version 22 (SPSS, Inc., Chicago, IL, USA).
**Results**

A total of 364 newborns were included in the study. There were 187 male newborns (51.4%) and 177 female newborns (48.6%). The mean ± SD gestational age was 38.69 ± 1.09 weeks (range, 37–41 weeks). Anthropometrics data including weight, length, and head circumference are not significantly different between males and females by Student’s t-test. The mean ± SD AGD_A, AGD_AP, and AGD ratio in males were 25.20 ± 4.80, 52.60 ± 6.90 and 0.48 ± 0.08 mm, respectively. The mean ± SD AGD_A, AGD_AP, and AGD ratio in females were 16.5 ± 3.9, 42.6 ± 6.2 and 0.39 ± 0.08 mm, respectively (Table 1). From the Student’s t-test, there were significant difference between AGD_A and AGD_A, AGD_AP and AGD_AC, and AGD ratio between males and females (p < 0.001). The AGD percentiles of subjects are shown in Table 2 at five percentiles (3%, 10%, 50%, 90%, 97%).

Table 3 presents the findings of Pearson’s correlation analysis used to determine the correlation between AGD parameters and anthropometrics. There was no correlation between the AGD ratio and birth weight or length in both sexes. The AGD_A, AGD_AP, AGD_A, and AGD_AC, however, had a statistically significant but weak positive correlation with birth weight and length.

**Discussion**

AGD is a marker to assess under-androgenization or over-androgenization. Male infants tend to have a shorter AGD than predicted if they have undescended testes, hypospadias, or prenatal exposure to antiandrogenic endocrine disrupting chemicals such as phthalate [9, 10]. In contrast, longer AGD can be found in virilized females. Female babies with virilizing congenital adrenal hyperplasia were found to have labioscrotal fusion and an increased anogenital ratio [3]. A study demonstrated that maternal cigarette smoking is associated with increased weight-adjusted AGD in female neonates [11]. This is likely due to the suppression of

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**Table 1** Anthropometric parameters and anogenital distances of the participants (N= 364)

| Parameter                  | Mean  | SD   | Median | Min | Max  |
|---------------------------|-------|------|--------|-----|------|
| Birth weight, gm          |       |      |        |     |      |
| Male                      | 3190.85 | 390.28 | 3150.00 | 2330.00 | 4380.00 |
| Female                    | 3151.29 | 414.30 | 3110.00 | 2260.00 | 4400.00 |
| Length, cm                |       |      |        |     |      |
| Male                      | 51.4  | 1.9  | 51.0   | 45.5 | 57.0 |
| Female                    | 51.0  | 2.0  | 51.0   | 47.0 | 57.0 |
| Head circumference, cm    |       |      |        |     |      |
| Male                      | 33.4  | 1.3  | 33.5   | 23.5 | 37.0 |
| Female                    | 33.2  | 1.2  | 33.0   | 30.0 | 37.0 |
| Chest circumference, cm   |       |      |        |     |      |
| Male                      | 32.4  | 1.49 | 32.0   | 29.0 | 37.0 |
| Female                    | 32.3  | 1.52 | 32.0   | 29.0 | 37.0 |
| Anogenital distances      |       |      |        |     |      |
| Male (N= 187)             |       |      |        |     |      |
| AGD_A*, mm                | 25.2  | 4.8  | 25.0   | 10.0 | 40.0 |
| AGD_AP**, mm              | 52.6  | 6.9  | 50.0   | 34.0 | 80.0 |
| AGD ratio***              | 0.48  | 0.08 | 0.50   | 0.25 | 0.67 |
| Female (N= 177)           |       |      |        |     |      |
| AGD_AF*, mm               | 16.5  | 3.9  | 15.0   | 6.0  | 25.0 |
| AGD_AC**, mm              | 42.6  | 6.2  | 40.0   | 27.0 | 60.0 |
| AGD ratio***              | 0.39  | 0.08 | 0.38   | 0.20 | 0.57 |

*, **, ***: Student’s t-test; p < 0.001
placental aromatase, a key enzyme responsible for the conversion of androgen to estrogen [12, 13].

The present study showed that there was a significant difference between AGD parameters in males and females. Sexual dimorphism of AGD is observed as males having a longer AGD than females. These findings may be explained by the fact that external genitalia are developed under the influence of sex hormones [14], particularly during the masculinization programming window which occurs between 8 and 14 weeks of pregnancy in humans [15]. Because of this period of prenatal androgen action, the distance between the anus and the base of the genital tubercle is approximately twice as long in males as females in both rodents and humans [16]; however, there are some variations, as Özkan B. et al. [17] found that the ratio of AGDAS in males to AGDAF in females was 2.2, while it is around 1.6 in our study.

The AGD measurements in term newborns from different countries are presented in Table 4. The mean AGDAS in male (25.2 mm.) and AGDAF in female (16.5 mm.) in the present study are comparable with the data from the USA reported by Sathyanarayana S and colleagues (mean AGDAS 24.7 mm. and mean AGDAF 16.0 mm.). [8], but much higher than the mean value reported by Shah R. et al. [18] (mean AGDAS 21.0 mm. and mean AGDAF 13.0 mm.) and the UK study [16] (mean AGDAS 19.8 mm. and mean AGDAF 9.1 mm.). The variation in AGD parameters between studies may be explained by ethnic differences, equipment, systematic errors in measurement, and measurement protocols.

Various infant positionings were used during AGD measurement in previous studies, which may result in different AGD values. The two most common technique from the literature are the TIDES method and the Cambridge method. Both protocols require the assistance of another person to hold the infant in a supine position, but the differences are in the posture of the lower half of the body. The TIDES method places the newborn's legs in a frog-like position and pulls the knees back toward the shoulders, whereas the Cambridge method places both hips in flexion, puts the feet on the surface, and exerts light pressure onto the thighs [22]. The TIDES method creates a slight stretch of the newborn's perineum during measurement, resulting in a longer AGD value than the

| Table 2 Percentiles of AGD among males and females |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Percentile | Male | Female | Male | Female | Male | Female |
| AGDAS (mm) | AGDAP (mm) | AGD ratio | AGDAS (mm) | AGDAP (mm) | AGD ratio |
| 3 | 15 | 40 | 0.33 | 10 | 30 | 0.22 |
| 10 | 20 | 45 | 0.38 | 10 | 35 | 0.25 |
| 50 | 25 | 50 | 0.50 | 15 | 40 | 0.38 |
| 90 | 30 | 60 | 0.60 | 20 | 50 | 0.50 |
| 97 | 35 | 70 | 0.62 | 25 | 55 | 0.50 |

Abbreviations: AGDAS Anopenile distance, AGDAP Anoscrotal distance, AGDAC Anoclitoral distance, AGDAF Anofourchette distance
AGD ratio: AGDAS/AGDAP (male), AGDAP/AGDAC (female)

| Table 3 Correlation of AGD and AGD ratio with anthropometric parameters |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| BW (Kg) | Length (cm) | HC (cm) |
| **Male** | | | | | |
| AGDAS (mm) | 0.265 | <0.001 | 0.259 | <0.001 | 0.032 | 0.665 |
| AGDAP (mm) | 0.235 | <0.001 | 0.260 | <0.001 | 0.076 | 0.299 |
| AGD ratio | 0.104 | 0.153 | 0.077 | 0.295 | -0.039 | 0.592 |
| **Female** | | | | | | |
| AGDAS (mm) | 0.433 | <0.001 | 0.311 | <0.001 | 0.146 | 0.05 |
| AGDAP (mm) | 0.340 | <0.001 | 0.249 | 0.001 | 0.232 | 0.002 |
| AGD ratio | 0.081 | 0.276 | 0.070 | 0.352 | 0.166 | 0.025 |

Pearson’s correlation coefficient (r) was used to assess the correlation
Abbreviations: AGDAS Anopenile distance, AGDAP Anoscrotal distance, AGDAC Anoclitoral distance, AGDAF Anofourchette distance
AGD ratio: AGDAS/AGDAP (male), AGDAF/AGDAC (female)
Table 4  Literature review of AGD reference ranges in term newborns of both sexes among different countries

| Countries                        | Female | Male          |                |                |
|----------------------------------|--------|---------------|----------------|----------------|
|                                  | AGDAF (mm) | AGDAC (mm) | AGD ratio      | AGDAF (mm) | AGDAC (mm) | AGD ratio |
| USA [18] (2021)                  | 13.0±2.0 | 35.0±3.0  | 0.37±0.07      | 21.0±4.0    | 50.0±4.0   | 0.42±0.07 |
| USA [8] (2015)                   | 16.0±3.2 | 36.7±3.9  | -              | 24.7±4.5    | 49.6±5.9   | -         |
| European countries (Multicenter) [19] (2020) | 14.8±3.5 | 37.8±4.5  | 0.39±0.10      | 24.6±4.7    | 47.6±5.8   | 0.49±0.10 |
| Turkey [17] (2011)              | 10.3±0.2 | 30.0±0.2  | 0.30±0.10      | 23.0±0.6    | 56.0±1.0   | 0.48±0.80 |
| Ghana [5] (2017)                 | 13.6±2.7 | 34.2±3.3  | -              | 25.5±5.0    | 48.9±5.6   | -         |
| UK [16] (2009)                  | 9.1±2.8  | -           | -              | 19.8±6.1    | -          | -         |
| Nigeria [20] (2019)             | -       | -           | -              | 25.5±3.9    | 48.7±3.9   | -         |
| Korea [21] (2015)               | -       | -           | -              | 23.0±2.0    | 42.0±3.0   | -         |
| Present study                    | 16.5±3.9 | 42.6±6.2  | 0.39±0.08      | 25.2±4.8    | 52.6±6.9   | 0.48±0.08 |
| Summary*                         | 13.2±2.9 | 35.9±3.9  | 0.37±0.09      | 23.7±4.8    | 49.7±5.2   | 0.47±0.36 |

Values are expressed as mean ± SD

* Weighted mean ± pooled variance

Abbreviations: AGDAF, Anoscrotal distance; AGDAC, Anocrotal distance; AGDAC, Anoclitoral distance; AGDAF, Anofourchette distance

AGD ratio: AGDAF/AGDAC (female)

Cambridge method [22]. In this study, AGD parameters were assessed using only a position similar to the TIDES technique; therefore, future research may be needed to develop the position-specific reference ranges for AGD.

Our findings support previous study that found significant but weak positive correlations between AGDAF, AGDAC, AGDAF, AGDAC, and birth weight and length in term newborns [5]. In that study, the strongest correlation was found between AGDAF and birth weight in male newborns (r=0.306; p<0.001), whereas our study found the strongest correlation between AGDAF and birth weight in female newborns (r=0.433; p<0.001). Mondal, et al. [23] measured the AGDAF in term and preterm female newborns in India and found a weak positive correlation between the AGDAF and birth weight (r=0.232, p<0.001), length (r=0.165, p=0.008), and head circumference (r=0.225, p<0.001). A number of studies have also discovered variable positive correlations between AGD and birth weight and length [3, 16–21].

Despite the fact that the mean AGDAF, AGDAC, AGDAF, and AGDAC varied across studies, we found a consistent AGD ratio [8, 16–21]. Ranges of mean AGD ratio in female and male newborns were 0.30–0.39 and 0.42–0.49, respectively. In our study, the mean (±SD) AGD ratio is 0.48±0.08 and 0.39±0.08 in males and females, closest to what was reported in a European multicenter study of 668 term babies which also found a significant difference between AGD ratios in males (0.49±0.1) and females (0.39±0.1) and intermediate values in differences of sex development (DSD) (0.43±0.1) [19]. Moreover, AGD ratio, unlike other AGD parameters, was unaffected by birth weight or length, suggesting that it might be a better marker for determining the degree of prenatal androgen exposure in a full-term newborn than the distance measures alone. However, the AGD ratios in Table 4 were largely from southeastern Europe and southwestern Asia, and the availability of the data in full-term newborns from other regions are limited, which could impact the generalizability of the AGD ratio values, therefore, more study on the AGD ratio is necessary.

AGD can be utilized to gain insight into the effect of androgens during pregnancy, to assess newborns with DSD, and to assess the health implications of endocrine disruptors in environmental toxicology. Normative data on AGD ranges for local references should be established as standards for comparison in clinical practice. We offered five percentile thresholds in our results as an initial standard for Thailand.

This study is the first to present data and standards for term newborns in Thailand. The limitation of our research is the single-center study and the reliance on a single measurement of AGD. However, the reliability of this measurement in humans has been well-established, and since one examiner performed all of the measurements, interobserver errors were minimized.

Conclusion
The present study provided sex-specific ranges and ratio for AGD in Thai healthy full-term newborns. We proposed using the AGD ratio, instead of individual AGD, as an indicator of prenatal androgen exposure.
Abbreviations
AGDAP: Anopenile distance; AGDAS: Anoscrotal distance; AGDAC: Anoclitoral distance; AGDDe: Anofourchette distance; AGD ratio: AGDAS/AGDAP (male), AGDDe/AGDAC (female).

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Authors’ contributions
V.S. designed the study and revised the final manuscript. N.N. reviewed the literature, analyzed the data, and wrote the first draft of manuscript. K.S., C.P. and S.P. enrolled participants, collected clinical data and measured AGD. All of the authors have read, discussed and approved the final version of the manuscript.

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Availability of data and materials
The datasets generated and/or analyzed during the current study are not publicly available due ethical issue but are available from the corresponding author on reasonable request.

Declarations
Ethics approval and consent to participate
Written informed consent were obtained from the parents or guardians of the study participants. All methods in this study were performed in accordance with the Declaration of Helsinki. This study was approved by the Institutional Review Board of the Faculty of Medicine, Chulalongkorn University (IRB No. 401/60).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1 Division of Pediatric Endocrinology, Department of Pediatrics, Faculty of Medicine, Chulalongkorn University, Bangkok 10330, Thailand. 2 Police General Hospital, Bangkok, Thailand.

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References
1. 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(6):2230–8.
2. Glintborg D, Jensen RC, Schmedes AV, Brandslund I, Kyhl HB, Jensen TK, et al. Anogenital distance in children born of mothers with polycystic ovary syndrome: the Odense Child Cohort.Hum Reprod. 2019;34(10):2061–70.
3. Callegari C, Everett S, Ross M, Brasel JA. Anogenital ratio: measure of fetal virilization in premature and full-term newborn infants. J Pediatr. 1987;111(2):240–3.
4. Salazar-Martínez E, Romano-Riqueur 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(1):8.
5. Asafo-Agyei SB, Ameyaw E, Chanoine JP, Zacharin M, Ngubu SB, Jarrett OO. Anogenital Distance in Term Newborns in Kumasi. Ghana Horm Res Paediatr. 2017;88(6):396–400.

6. Liu C, Xu X, Huo X. Anogenital distance and its application in environmental health research. Environ Sci Pollut Res Int. 2014;21(8):5457–64.
7. WW. L. Sample Size for One Sample, Continuous Outcome: Boston University School of Public Health. 2020.
8. Sathyarayana S, Grady R, Redmon JB, Ivcek K, Barrett E, Janssen S, et al. Anogenital distance and penile width measurements in The Infant Development and the Environment Study (TIDES): methods and predictors. J Pediatr Urol. 2015;11(2):76 e1-6.
9. Swan SH, Main KM, Liu F, Stewart SL, Kruse RL, Calafat AM, et al. Decrease in anogenital distance among male infants with prenatal phthalate exposure. Environ Health Perspect. 2005;113(8):1056–61.
10. Zarean M, Keiltha M, Feizi A, Kazemitabae M, Kelishadi R. The role of exposure to phthalates in variations of anogenital distance: a systematic review and meta-analysis. Environ Pollut. 2019;247:172–9.
11. Kizilay D, Aydin C, Aygün AP, Tuhan H, Okumakan O. Prenatal smoke exposure is associated with increased anogenital distance in female infants: a prospective case-control study. J Pediatr Endocrinol Metab. 2021;34(1):79–88.
12. W. H. Sex Hormones and Human. Principle and Practice of Endocrinology and Metabolism 3rd ed. 2002. p. 219–359.
13. Tortola AT, Väärsämäki M, Lehtinen M, Zeleniuch-Jacquotte A, Lundin E, Rodgers KG, et al. Determinants of maternal sex steroids during the first half of pregnancy. Obstet Gynecol. 2011;118(5):1029–36.
14. Jain VQ, Singal AK. Shorter anogenital distance correlates with undescended testis: a detailed genital anthropometric analysis in human newborns. Hum Reprod. 2013;28(9):2343–9.
15. Welsh M, Suzuki H, Yamada G. The masculinization programming window. Endocr Dev. 2014;27:17–27.
16. Thankamony A, Öng KK, Dungar DB, Acerini CL, Hughes IA. Anogenital distance from birth to 2 years: a population study. Environ Health Perspect. 2009;117(11):1786–90.
17. Özkán B, Konak B, Cayer A, Konak M. Anogenital distance in Turkish newborns. J Clin Res Pediatr Endocrinol. 2011;3(3):122–5.
18. Shah R, Alishakib B, Schall JI, Kelly A, Ford E, Zemel BS, et al. Endocrine-sensitive physical endpoints in newborns: ranges and predictors. Pediatr Res. 2021;89(3):660–6.
19. van der Straaten S, Springer A, Zecic A, Hebenstreit D, Toninhofer U, Gawlik A, et al. The External Genitalia Score (EGS): a European multicenter validation study. J Clin Endocrinol Metab. 2020;105(3):dgz142.
20. Adebayo AO, Feturuga MB, Jarrett OX, Ogunlesi TA, Chanoine JP, Oba-Daini O. Normative data on penile and anogenital measurements of term male infants in Sagamu, Nigeria Acta Paediatr. 2019;108(11):2041–7.
21. Park JY, Lim G, Oh KW, Ryu DS, Park S, Jeon JC, et al. Penile length, digit length, and anogenital distance according to birth weight in newborn male infants. Korean J Urol. 2015;56(3):248–53.
22. Fischer MB, Ljubicic ML, Hagen CP, Thankamony A, Öng K, Hughes I, et al. Anogenital distance in healthy infants: method-, age- and sex-related reference ranges. J Clin Endocrinol Metab. 2020;105(9):3096–3004.
23. Mondal R, Chattjeee K, Samanta M, Hazra A, Ray S, Sabui TK, et al. Clitoral length and anogenital ratio in Indian newborn girls. Indian Pediatr. 2016;53(4):299–303.

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