Levonorgestrel correlates with less weight gain than other progestins during hormonal replacement therapy in Turner Syndrome patients

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Turner Syndrome (TS) is associated with an increased risk of cardiovascular and metabolic complications. Furthermore, TS women need hormone replacement therapy (HRT), of which progestins can influence body weight. We aimed to analyze the metabolic and weight profile in a cohort of 111 TS women. They started receiving estrogen at 15.8 (± 3.6) years old, with no change in hypertension, dysglycemia, and dyslipidemia incidence but with a tendency to increase overweight (p = 0.054). As the first used type of progestin, most had received cycles of 10 days per month of medroxyprogesterone (MPA) or levonorgestrel (LNG), then shifted to micronized progesterone (MP), which has currently become the most used one. By multiple linear regression analysis, we found that the prolonged use of MPA, LNG, or MP showed no metabolic change except for weight gain. The percentage of annual BMI increment was positive for all progestins used in TS women (MPA 2.2 ± 2.2; LNG 0.2 ± 1.2; and MP 2.2 ± 2.6 kg/m2), but LNG seemed to best prevent on weight gain over time (p < 0.05). In conclusion, metabolic comorbidities are prevalent in TS even before the HRT regimen, and LNG performed better on less weight gain than MPA and MP in our cohort of the TS population.

Turner Syndrome (TS) is a chromosomal sex disease that affects 1:2,500–1:4,000 females. It is characterized genetically by missing or structurally altered one of the X chromosomes, and clinically by ovarian failure, infertility, and short stature. TS diagnosis is confirmed by the typical karyotype 45,X, or partial monosomy with or without mosaicism1–4. The X chromosome monosomy incurs in gonadal dysgenesis, comprising 85% of patients with primary amenorrhea and 98% infertility. Although Turner girls may present complete puberty spontaneously in 15%, breast development may reach 48%5. TS adolescents present marked hypergonadotropic hypogonadism, so puberty development usually needs pharmacological induction at about the age of 126.

TS per se is associated with a three to fourfold increase in mortality related to cardiovascular complications and increased risk of metabolic diseases such as dysglycemia, dyslipidemia, hypertension, and obesity2,4,6. Hypertension affects up to 60% of this population, and there is a type 2 diabetes (T2D) risk increase in two to four times. The body mass index (BMI) is higher than the general population not only because of the short stature but also related to more visceral adiposity. Furthermore, some studies found an increased incidence of hypertriglyceridemia, low HDL, elevated and more atherogenic LDL, independently of being already obese1,2. Nonalcoholic fatty liver disease (NAFLD) is often found, and it is related to insulin resistance (IR)7.

Many factors have been studied as possible influences for developing metabolic comorbidities. Hypogonadism is a factor that contributes to the increase in metabolic diseases in TS women. Besides, it is associated with low bone mineral density and psychological variations1. Hypothyroidism is the most frequent autoimmune disease affecting this population, but it does not seem to modify the lipid profile or BMI. Also, the treatment with

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somatotropin improves stature without changing corporal composition, independently of dose, duration, or age at onset of the therapy’s beginning.

The hypogonadism treatment is based on hormonal replacement therapy with estrogen (E) plus progestin (P). It is well known that E is essential not only for puberty induction and maintenance but also for bone mass gain and metabolic improvement. Both oral and transdermal routes have the same effect over corporal composition, blood pressure, hepatic enzymes, bones, carbohydrate metabolism, and lipidic profile, except for one study reporting higher HDL after taking oral estradiol. Transdermal estradiol (TE2) has shown to be the most appropriate E for replacement therapy and its pharmacokinetics prevents hepatic first-pass and achieves physiological E2 serum concentration with a lower risk of thrombosis.

Progestins input should start around two years after estrogen replacement or after the first induced menes. However, progestins may have effects on androgenic, glucocorticoid, and mineralocorticoid receptors other than progesterone receptor (PR). As a result, the continuous use of progestin (P), substantially present in contraceptive pills, more markedly when with medroxyprogesterone acetate (MPA), is associated with weight gain and decreases in lean mass.

Therefore, it is of great importance to ensure that the hormone replacement therapy (HRT) offered to TS women is no longer a factor of cardiometabolic risk increase as it is going to be used for a long time, at least until the usual menopause age of general population.

Based on this, we performed a retrospective study to analyze the correlation between TS’s metabolic profile and the use of oral (P), and to examine the best therapeutic alternative to be employed in HRT in this population. The hypothesis addressed in this study is that MPA would contribute to the worsening of metabolic risks in TS patients, given that it also has glucocorticoid activity.

Patients and Methods

Patients. We performed a cross-sectional and non-controlled retrospective study of the TS cohort followed at the Endocrinology Outpatient Clinics of Hospital São Paulo from the Universidade Federal de São Paulo. All of them have a diagnosis based on the clinical profile and karyotype. They signed the informed consent to participate in this study after the protocol was properly approved by the Ethics Committee in Research (research number 1146/2015). All methods were performed in accordance with the relevant guidelines and regulations.

All enrolled TS patients were on HRT. Of note, not having an ideal control group is justified because of the limitation of having TS adults who had not received HRT, which of course it would be non-ethical. We excluded TS patients who had used norethisterone, gestodene, cyproterone acetate, and drospirenone as contraceptive pills prescribed by other services.

We evaluated retrospectively 123 medical records from TS patients that started HRT. However, 12 of them were excluded due to inconsistent clinical information (addressed from other services), irregular follow-up, still on growth hormone or non-compliance with levothyroxine (LT4) therapy. This cross-sectional analysis comprises Phase 1, with which we identified general clinical information regarding the percentage of 45,X karyotype, current age in years, if hypothyroid on regular treatment, GH therapy before the start of HRT, metabolic data (anthropometric, hypertension, lipids profile, glycemic measurements), and the age at estrogen and progestin onset.

We further stratified them in three other retrospective analytical phases. Phase 2.1 comprised those from the very beginning of estrogen introduction until the start of the progestin association. Then, we only considered patients in regular HRT and with documented estrogen and progestin start time to be able to precisely evaluate the type and age at HRT onset used to pubertal induction and the time of the last outpatient follow-up visit.

Moreover, we selected those to study the progestin effect on metabolic comorbidities over time, being Phases 2.2 and 2.3 the short and long use of progestin, respectively. Patients who had changed their type of progestin entered Phase 2.2 and Phase 2.3 study more than once. Figure 1 describes the number and characteristics of TS patients enrolled in each analysis. For the Phase 2.2, characterized by HRT with estrogen and within the first year of progestin, we included those patients with available metabolic data from the last consultation before using progestin and exactly 1 year after use. For Phase 2.3, the long retrospective analysis of estrogen and progestin HRT, we included those patients in regular HRT with available metabolic data from the time just before starting progestin until the last examination. TS patients without progestin or less than one year use, in irregular use, or with any medical condition or medication, such as corticosteroids or insulin, which could have interfered with metabolic parameters, were excluded.

Clinical and lab evaluation. All data were collected from the medical records including karyotype, age at which estrogen and progestin started, type of estrogen used to induce puberty and at the last medical visit, whether progestin was changed, and what had been used until last outpatient visit, the period of time between estrogen and progestin introduction, time of progestin exposure (total and for each progestin used), and metabolic parameters before HRT, after 1 year of each progestin, and longtime use based on the last follow-up visit.

For each situation, clinical parameters were evaluated by measuring the weight and height for body mass index (BMI) calculation (weight square of height) or for the BMI z-score (Standard Deviation Score) if under 20 yo; also the systolic blood pressure (SBP) and diastolic blood pressure (DBP) were noted. The z-score was calculated with the WHO app, Anthro-Plus (http://www.who.int/childgrowth/software/en/). Laboratory data were retrieved for fasting glycemia (FG), triglycerides (TG), high-density lipoprotein (HDL), low-density lipoprotein (LDL), TSH, and renal function.

We followed standard altered clinical and lab metabolic parameters as defined: hypertension when using of anti-hypertensive drugs or SBP ≥ 140 mmHg or DBP ≥ 90 mmHg in adults or if greater than 95th Percentile when age under 20yo; overweight when BMI > 25Kg/m² in adults or z-BMI > 1 if age inferior to 20yo; obesity was defined as BMI ≥ 30Kg/m² in adult patients or if standard deviation score body mass index (z-BMI) ≥ 2 if age inferior to
20 yo; dysglycemia with pre-diabetes when FG 100–125 mg/dl or HbA1C 5.7–6.4%, or diabetes when FG ≥ 126 mg/dl or HbA1C > 6.5% or in use of an oral antidiabetic [10]; dyslipidemia when HDL-C < 50 mg/dl (HDL < 45 before 19 years old), LDL-C ≥ 160 mg/dl, TG ≥ 150 mg/dl (TG ≥ 130 before 19 years old) or in statin use [13].

Progestin use evaluation. Since TS patients used more than one progestin in different periods of the HRT follow-up, either due to health system availability, personnel or lack of family financial resources, and adverse side effects, we have developed a method to combine and normalize the period and the dose of progestin use by creating a unit for measuring the progestin exposure by the Standard Progestin Cumulative Dose (SPCD).

We applied SPCD rule to normalize combined and interposed most common oral progestins used in each HRT for 10-day-per-month cycles within one year: Medroxyprogesterone Acetate (MPA), Levonorgestrel (LNG) and Micronized Progesterone (MP). Also, we assumed that the daily dose, 10 days of use per month, and years of use have equal influence as progesterone effect. One SPCD unit for each MPA (5 mg/day × 10 days per month × 1 year = 5 mg/10d/yr.), LNG (250 mg/day × 10 days per month × 1 year = 250 mg/10d/yr.), and MP (100 mg/day × 10 days per month × 1 year = 100 mg/10d/yr.) was considered equalized. As a result, a hypothetical patient who used MPA on the scheme of 10 mg/10d/month for 1 yr has an SPCD of 2, which was the same as a patient who has 10 mg/5d/month for 2 yr, LNG 0.25 mg/10d/month for 2 years, or MP 200 mg/10d/month for 1 year.

Statistical analysis. Data were presented as Mean ± Standard Deviation for numeric parametric or Median (Minimum-Maximum) for non-parametric variables and by number and percentage for qualitative variables. Shapiro Wilk test was applied to check for normal population distribution, with equal variances among individuals (p-value 0.05). A comparison of means was performed using Student’s paired t-test or One Way ANOVA complemented with Bonferroni’s Test for all pairwise comparison, when there was normal data distribution. Multiple Linear Regression was applied for annual variation in body mass index relative to pre-progestin BMI versus type of progestin. Wilcoxon Signed Rank Test was applied for before and after treatments comparison. The Contingency Tables were used to determine whether or not the distribution of each group was contingent on the categories it fell in, and the Fisher Exact Test was applied. Yates Correction Factor was used for a more accurately computed p-value in the significance cases. A significance level of 0.05 was adopted for all analyses. The power considered was greater than 0.80. The analyses were performed using the software’s Sigma Plot 13.2 for Windows (Systat Software Inc.), Statistical Package for Social Sciences (SPSS version 24), Minitab 16, and depicted in Excel Office 2010.

Results

General TS cohort features on hormone replacement therapy (HRT). At the time of puberty induction, based on 73 TS patients with pediatrics available documentation, most TS patients (45/73; 62%) received conjugated equine estrogens (CEE), followed by estradiol valerate (VE2). Nowadays, among 111 patients using estrogen replacement, 56 (50%) are using transdermal 17β-estradiol (TE2), 51 (46.4%) VE2, and 4 (3.6%) Ethinyl estradiol (EE2). Regarding the type of progestin, 62/73 (56%) used medroxyprogesterone acetate (MPA) at the
They were more exposed to the progestin as shown in Table 3. We corrected this bias in body weight change for with MPA and MP was similar, as shown in Fig. 2.

Percentage variation on BMI was significantly lower with LNG than those who had used MPA or MP. Weight gain during the time evaluated, dividing weight variation by the years of exposition. We then observed that the mean annual change in these parameters after one year-use of each SPCD.

The association between the type of progestin and weight change was confirmed by using a multiple linear regression model. For that, we applied the annual percentual variation in BMI as the dependent variable and logistic variables LNG, age, duration of treatment, and cumulated exposition (SPCD) as independent ones. Annual BMI variation \( \frac{\text{BMI}_{\text{after}} - \text{BMI}_{\text{before}}}{\text{years}} \) correlated positively with BMI (rs 0.30; \( p = 0.0144 \)).

Table 1. Overall clinical features of the cohort at basal analysis - Phase 1. Karyotype distribution, age (mean ± SD), the prevalence of thyroid dysfunction, GH and HRT use in the 111 girls with Turner syndrome. Data showed either as mean ± SD or N/total (%).

| Clinical feature | Number of subjects |
|------------------|--------------------|
| 45, X Karyotype (\%) Non-45, X, n (%) | 79/111 (71.2)/32/111 (28.8) |
| Current age in years | 25.7 ± 12.2 |
| Hypothyroidism on regular treatment | 29/111 (26.1) |
| GH therapy before HRT | 75/111 (67.6) |
| Age at estrogen onset in years (range) | 15.8 ± 5.6 (9.6–29.2) |
| Age at progestin onset in years (range) | 18.2 ± 5.7 (12.5–29.6) |
| Time between estrogen and progestin onset, in years | 2.17 ± 2.1 |

Weight gain and the type of used progestin. Initially, we studied the weight change of 73 TS patients with a mean age of 22.0 ± 6.3 yo, regardless of the type of used progestin in HRT. Then, we were able to select 61/73 TS patients who had information of the precise period of each type of progestin, in an attempt to stratify the analysis per subtype of progestin concerning its influence in weight gain. We found that using MPA increased weight more than LNG or MP (3.9 ± 5.81 vs. 3.0 ± 2.75 and 1.5 ± 2.24, respectively; \( p < 0.05 \)). However, evaluating only young TS patients 24/73 (<19 yo), with a mean age of 16.6 ± 1.41 yo, using progestin for 3.6 ± 2.9 years (from 1 to 8.8 y), there was no significant change in z-BMI scores (Mi = 0.04, from −0.79 to 3.50). In this young subgroup, 16 were taking MPA, 3 LNG and 5 MP, and comparing the rate of z-BMI change for each type of P (from 1 to 8.8 y), there was no significant change in z-BMI scores (Mi = 0.04, from −0.79 to 3.50). SPCD for each evaluated progestin (MPA, LNG, and MP) correlated positively with BMI (rs 0.30; \( p = 0.02 \)).

Since some patients had used more than one type of progestin during the follow-up, they were analyzed more than once considering the SPCD before and after each progestin applied in the HRT, being 57 TS patients in an overlapped analysis. Some patients using MP nowadays were those who had used either MPA or LNG in the past. They presented similar results in lab tests such as HbA1C, FG, HDL, LDL, and TG. There was no significant change, we observed that TS patients tended to present more hypertension (\( p = 0.071 \)) (Table 2D). Besides, we were able to demonstrate a significant correlation of SPCD with improving HDL level (rs 0.38; \( p = 0.01 \)) and worsening BMI (rs −0.32; \( p = 0.01 \)). SPCD for each evaluated progestin (MPA, LNG, and MP) correlated positively with BMI (rs 0.30; \( p = 0.02 \)).

When analyzing only the TS adult (>19 yo) group (n = 37/61), with a mean age of 25.8 ± 5.8 yo, who had used progestin for 3.6 ± 2.5 years (from 1 to 8.9 y), those using MPA were significantly younger than MP ones, so they were more exposed to the progestin as shown in Table 3. We corrected this bias in body weight change for the time evaluated, dividing weight variation by the years of exposition. We then observed that the mean annual percentage variation on BMI was significantly lower with LNG than those who had used MPA or MP. Weight gain with MPA and MP was similar, as shown in Fig. 2.

The association between the type of progestin and weight change was confirmed by using a multiple linear regression model. For that, we applied the annual percentual variation in BMI as the dependent variable and logistic variables LNG, age, duration of treatment, and cumulated exposition (SPCD) as independent ones. Annual BMI variation \( \frac{\text{BMI}_{\text{after}} - \text{BMI}_{\text{before}}}{\text{years}} \) correlated positively with BMI (rs 0.30; \( p = 0.0144 \)).
Discussion

Our study showed, for the first time, that neither the use of estrogen alone nor ten-day monthly use of progestin replacement, independently of its type, during the first year of HRT, has modified any metabolic parameters in TS. However, we found a significant effect concerning a type of progestin for weight gain when used for a long time. The HRT with LNG showed a significantly less weight gain than with MPA and natural MP, beyond being better for weight maintenance.

According to Trolle et al., most treatment recommendations directed to the TS population are based on expert opinion and not evidence-based. Since TS implies in increased metabolic risk per se, revisiting the long-term HRT with estrogen and progestin, mimicking the female physiological pattern the from pharmacological puberty induction until the age of menopause, it is of great importance to look for a safer sex steroid treatment for them. Considering that estrogenization can improve the metabolic profile and it is not related to weight gain independently of the route of administration, our cohort was not divided per subtype of used estrogen. We found no significant weight alteration during estrogen replacement alone. As shown by Baldin et al., the use of growth hormones also did not change body composition, so our group was also not divided by previous use or not of GH. All TS patients were euthyroid even though some on levothyroxine replacement, which prevented us from separate them in a different group for thyroid status.

## Table 2.
Prevalence of metabolic changes observed in Turner Syndrome patients along with different hormone replacement therapy regimens. A and B, Phase 2.1, HRT before and after estrogen-only; C, Phase 2.2, HRT with estrogen and within one year of progestin. D, Phase 2.3, HRT with estrogen and long-term progestin. HRT, Hormone Replacement Therapy; *time evaluated as the average and minimum-maximum. In parenthesis is represented the number of patients with available data regarding that aspect analyzed.

| Metabolic changes | N (total) | %  | P-value |
|-------------------|----------|----|---------|
| A-Before HRT      |          |    |         |
| Hypertension      | 4 (38)   | 10.5 |         |
| Dysglycemia       | 3 (28)   | 10.7 |         |
| Dyslipidemia      | 8 (23)   | 34.8 |         |
| Overweight/Obesity| 10 (43)  | 23.2 |         |
| B-HRT with estrogen-only | | | |
| Hypertension      | 6 (28)   | 20.7 | 0.222   |
| Dysglycemia       | 2 (13)   | 15.4 | 0.670   |
| Dyslipidemia      | 6 (12)   | 50.0 | 0.383   |
| Overweight/Obesity| 13 (29)  | 44.8 | 0.054   |
| C-HRT with estrogen and within the first year of progestin | | | |
| Hypertension      | 3 (25)   | 12.0 | 0.855   |
| Dysglycemia       | 0 (0)    | 0   |         |
| Dyslipidemia      | 9 (24)   | 37.5 | 0.866   |
| Overweight/Obesity| 12 (29)  | 41.3 | 0.101   |
| D-HRT with estrogen and after long-term of progestin | | | |
| Time evaluated (years)* | 2.67 y (1.00 to 14.08 y) | | |
| Hypertension      | 15 (59)  | 25.4 | 0.071   |
| Dysglycemia       | 6 (59)   | 10.2 | 0.937   |
| Dyslipidemia      | 18 (66)  | 27.3 | 0.495   |
| Overweight/Obesity| 43 (73)  | 58.9 | 0.00019*|

## Table 3.
Body mass index variation in adult Turner Syndrome patients stratified by the types of progestin used in the hormone replacement therapy. Data expressed as mean ± SD for parametric and as median – minimum/maximum for nonparametric data distribution by different types of progestins during the long-retrospective assessment. This subgroup comprises 37/73 TS adult patients enrolled in Phase 2.3. MPA medroxyprogesterone, LNG levonorgestrel. MP micronized progesterone; SPCD standard progestin cumulative dose; BMI body mass index. Differences in mean values among the groups are greater than would be expected by chance (p < 0.05); *MPA vs MP, **LNG vs MP.

| Progestin | N  | Ageyo. | Timeyr. | SPCD | Initial BMIKg/m² | After BMIKg/m² |
|-----------|----|--------|---------|------|-----------------|----------------|
| MPA       | 12 | 22.6 ± 3.6* | 5.5 ± 2.0* | 6.0*(1.0 to 11.9) | 22.9 ± 4.1* | 25.5 ± 5.9 |
| LNG       | 14 | 25.0 ± 6.1 | 3.4 ± 1.9 | 2.8(1.3 to 7.5) | 24.5 ± 4.4** | 24.9 ± 4.3 |
| MP        | 11 | 29.3 ± 5.4 | 1.8 ± 0.7 | 1.6(1.1 to 4.3) | 28.7 ± 4.0 | 29.7 ± 4.2 |
| Total     | 37 | 25.5 ± 5.8 | 3.6 ± 2.5 | 2.6(1.0 to 11.9) | 25.3 ± 4.7 | 26.5 ± 5.2 |
The benefits of TE2 replacement are better than with EE2 or even oral E2, as consolidated by literature²,¹⁷, so we also prefer this subtype and route of E to start puberty with low-dose and gradual increases¹,²,¹¹. Likewise, when possible, and during follow-up, we changed the oral to transdermal route to decrease thromboembolic and stroke risk¹,¹¹,¹⁷. Skin allergy problems or financial constraints are the reasons for those patients still in use of oral E2. We do not advocate for the use of EE2 as HRT in TS patients.

As far as we know, no published work accessed the cyclical replacement of progestin effect over metabolic parameters, neither in normal nor in the Turner population. The main articles related to this topic addressed contraception, breast cancer, polycystic ovary syndrome, menopause women or animal models, situations related to daily progestin use, different from the 10-day per month regimen. TS is an interesting population to be studied concerning HRT effects once they are free from their ovarian hormones due to primary ovarian failure²³. We focused our study on the progestin influence since previous studies showed no difference in metabolic variables with estrogens¹,¹¹.

As progestin is used in HRT only for an average of 10 days per month but for so many years, it is essential to consider that it may have a cumulative effect on the metabolic health, especially in TS women, since it totalizes a huge P exposure time along with the basal risk for metabolic comorbidities. Our study is the first to investigate the metabolic effect of three types of P for HRT in TS. To overcome the bias of overlapping P exposure, we have created a standardizing tool (SPCD) to measure the cumulative exposure to different P. We noted that SPCD was higher for MPA than MP and LNG. We hypothesized that MPA worsens the metabolic risk of TS women based on its cumulative glucocorticoid activity²².

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It is well established in cell cultures that MPA has glucocorticoid properties²⁴. Besides, in ovariectomized female mice, MPA produced a pro-thrombotic effect²⁵. Arias-Loza et al. have also found that MPA in animal model abrogates protective vascular effects of 17β-estradiol because it acts not only on PR but also on AR and GR¹⁵. Otto et al. compared MPA with Drospirenone, a synthetic progesterone hormone, and found that mitogenic activity of MPA on the mammary gland is higher than drospirenone and possibly because of its greater glucocorticoid activity²⁶.

In humans, MPA effects on weight were studied in the context of contraceptive therapy²⁷,²⁸. Yancey and Raleigh found evidence that progestin-only contraceptives cause weight gain with an increase in fat deposition and decreases in lean mass²⁸. Regarding P and changing metabolic profile, Ozdemir et al., studied the effect of MPA in women with polycystic ovary syndrome and found that there were no changes in carbohydrate or lipid metabolism²⁹. Moreover, P has different effects on human breast cancer because of its pharmacological ductal growth properties. Comparing MPA, natural MP, Drospirenone, and Nestorone, Fu et al., concluded that all of them enhanced breast cell migrations and invasion when PR immunostaining is positive (PR⁺)³⁰. They found that the spectrum of action of P over other targets on GR, MR, and AR makes MPA the most potent on the progression of PR⁺ breast cancer³⁰.

Considering the evidence that MPA has higher action over GR than LNG and natural MP³¹, the vast majority of our patients are nowadays on MP or LNG. In our study, we found that the prolonged use of P has an impact on weight gain without worsening blood pressure, lipidic, and glycemic profile. Meanwhile, as the SPCD of the three studied progestins showed a correlation with BMI over time, we proceed to investigate the effect of MPA, LNG, and MP on worsening weight.

During the estrogen and long-use of progestin HRT assessment (phase 2.3), all types of progestins increased BMI. However, no marked difference occurred in the young group, even correcting BMI for z-score, being only evident the BMI change among adult TS patients. Also, MPA impact was more prominent than with LNG and MP. Notably, we also verified that the weight gain was clearly lower with LNG than MPA, as expected due to its less glucocorticoid activity, and surprisingly, TS patients on HRT with MP also gained more weight over time similar to MPA.
Studies assessing LNG and weight gain were all in the field of contraception or menopausal hormone therapy. Modesto et al. compared depot-MPA, LNG-IUD, and TCu380A IUD action on weight and found that all of them increased weight after the first year and after ten years, but LNG and TCu380A devices gained less weight than depot-MPA86. Besides, a systematic review and meta-analysis evaluating thromboembolism found that combined oral contraceptives containing LNG have less risk for venous thromboembolism87, as primarily demonstrated in the Danish cohort study88.

Concerning natural MP, a 3-month study from Casanova and Spritzer did not found that cyclic vaginal MP alter weight or other metabolic variables in the context of menopausal period89. Nevertheless, literature is scarce in MP study related to metabolic parameters, especially in evaluating of TS population. Although our study was carefully conducted, there were some limitations. The lack of baseline clinical and lab data from the beginning of HRT registered in patient’s medical records and in addressing letter from pediatric services to our adult unit reduced a lot our number of samples. Also, the retrospective nature of the study cannot provide a cause-effect relationship and make it difficult to assess the correlation of each P since patients used more than one type of P onwards. We reinforce that the majority of our TS patients had used MPA for some period, and they present an inherent risk for metabolic disorders when compared to the general female population90.

In conclusion, metabolic morbidities are prevalent in TS even before HRT and LNG seems to be a better progestin choice in comparison with MPA and MP. Prolonged HRT with LNG may have a positive impact on metabolic features. Although, LNG did not impair blood pressure, lipid and glycemic profile, and correlated with less weight gain in our study, further research is needed to indicate a recommendation.

Data availability
All raw data are available for further analysis upon request to corresponding author.

References
1. Gravholt, C. H. et al. Clinical practice guidelines for the care of girls and women with Turner syndrome: proceedings from the 2016 Cincinnati International Turner Syndrome Meeting. Eur J Endocrinol 177, G1–G70, doi:10.1530/EJE-17-0430 (2017).
2. Collett-Solberg, P. F. et al. Endocrine diseases, perspectives and care in Turner syndrome. Arq. Bras. Endocrinol. Metab. 55, 550–558 (2011).
3. Gonzalez, L. & Witchel, S. F. The patient with Turner syndrome: puberty and medical management concerns. Fertil. Steril. 98, 780–786, https://doi.org/10.1016/j.fertnstert.2012.07.1104 (2012).
4. Gravholt, C. H. Epidemiological, endocrine and metabolic features in Turner syndrome. Arq. Bras. Endocrinol. Metab. 49, 145–156, https://doi.org/10.1590/s0004-27302005000100019 (2005).
5. Hankus, M. et al. Prediction of Spontaneous Puberty in Turner Syndrome Based on Mid-Childhood Gonadotropin Concentrations, Karyotype, and Ovary Visualization: A Longitudinal Study. Horm. Res. Paediatr. 89, 90–97, https://doi.org/10.1159/000483521 (2018).
6. Giordano, R. et al. Metabolic and cardiovascular outcomes in a group of adult patients with Turner’s syndrome under hormonal replacement therapy. Eur. J. Endocrinol. 164, 819–826, https://doi.org/10.1530/EJE-11-0002 (2011).
7. Gravholt, C. H., Poulsen, H. E., Ott, P., Christiansen, J. S. & Vilstrup, H. Quantitative liver functions in Turner syndrome with and without hormone replacement therapy. Eur. J. Endocrinol. 156, 679–686, https://doi.org/10.1530/EJE-07-0070 (2007).
8. Lichardopol, C. & Mota, M. Hypothyroidism in Turner syndrome. Rom. J. Intern. Med. 45, 177–182 (2007).
9. Baldin, A. D. et al. Growth hormone effect on body composition in Turner syndrome. Endocrine 40, 486–491, https://doi.org/10.1007/s12020-011-9504-z (2011).
10. Gawlik, A. & Malecka-Tendera, E. Transitions in endocrinology: treatment of Turner’s syndrome during transition. Eur. J. Endocrinol. 170, R57–74, https://doi.org/10.1530/EJE-13-0900 (2014).
11. Klein, K. O. et al. Estrogen Replacement in Turner Syndrome: Literature Review and Practical Considerations. J. Clin. Endocrinol. Metab. 103, 1790–1803, https://doi.org/10.1210/jc.2017-02183 (2018).
12. Vickery, Z. et al. Weight change at 12 months in users of three progestin-only contraceptive methods. Contraception 88, 503–508, https://doi.org/10.1016/j.contraception.2013.03.004 (2013).
13. Lopez, L. M. et al. Progestin-only contraceptives: effects on weight. Cochrane Database Syst. Rev, CD008815, https://doi.org/10.1002/14651858.CD008815.pub3 (2013).
14. Schindler, A. E. et al. Classification and pharmacology of progestins. Maturitas 61, 171–180, https://doi.org/10.1016/j.maturitas.2008.11.013 (2008).
15. Arias-Loza, P. et al. Medroxyprogesterone acetate but not drospirenone ablates the protective function of 17 beta-estradiol in aldosterone salt-treated rats. Hypertension 48, 994–1001, https://doi.org/10.1161/111 HYP00042482.57186.e8 (2006).
16. Trolle, C., Mortensen, K. H., Hjerrild, R. E., Cleemann, L. & Gravholt, C. H. Clinical care of adult Turner syndrome–new aspects. Arq. Bras. Endocrinol. Metab. 98, 2716–2724, https://doi.org/10.1590/s0004-27302013000400005 (2013).
17. Rebholz, Y. et al. The Natural History of Metabolic Comorbidities in Turner Syndrome from Childhood to Early Adulthood: Comparison between 45,X Monosomy and Other Karyotypes. Front. Endocrinol. 9, 27, https://doi.org/10.3389/fendo.2018.00027 (2018).
18. Roulot, D. Liver involvement in Turner syndrome. Liver Int. 33, 24–30, https://doi.org/10.1111/liv.12007 (2013).
19. Minzioni, G., Poulakos, P., Tsiamitis, C. & Goulis, D. G. Hypogonadism and non-alcoholic fatty liver disease. Minerva Endocrinol. 42, 145–150, https://doi.org/10.23736/S0391-1977.16.02570-0 (2017).
20. Reinehr, T. et al. Weight gain in Turner Syndrome: association to puberty induction? - longitudinal analysis of KIGS data. Clin. Endocrinol. 85, 85–91, https://doi.org/10.1111/cen.13044 (2016).
21. Taboada, M. et al. Pharmacokinetics and pharmacodynamics of oral and transdermal 17beta estradiol in girls with Turner syndrome. J. Clin. Endocrinol. Metab. 96, 3502–3510, https://doi.org/10.1210/jc.2011-1449 (2011).
22. Sweeney, E. E., Fan, P. & Jordan, V. C. Molecular modulation of estrogen-induced apoptosis by synthetic progestins in hormone replacement therapy: an insight into the women’s health initiative study. Cancer Res. 74, 7060–7068, https://doi.org/10.1158/0008-5472.CAN-14-1784 (2014).
25. Freudenberger, T. et al. Synthetic gestagens exert differential effects on arterial thrombosis and aortic gene expression in ovariecetomized apolipoprotein E-deficient mice. Br. J. Pharmacol. 171, 5032–5048, https://doi.org/10.1111/bph.12814 (2014).
26. Otto, C. et al. Comparative analysis of the uterine and mammary gland effects of drospirenone and medroxyprogesterone acetate. Endocrinology 149, 3952–3959, https://doi.org/10.1210/en.2007-1612 (2008).
27. Lopez, L. M. et al. Progestin-only contraceptives: effects on weight. Cochrane Database Syst Rev, CD008815, https://doi.org/10.1002/14651858.CD008815.pub4 (2016).
28. Yancey, J. R. & Raleigh, M. Progestin-only contraceptives: effects on weight. Am. Fam. Physician 89, 715–716 (2014).
29. Ozdemir, S., Gorkemli, H., Gezginc, K., Ozdemir, M. & Kiyici, A. Clinical and metabolic effects of medroxyprogesterone acetate and ethinyl estradiol plus drospirenone in women with polycystic ovary syndrome. Int. J. Gynaecol. Obstet. 103, 44–49, https://doi.org/10.1016/j.ijigo.2008.05.017 (2008).
30. Fu, X. D. et al. Comparative actions of progesterone, medroxyprogesterone acetate, drospirenone and nestorone on breast cancer cell migration and invasion. BMC Cancer 8, 166, https://doi.org/10.1186/1471-2407-8-166 (2008).
31. Modesto, W., de Nazare Silva dos Santos, P., Correia, V. M., Borges, L. & Bahamondes, L. Weight variation in users of depot-medroxyprogesterone acetate, the levonorgestrel-releasing intrauterine system and a copper intrauterine device for up to ten years of use. Eur. J. Contracept. Reprod. Health Care 20, 57–63, https://doi.org/10.1016/j.ejchc.2014.09.006 (2015).
32. Dragoman, M. V. et al. A systematic review and meta-analysis of venous thrombosis risk among users of combined oral contraception. Int. J. Gynaecol. Obstet. 141, 287–294, https://doi.org/10.1002/ijgo.12455 (2018).
33. Lidegaard, O., Nielsen, L. H., Skovlund, C. W., Skjeldestad, F. E. & Lokkegaard, E. Risk of venous thromboembolism from use of oral contraceptives containing different progestogens and oestrogen doses: Danish cohort study, 2001-9. BMJ 343, d6423, https://doi.org/10.1136/bmj.d6423 (2011).
34. Casanova, G. & Spritzer, P. M. Effects of micronized progesterone added to non-oral estradiol on lipids and cardiovascular risk factors in early postmenopause: a clinical trial. Lipids Health Dis. 11, 133, https://doi.org/10.1186/1476-511X-11-133 (2012).
35. Schoemaker, M. J. et al. Mortality in women with Turner syndrome in Great Britain: a national cohort study. J. Clin. Endocrinol. Metab. 93, 4735–4742, https://doi.org/10.1210/jc.2008-1049 (2008).

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Author contributions
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