INTRODUCTION

The human X and Y chromosomes have evolved from a pair of ancestral chromosomes. The X chromosome has retained many properties of an autosome, containing 2218 genes which is one of the lowest gene density in the human genome (Homo sapiens GRCh38.p14). Thus, it is an interesting subject as it has a wide spectrum of clinical manifestations. 49,XXXXY syndrome is a rare sex chromosome polysomy with an approximate incidence of 1 in 85,000 male births. In 1960, Fraccaro described this clinical entity. Fraccaro aneuploidy maybe emanates from the nondisjunction of the X chromosome during the meiosis division. These consecutive nondisjunctions will produce an egg with four X chromosomes when fertilized by a Y generation sperm, results in an embryo with 49,XXXXY syndrome. Interestingly, the appearance of this syndrome does not depend on the age of the mother. The chromosomal aberration found in Klinefelter syndrome is due to either meiotic or mitotic nondisjunction, leading to a sex-chromosomal aneuploidy. Two genetic variations exist in Klinefelter syndrome. The majority of cases more than 90% would be presented as a pure form with a 47,XXY karyotype, whereas the remaining 10% include the following sex-chromosomal abnormalities: Mosaic karyotypes such as 46,XY/47,XXY, higher-grade aneuploidy such as 48,XXXXY; 49,XXXXY, and structurally abnormal X chromosomes. Klinefelter syndrome might be detected during the prenatal, prepubertal, adolescent, or adult period. The physical manifestations of Fraccaro syndrome are often variable; Clinical features of Fraccaro syndrome are hypergonadotropic hypogonadism; moreover, the risk of testicular tumorigenesis and testicular degenerative changes mental retardation and radioulnar synostosis. In this study, we report clinical presentation of a 19-year-old male patient with Fraccaro syndrome. We also compared all 49,XXXXY reported patients among the Iranian population. This study provides new insights into the association between maternal age at birth time and the occurrence of Fraccaro syndrome.
2 | CASE REPORT

A 19-year-old male patient was referred to the Department of Genetics, Razi Pathobiology & Genetic center for karyotyping. He has been the second child of healthy unrelated parents. His mother was 21 years old and his father was 25 years old. There was no evidence of intellectual disability or mental illness in the family. His older sister was a healthy individual with normal development and she has also two normal children.

The patient’s clinical examinations revealed several congenital abnormalities such as specific bone malformation, microcephaly, dental issues, muscular hypotonia, gynecomastia, tall stature disorder, small hands, low nasal bridge, low set ears, unilateral dysplasia of the hip, and azoospermia (Figure 1). The patient had suffered from hypogonadism, micropenis, and masturbation. He also had some learning disabilities, speech impediment which increased the risk of low-quality friendships and social difficulties in our patient with his peers. In our case, some specific symptoms were detected which includes mild anemia. He also suffered from recurrent bacterial infections during his growth.

3 | MATERIALS AND METHODS

Peripheral lymphocytes were cultured and GTG-banded using standard techniques and 50 metaphases were analyzed.

A multiplex quantitative fluorescent polymerase chain reaction (QF-PCR) was performed using Devyser Compact v3 kit according to the manufacturer’s protocol. DNA was extracted from blood samples of the patient using QIAamp DSP DNA Blood Mini kit (QIAGEN). The QF-PCR products were analyzed by capillary electrophoresis on an ABI 3500 automated DNA Sequencer.

Fluorescence in situ hybridization (FISH) analysis of 100 interphase nuclei, using Cytocell probes for chromosomes X, Y and 18 indicated a line proportion for 49,XXXXY (Figure 3). Analysis of the karyotyping and FISH results were performed using GeneASIS 7.2 software (Applied Spectral Imaging) and reported based on the International System for Human Cytogenetic Nomenclature (ISCN)-2013.

The levels of follicle stimulating hormone (FSH), luteinizing hormone (LH), and total testosterone were measured in the 49,XXXXY patient as he referred to the laboratory, without testosterone therapy. The serum level of LH and FSH were assayed by the LIAISON kits (Diasorin-Saluggia) and testosterone was also assayed by the Elecsys TESTO II (Roche Diagnostics Gmbl) kits.

To investigate the reported males with Fraccaro syndrome among the Iranian population, we systematically reviewed PubMed, Google scholar, and magiran databases to find related articles in both English and Persian languages, on April 20, 2022. To collect these articles, “Advance search” was used in the PubMed database by applying keywords “Fraccaro Syndrome “and” 49,XXXXY” in (Title/Abstract) combined with “Iran” in (Affiliation). To compile Persian articles, we checked magiran database (http://www.magiran.com) by applying “Fraccaro Syndrome” and “49,XXXXY” as a keyword in (search). Furthermore, Fraccaro Syndrome” and
“49,XXXXY” were applied in Google scholar as search terms.

4 | RESULT

In the karyotype study, all metaphases showed a 49,XXXXY syndrome (Figure 2). Investigation of the chromosomal status of the patient’s father indicated a normal 46,XY karyotype and it was also the case for his mother 46,XX karyotype. The finding of this karyotype test was also confirmed by both FISH (Figure 3) and QF-PCR techniques (Figure 4). All short tandem repeat markers for chromosomes 13, 18, and 21 (autosomal) were observed in a normal diallelic pattern. For chromosomes X and Y copy number, the amplification of the homologous gene AMELX/Y indicated a diallelic pattern with an unusual ratio between fluorescent peak areas of 4:1 (Figure 4).

Although the karyotype as a gold standard method, revealed 49,XXXXY in our case, the FISH technique was applied to detect possible low-level X chromosome mosaicism.11 100 metaphases were studied, and no mosaicism was detected.

The X-specific peak of the AMELXY is in a ratio of 4:1 if compared with the Y-specific peak. Electrophoretograms of the QF-PCR products show four microsatellite markers on chromosomes X (DXS2390, DXS1187, DXYS267, DXYS218, and XHPRT) of the patient. AMELXY and ZFYX (on chromosome X and Y) and SRY (only on chromosome Y) region were used for the detection and comparison of the number of X and Y chromosomes. The box under each peak included molecular size (bp) and the area of the peak. Ratio of the peak areas of AMELXY and DXYS218 had been approximately 4:1 in the patient and indicated the presence of four X and one Y chromosomes.

Results of the serum LH, FSH and testosterone levels had been summarized in Table 1. Additionally, Hemoglobin level was measured 10.6 g/dL; Hematocrit examination revealed dimorphic anemia with lymphocytosis and thrombocytosis. Clinical investigations including serum cholesterol, fasting blood sugar, urine analysis, blood urea nitrogen, ferritin, 25-OH-vitamin D, creatinine,
calcium, phosphorus, potassium, sodium serum, and alkaline phosphatase levels were measured and all indicated the normal range.

According to the search strategy mentioned in the method section, two articles in Persian (Magiran) and two articles in English (PubMed) were eventually collected and reviewed. Finally, five male patients with Fraccaro syndrome were compiled among the Iranian population (Table 2). This population has been identified in different regions throughout the country that involved cases in Isfahan, Mazandaran, Tehran, and Hamadan provinces.

### DISCUSSION

49,XXXXY is the result of X chromosome mis-segregation in both meiosis I and meiosis II. Fraccaro’s syndrome is caused by 49,XXXXY chromosomal aneuploidy and is often classified as a Klinefelter’s syndrome variant. Two prevailing theories have been made to account for the phenotype associated with a 49,XXXXY genotype as well as for the other X chromosome aneuploidies. First, an increased dosages of active genes in regions that escape X inactivation, and second, asynchronous replication of the extra X chromosomes.16

The 49,XXXXY syndrome is a very rare but a distinct clinical entity. Typical clinical symptoms include hypogonadism, mental retardation with severe learning difficulties, craniofacial and skeletal abnormalities.17–19 Along with classical clinical features of Fraccaro syndrome, our case further presented mild anemia, bacterial infections, and sexual masturbation behavior.

The hemoglobin level was in the low range in our patient (10.6 g/dl), but his father had also suffered from a mild anemia. Therefore, anemia is not apparently a clinical symptom of 49,XXXXY syndrome in this case; However, it is worth to mention that a low level of Hemoglobin (10.2 g/dl) had been reported previously in a patient with Fraccaro syndrome.20 We could not find any research which addressed the association between hemoglobin level and Fraccaro syndrome. Therefore, this would be an issue for future studies.
In the present study, hormone levels (LH, FSH, and testosterone) were in the abnormal range. This is also in accord with Wei and et al. (2019) findings who obtained the same results.

In Table 2, all would be 49,XXXXY syndrome reported so far in Iran had already been compiled. As far as we know, this study is the first one in which this genetic abnormality type in Iranian population has been compiled. Although the high maternal age would increase the risk of childbirth with chromosomal abnormality and nondisjunction during meiosis, our finding is somewhat different and is not completely in accordance with the previous results. In fact, maternal age in 3 out of 5 mothers was lower than 30 years. Indeed, our finding is in line with those previous studies which indicated that there is no association between maternal age and Fraccaro syndrome. Moreover, Lia and colleagues that reported that 49,XXXXY is not associated with maternal age. Therefore, it is an open question and further studies need to be undertaken in a larger population.

Furthermore, the symptoms including behavioral issues, tall stature with long legs and arms, speech impediments, kidney anomalies, hypogonadism, specific bone malformation and micropenis in an undiagnosed male should be followed by a karyotype analysis and a subsequent appropriate treatment for adequate growth and pubertal development.

### Table 2

| Reported year | Age of diagnosis | Maternal age | Phenotypic and pathological summary | City          | Reference |
|---------------|------------------|--------------|-------------------------------------|---------------|-----------|
| 2010          | 50 days          | 24           | Hypospadiasis, hypotonia, hypertelorism, atrial septal defect, patient ductus arteriosus, speech impediment | Isfahan       | 12        |
| 2012          | 11 month         | 36           | Hypogonadism, hypo plastic scrotum, hypothyroidism, hypotonia, low set ears, hypertelorism | Babol         | 13        |
| 2013          | 45 years         | NA           | Mild mental retardation and infertility | Tehran       | 14        |
| 2013          | 10 month         | 32           | Hypotonia, microcephaly, low set ears, micrognathia and congenital heart disease, kyphoscoliosis, clinodactyly of the fourth and fifth fingers of both hands, bilateral clubfoot and unilateral dysplasia of the hip | Tehran       | 14        |
| 2015          | 2 month          | 26           | Intrauterine growth restriction, low birth weight, facial dysmorphism, Clinodactyly in feet, microphallus and right undescended testis | Hamadan      | 15        |
| Current study | 19 years         | 21           | Hypogonadism, specific bone malformation, microcephaly, dental issues, Muscular hypotonia, behavioral issues, gynecomastia, tall stature, small hands, speech impediment, azoospermia | Babol        | -         |

Abbreviation: NA, not available.

### Author Contributions

MR performed blood karyotype and participate in manuscript writing. NT performed QF test and analysis. OJ supervised the research, collect the clinical features of patient, and wrote the manuscript.

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### Conflict of Interest

The authors declare no conflicts of interest.

### Data Availability Statement

Data are available from the corresponding author upon reasonable request.

### Consent

Written informed consent was previously obtained from the patient to publish this report in accordance with the journal’s patient consent policy.

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REFERENCES

1. Ross MT, Graffham D, Coffey A, et al. The DNA sequence of the human X chromosome. Nature. 2005;434(7031):325-337.
2. Bearelly P, Oates R. Recent advances in managing and understanding Klinefelter syndrome. FI000Res. 2019;8(112):112.
3. Kleczkowska A, Fryns JP, Berghe H. X-chromosome polysomy in the male. Hum Genet. 1988;80:16-22.
4. Fraccaro M, Kaijser K, Lindsten GJA. Child with 49 chromosomes. Lancet. 1960;2:899-902.
5. Mazzilli R, Delfino M, Elia J, et al. Testosterone replacement in 49,XXXXY syndrome: andrological, metabolic and neurological aspects. Endocrinol Diabetes Metab Case Rep. 2016;2016:150114.
6. Pascanu I, Butila Todoran AM, Csef K, Banescu C, Toganel R. A case of 49,XXXXY syndrome in endocrine practice. Acta Endocrinol. 2008;4:455-464.
7. Bearelly P, Oates R. Evaluation and Management of Klinefelter Syndrome. FI000Res. 2019;8. doi:10.12688/f1000research.16747.1
8. Burgemeister AL, Daumiller E, Bois G, et al. Clinical report of 8 patients with 49,XXXXY syndrome: delineation of the facial gestalt and depiction of the clinical spectrum. Eur J Med Genet. 2019;62(3):210-216.
9. Maqdasy S, Bogenmann L, Batisse-lignonier M, et al. Testosterone replacement in 49,XXXXY syndrome: andrological, metabolic and neurological aspects. Endocrinol Diabetes Metab Case Rep. 2016;2016:150114.
10. Linden MG, Bender BG, Robinson A. Sex chromosome tetrasomy and pentasomy. Pediatrics. 1995;96:672-682.
11. Lakhil B, Braham R, Berguiquia R, et al. Cytogenetic analyses of premature ovarian failure using karyotyping and interphase fluorescence in situ hybridization (FISH) in a group of 1000 patients. Clin Genet. 2010;78:181-185.
12. Hashemi Dehkordi E, Salek M, Hashemipour M, Moaddab MH. A rare case with sex developmental disorder. J Isfahan Med Sch. 2010;28(104):57-62. [In Persian].
13. Alijanpour M, Hadipour A, Taghavi M. A rare case with 49,XXXXY syndrome. J Babol Univ Med Sci. 2012;14(6):102-106. [In Persian].
14. Hadipour F, Shafeghati Y, Bagherizadeh E, Behjati F, Hadipour Z. Fraccaro syndrome: report of two Iranian cases: an infant and an adult in a family. Acta Med Iran. 2013;51:12.
15. Etemadi K, Basir B, Ghahremani S. Neonatal diagnosis of 49,XXXXY syndrome. Iran J Reprod Med. 2015;13:181-184.
16. Hou JW. 49,XXXXY syndrome. Chang Gung Med J. 2004;27:551-554.
17. Tartaglia N, Ayari N, Howell S, D’Epagnier C, Zeitler P. 48,XXYY, 48,XXXY and 49,XXXXY syndromes: not just variants of Klinefelter syndrome. Acta Paediatr. 2011;100(6):851-860.
18. Naotunna N, Liyanage C, Atapattu N. An infant with 49,XXXXY syndrome: a case report. J Med Case Reports. 2021;15.6:30.
19. Hammami MB, Elkhapery A. Sexual and developmental aspects of 49,XXXXY syndrome: a case report. Andrologia. 2020;52(10):e13771.
20. Nerli R, Magdum P, Mungawadi A, Ghagane S, Hiremath M. 49,XXXXY syndrome: an infant presenting with ambiguous genitalia. Indian J Health Sci Biomed Res. 2016;9:342-344.
21. Celik A, Eraslan S, Gökçöğüz N, et al. Identification of the parental origin of polysomy in two 49,XXXXY cases. Clin Genet. 1997;51:426-429.
22. Peet J, Weaver DD, Vance GH. 49,XXXXY: a distinct phenotype. Three new cases and review. J Med Genet. 1998;35:420-424.
23. Lia EN, Otero SAM, Ferraz M, Gonçalves LPV. Oral aspects of 49,XXXXY syndrome: a case report. J Dent Child (Chic). 2007;74(2):136-139.
24. Peitsidis P, Manolakos E, Peitsidou A, et al. Pentasomy 49,XXXXY diagnosed in utero: case report and systematic review of antenatal findings. Fetal Diagn Ther. 2009;26:1-5.

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