INTRODUCTION

Liaoning Province is located in the North East of the People’s Republic of China. In Chinese, it is known as “the Golden Triangle” because of its strategic location. Liaoning Province was established around 1907 with the name “Fengtian” and later in 1929, it was changed to Liaoning (Rhoads, 2000). The estimated population size is approximately 43.91 million with Han (83.94%) as the dominating ethnic group followed by minorities such as...
as Manchus (12.88%), Mongols (1.60%), Hui (0.632%), Koreans (0.576%), and Xibe (0.317%; www.stats.gov.cn). With the advancement in forensic genetics, short tandem repeats (STRs) are used for investigations such as rape cases, paternity testing, Kinship analysis, familial search, and missing person investigations because these genetic markers are highly informative due to their polymorphic nature (Adnan et al., 2017; Xing et al., 2019; Zhan et al., 2018). A 5-dye GoldenEye™ 20A kit (Beijing PeopleSpot Inc), which contains 13 combined DNA index system core STR loci and six additional STRs (PentaE, PentaD, D2S1338, D19S433, D12S391, and D6S1043) which can be amplified simultaneously (Huang et al., 2013). However, forensic statistical parameters, allelic frequencies, phylogenetic relationship, and population structure still needs to be investigated on an extended set of marker such as Goldeneye 20A for Liaoning Han population in comparison with local and worldwide populations.

1.1 | Sample

Blood Samples were collected from a total of 1138 unrelated Han individuals (667 males and 471 females) residing in the Liaoning province of people’s republic of china at least from three generations with written informed consent. This project was approved by the institutional review boards of China Medical University, Shenyang, PR China. ReliaPrep™ Blood gDNA Miniprep System (Promega) was used to isolate the DNA, and Nanodrop-2000 (Thermo Fisher Scientific) was used to quantify the DNA concentration according to the manufacturer’s instructions. Later on, DNA has diluted accordingly to approximately 1 ng/μl.

1.2 | PCR amplification and genotyping

PCR co-amplification of 20 autosomal STR loci (D19S433, D5S818, D21S11, D18S51, D6S1043, D3S1358, D13S317, D7S820, D16S539, CSF1PO, Penta D, vWA, D8S1179, TPOX, Penta E, TH01, D12S391, D2S1338, FGA, and Amelogenin) were performed in a fluorescence-based multiplex reaction using the Goldeneye® 20A STR Kit. About 1 to 2 ng of the template DNA was used to amplify according to the manufacturer’s instructions in GeneAmp PCR System 9700 thermal cycler (Applied Biosystems). The PCR products were analyzed using an eight capillary ABI 3500 DNA Genetic Analyzer with the POP-4 TM polymer (Life Technologies). GeneMapper Software version 4.0 (Life Technologies) was used for genotype assignment. Allelic nomenclatures were based according to the allelic ladder provided by GoldenEye™ 20A kit (Beijing PeopleSpot Inc) in accordance with ISFG recommendations [9, 10].

1.3 | Statistical analysis

Allele frequencies and other important forensic statistical parameters and principal components analysis (PCA) were computed using the STRAF (Gouy & Zieger, 2017). Hardy-Weinberg equilibrium (HWE) and linkage disequilibrium (LD) and observed heterozygosity (Ho) were estimated using Arlequin 3.5 (Excoffier et al., 2007). Pairwise Fst genetic distance between Liaoning Han and the other seven populations (Ye et al., 2017; Zhan et al., 2018) was calculated using the STRAF. Nei genetic distances between Liaoning Han and other 71 Chinese reference populations and 81 worldwide populations were estimated according to the Phylip3.695 package. The phylogenetic tree was visualized with Mega7 software (Kumar et al., 2016). The STRUCTURE v.2.3.4 software (Falush et al., 2003) was used to calculate the ancestry component. The model-based analysis employed the length of burnin period of 100,000 and Markov Chain Monte Carlo (MCMC) step of 100,000 under the “independent allele frequencies” and “LOCPRIOR” models with the k values ranging from 2 to 10 with five repeats each run.

2 | RESULTS AND DISCUSSION

We have successfully generated genotype profiles of 20 autosomal STRs using the GoldenEye™ 20A kit (Beijing PeopleSpot Inc) and the allelic frequencies of these STRs are summarized in Table S1. A total of 253 alleles were observed while PentaE was the most polymorphic with 25 alleles while TH01 was the least polymorphic with six alleles. Allelic frequencies ranged from 0.00043 to 0.5369 while gene diversity (GD) values ranged from 0.9198 (PentaE) to 0.6080 (TPOX). Polymorphism information content (PIC) also followed the above trend where PentaE (0.9137) showed the highest value while TPOX (0.5452) showed the lowest value whereas the typical paternity index (TPI) diversified from 1.2757 (TPOX) to 5.5784 (PentaE). Probability of match (PM), power of discrimination (PD) and power of exclusion (PE) ranged from 0.0130 (PentaE) to 0.2180 (TPOX), 0.7819 (TPOX) to 0.9869 (PentaE) and 0.3006 (TPOX) to 0.8166 (PentaE), respectively. Observed heterozygosity values ranged from 0.6080 (TPOX) to 0.9103 (PentaE). Detailed forensic parameters are summarized in Table 1. The combined discrimination power (CPD) was 99.999999999999997% and the combined exclusion power (CPE) was 99.99998231%. The result of these forensic parameters showed that the Goldeneye 20A kit is suitable for forensic investigations such as paternity testing, personal identification, and familial search in the Liaoning Han population. Most of the loci were in Hardy-Weinberg Equilibrium (HWE) except for Penta E, vWA, and D21S11. Consequently, when we applied sequential Bonferroni
correction (Benjamini & Hochberg, 1995) to mitigate against the so-called “multiple comparison problem,” only one locus (D21S11) was found to be out of HWE (Table S2). Various circumstances can result in deviation from HWE such as systemic factors that consist of migration, mutation, and natural selection, while dispersive factors result from genetic drift by the force of chance and sample size of a population. Linkage disequilibrium (LD) implies an association between the qualitative random variables corresponding to alleles at different STRs. Measuring the levels of linkage disequilibrium is important for gene mapping and it helps in the understanding of genome structure (Chen et al., 2006). Exact tests for linkage equilibrium (LE) showed that the $p$-values of 25 pairs of STR loci were below 0.05, and thus displaying LD, which is summarized in Table S3. After a sequential Bonferroni correction (Benjamini & Hochberg, 1995), only five pairs were out of LE which were (vWA/D18S51), (FGA/D12S391), (D8S1179/Penta D), (Penta E/D6S1043), and (D3S1358/D6S1043). This Linkage disequilibrium (LD) may be the result of the association between adjacent alleles co-inherited.
from single, ancestral chromosomes but may also be a result of selection, random genetic drift, the rate of mutation or recombination, nonrandom mating, founder effects, sampling effects, recent admixture, and population substructure (Chakravarti, 1999).

To check the hierarchy existence of the population in Liaoning Han and seven other populations (Uzbek, Kyrgyz, Manchu, Mongol, Tibetan, Ili Kazakhs, and Yanbai Korean) with raw genotypic data, we explored the genetic heterozygosity or homozygosity via principal component analysis (PCA).
A total of 3.01% genetic variations can be extracted by the first three PCs (Figure 1). Above mentioned PCA results are later confirmed with pairwise $F_{st}$ genetic distances (Table S4) and phylogenetic relationship reconstruction (Figure S1). We set k-values varying from 2 to 10 to get information on ancestry related to the Liaoning Han population (Figure 2). As shown in Figure 2, we identify the best optimal predefined populations in five ($K = 5$). Liaoning Han shared their genetic component with Manchu and Korean (yellow component) while the other four populations (Kazakh, Uzbek, Kyrgyz, and Mongolian) make a separate component (green component). We can also identify a common blue component existing in all of these populations with different proportions (blue and red component). In total, two genetical clusters were observed: one comprises Liaoning Han Chinese, Manchu, Jilin Korean, and Tibetans, which are typical East Asian populations with the dominant East Asian ancestry components; while the other one consists of Kazakh, Kyrgyz, Uzbek and Mongolian populations, which are descendants of ancient Altai-speaking populations residing in central and north Asia. Moreover, to check the genetic affinities of the Liaoning Han population we have compared with 71 local Chinese populations and 80 worldwide populations by calculating Nei’s genetic distances. According to Nei’s genetic distance (Table S5), among the Chinese populations Han population from Hubei (0.0021) showed the closest genetic distance and followed by the Han population from Yunnan (0.0022) while the Kyrgyz population from Xinjiang (0.4569) showed greatest genetic distance followed by Manchu population from Xinjiang (0.4010) and Uzbek population from Xinjiang (0.3382) among the studied populations. In neighbor-joining (NJ) Liaoning Han population formed a close cluster with other Han populations from the Northeast, East, and Southeast (Figure 3). In worldwide populations, the Korean population from Jilin (0.0064) showed the closest affinity, followed by the Han population
from central China (0.0287) while distant genetic distance showed the Mongol population from Mongolia (1.2226) followed by the Kadažan population (0.6203) from Malaysia and amaXhosa population (0.2638) from South Africa (Table S6). In neighbor-joining (NJ) Liaoning Han population formed a close cluster with local Chinese populations (Figure 4) such as Han and Koreans. A heat map of this genetic matrix showed that the Hans, Korean, and Japanese populations showed higher genetic similarities (Figure S2). Finally, in a previous study based on large-scale whole-genome variations in which researchers have collected 11,670 Han Chinese individuals samples across China to explore the Han Chinese population genetic structure (Chiang et al., 2018). Their finding suggested that there are differences among east to the west among Han Chinese which were not explored in depth previously and these results are in accordance with the north-south differentiation among Han Chinese previously suggested by Xu et al., (2009). Findings of our results based on STR variations also suggest the variations among north-south genetic structure variations but we didn't find any variations among east to west genetic structure changes and this may be the result of sample coverage.

2.1 Comments

Overall, our study demonstrates that the GoldenEye™ 20A kit (Beijing PeopleSpot Inc) showed a higher level of genetic diversity in the Liaoning Han population. These STRs which are included in the GoldenEye™ 20A kit can be used for forensic applications along with population genetic studies. Population genetic analysis showed that the Han population has significant genetic variations when compared with other Chinese minorities such as Uzbek, Kyrgyz, Kazakh, Uyghur, and Manchu populations of Xinjiang. The present study provides a precise reference database of Liaoning Han population on extended set of autosomal STRs for forensic applications and population genetic studies.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

AUTHOR CONTRIBUTION

A.A. and J.D. designed this study. J.D., A.A., and Y.D. collected the samples. J.D., A.A., and Y.D. performed experimental work. J.D., A.A., Y.D., A.R., F.A., and M.A. analyzed the results. A.A. wrote and modified the manuscript. All authors reviewed the manuscript.

COMPLIANCE WITH ETHICAL STANDARDS

All participants gave their informed consent in writing after the study aims and procedures were carefully explained to them in their language. The study was approved by the ethical review board of China Medical University, Shenyang Liaoning Province, People's Republic of China, and in accordance with the standards of the Declaration of Helsinki.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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