INTRODUCTION

Shanghai, located on the eastern coast, was listed as the largest city in China by population according to the 2010 population census. There are more than 26 million citizens in Shanghai as of 2019, and approximately 98.8% of the residents are of Han Chinese ethnicity. Approximately 9 million Shanghai residents are long-term migrants from Anhui (29.0%), Jiangsu (16.8%), and Henan (8.7%), among other regions (https://en.wikipedia.org/wiki/Shanghai). With a history of more than 2,000 years, Shanghai was a small fishing village until it began to serve as a major trading port during the Tang Dynasty. Before the liberation, Shanghai received a large Han population who migrated from the central plains due to the war. Since the founding of the People’s Republic of China, Shanghai has attracted a large number of migrants.
from diverse regions of China due to its rapid economic development. Mandarin is widely used as the official language in Shanghai, and the Shanghai dialect, a fusion of dialects brought by immigrants from the surrounding areas, is also widely used in Shanghai. Gene flow and exchange lead to continuous genetic admixture in Shanghai with the frequent inflow of populations. It is thus worthwhile to obtain an overview of the features of genetic polymorphism in the Han population in Shanghai.

Short tandem repeats (STRs) are DNA segments that consist of 2–100 nucleotides repeated in tandem. These repetitive elements make up approximately 3% of the human genome, and most of them are located in noncoding regions. STRs have relatively high estimated mutation rates ($10^{-3}$) (Brinkmann, Klintschar, Neuhuber, Huhne, & Rolf, 1998) compared with SNPs ($10^{-8}$) (Nachman & Crowell, 2000). They experience changes in their number of repeat units every 1,000–10,000 generations. This elevated capacity to change makes them powerful agents of genetic diversity. From an evolutionary perspective, the dynamic expansion and the mutator properties of STRs make them powerful generators of genetic variability for potential evolutionary change. In other words, they may provide latent selective advantages in short periods of time as opposed to geological time. In addition, due to the accumulation of length mutations during replication, highly polymorphic STRs are considered to be rapidly evolving sequences. On the other hand, as informative neutral markers, STRs are well suited for individual identification and could provide improved resolution in population genetic studies. Therefore, STRs have been adopted for routine forensic applications.

Short tandem repeats can be evaluated by multiplex PCR, which requires only a small amount of DNA. Allele sizing can be achieved with fluorescent primers and an automatic sequencer, providing highly reliable results. Forensic laboratories commonly use tetranucleotide repeats containing a four-base pair repeat structure, such as GATA. In 1997, the FBI Laboratory selected 13 STR loci (CODIS STRs) that form the backbone of the U.S. national DNA database. Many of these same STR loci are used by other countries around the world. Population data on CODIS STRs in the Shanghai_Han population have been reported and updated for routine forensic applications, such as personal identification and paternity testing. However, extra STRs are needed, since joint analysis of non-CODIS STRs and CODIS STRs has been reported to be significantly informative in identifying problematic kinship cases when a definite conclusion could not be obtained with only the profile of CODIS STRs (Asamura, Fujimori, Ota, & Fukushima, 2007; Rodovalho et al., 2015). Commercial STR kits enable consistency in marker use and allele nomenclature among laboratories and help improve quality control. In this study, the genetic polymorphism of 21 non-CODIS STR loci included in the AGCU® Expressmarker 21+1 kit were investigated in 255 unrelated individuals from a Shanghai_Han population. Phylogenetic analyses were also performed to assess the genetic structure among different populations from mainland China.

2 | MATERIALS AND METHODS

2.1 | Sample collection and DNA extraction

In the present study, blood stain samples were collected from 255 unrelated healthy Shanghai_Han individuals according to their household registration information. The tested samples were from 125 males and 130 females whose families resided in Shanghai for at least three generations. Written informed consent was obtained from all the individuals before sample collection. Genomic DNA was extracted from FTA cards using the Chelex-100 procedure (Walsh, Metzger, & Higuchi, 1991). NanoDrop-1000 spectrophotometry (Thermo Scientific) was employed to measure DNA concentration and quality based on the manufacturer’s instructions.

2.2 | PCR amplification and STR typing

The extracted DNA was amplified with the GeneAmp PCR System 9700 (Applied Biosystems) according to the manufacturers’ recommendations. The AGCU® Expressmarker 21+1 kit (AGCU ScienTech Corporation) includes 21 autosomal STR loci (D6S474, D22S1045, D12ATA63, D10S1248, D1S1677, D11S4463, D2S441, D16S217, D3S529, D6S1017, D4S2408, D19S133, D7S1301, D1GATA113, D18S853, D20S482, D14S1434, D9S1122, D2S1776, D10S1435, and D5S2500) and amelogenin. Amplified PCR products were separated by capillary electrophoresis in an ABI PRISM 3130xL Genetic Analyzer (Applied Biosystems). Alleles were determined according to the provided allelic ladders by GeneMapper® ID software v3.2 (Applied Biosystems). All experimental procedures were performed according to laboratory internal control standards.

2.3 | Quality control

We strictly followed the recommendations of the Chinese National Standards and Scientific Working Group on DNA Analysis Methods (SWGDAM, 2010) and the recommendations of the DNA Commission of the International Society of Forensic Genetics (ISFG) (Carracedo et al., 2013). Control DNA 9947A (AGCU ScienTech Corporation) was used as a positive control while sdH2O (AGCU ScienTech Corporation) was used as a negative control for each batch of amplification and genotyping. Moreover, the
laboratory has been accredited in accordance with ISO/IEC 17025:2005 and the China National Accreditation Service for Conformity Assessment (CNAS) (Registration No. CNAS L4476).

2.4  Statistical analysis

Allele frequencies and forensic parameters including the observed heterozygosity (HO), expected heterozygosity (HE), typical paternity index (PI), matching probability (MP), power of discrimination (PD), polymorphism information content (PIC), probability of paternity exclusion (PE), cumulative power of discrimination (CPD), cumulative probability of exclusion (CPE) and Hardy–Weinberg equilibrium (HWE) tests of the 21 STR loci were calculated by Modified-PowerStat software (version 1.2) (Zhao, Wu, Cai, & Xu, 2003). Linkage disequilibrium (LD) tests between all pairs of STR loci were performed using Arlequin v3.5.2 software (Excoffier & Lischer, 2010). To measure interpopulation differentiation, pairwise \(F_{ST}\) values and corresponding \(p\)-values were calculated by analysis of molecular variance (AMOVA) using Arlequin v3.5.2 software between the Shanghai_Han population and 23 other populations in mainland China for which data were available, including the Liaoning_Han (Xiao, Yu, & Zhou, 2015), Shandong_Han (Han et al., 2015), Henan_Han (Shen et al., 2017), Guanzhong_Han (Zhang, Tang, et al., 2015), Ningxia_Han (Wang, Liao, et al., 2013), Hunan_Han (Guo et al., 2015), Chengdu_Han (Li et al., 2018), Guangdong_Han (Lu, Qiu, Liu, Du, & Chen, 2017), Hainan_Han (Guo, Wang, Liu, Liu, & Deng, 2016), Xinjiang_Kazak (Yuan, Wang, et al., 2014), Xinjiang_Kyrgyz (Guo et al., 2018), Xinjiang_Uygur (Guo et al., 2018), Xinjiang_Xibe (Meng et al., 2015), Inner Mongolian_Russian (Wang, Shen, et al., 2013), Mongolian (Gao et al., 2014), Gansu_Yugu (Zhang, Ma, Sun, Yang, & Luo, 2015), Gansu_Tibetan (Zang et al., 2015), Qinghai_Salar (Teng et al., 2012), Hubei_Tujia (Yuan et al., 2012), Fujian_She (Yuan, Ou, et al., 2014), Yunnan_Bai (Shen et al., 2013), Yunnan_Yi (Zhu et al., 2013), and Hainan_Li (Guo, Guo, et al., 2016) populations. SPSS 24 software (IBM Corp., 2016) was used to calculate the Mantel test, which was used to test the correlation between the genetic distances and geographical distances using Molecular Evolutionary Genetics Analysis (MEGA X) software (Kumar, Stecher, Li, Knyaz, & Tamura, 2018).

3  RESULTS AND DISCUSSION

3.1  Allele frequencies and forensic parameters of the 21 non-CODIS autosomal STR loci

The allele frequencies and forensic parameters of the 21 non-CODIS autosomal STR loci are listed in Table S1. A total of 173 alleles were observed in the studied population, with corresponding allele frequencies from 0.0020 to 0.5512. There was no significant deviation from HWE after applying the Bonferroni correction, except for locus D1GATA113. This may be explained by the large-scale population mobility in Shanghai, given that HO (0.8402) was much higher than the HE (0.6767) at D1GATA113. Linkage disequilibrium (LD) tests were performed between each pair of STR loci before further analyses. As shown in Table S2, no significant linkage disequilibrium was observed in the 210 pairs of STR loci after Bonferroni correction \((p = 0.05/210 = .00024)\). Therefore, the 21 tested non-CODIS autosomal STR loci could be treated as independent loci in the following analyses. The lowest values of HE, HO, PE, and PIC were observed at locus D1S1627, equaling 0.6093, 0.6063, 0.2958, and 0.5539, respectively, while the highest values of these parameters were 0.7958, 0.8627, 0.7200, and 0.7681, respectively, at locus D19S433. The MP values ranged from 0.0061 at locus D5S2500 to 0.2469 at locus D1GATA113. In addition, the PD values ranged from 0.7531 at locus D1GATA113 to 0.9939 at locus D5S2500. The CPD and CPE values of these 21 STR loci could be treated as independent loci in the following analyses. The lowest values of HE, HO, PE, and PIC were observed at locus D1S1627, equaling 0.6093, 0.6063, 0.2958, and 0.5539, respectively, while the highest values of these parameters were 0.7958, 0.8627, 0.7200, and 0.7681, respectively, at locus D19S433. The MP values ranged from 0.0061 at locus D5S2500 to 0.2469 at locus D1GATA113. In addition, the PD values ranged from 0.7531 at locus D1GATA113 to 0.9939 at locus D5S2500. The CPD and CPE values of these 21 STR loci were 0.999999999999999999, 0.999999999999999999, 0.999999999999999999, and 0.999999999999999999, respectively, which indicates that the 21 non-CODIS autosomal STR loci were highly polymorphic and appropriate for individual identification and paternity testing in the studied population. On the other hand, the MP and CPE values of these 13 CODIS STRs in the studied population were 0.999999999999999999 and 0.999999999999999999, respectively, revealing that the utilization of non-CODIS STRs could significantly improve the discrimination efficiency of the testing system, thus providing solutions for intricate forensic cases. Raw genotyping data for the 21 tested non-CODIS STR loci are available upon request to daisydaisysun@163.com.
3.2 Multidimensional scaling and structure analyses

The pairwise $F_{ST}$ values and corresponding $p$-values are given in Table S3. The largest $F_{ST}$ value (0.00268) was observed between the Fujian_She and Gansu_Tibetan group, whereas the smallest $F_{ST}$ value (0.00000) was found three times: between the studied population and Henan_Han group, between the Xinjiang_Xibe and Chengdu_Han group, and between Mongolian and Gansu_Yugu group. In general, the $F_{ST}$ values increased with greater geographic distances. Based on the maximum value of corresponding $F_{ST}$, the interpopulation differentiation does not affect the effectiveness of this set of markers for forensic application in mainland China. The result of multidimensional scaling analysis is shown in Figure 1 to illustrate the genetic relationships among the studied population and the other 23 reference populations. In the MDS plot, Sino-Tibetan populations are labeled with yellow, Altaic populations are labeled with blue, and the Indo-European population (only Inner Mongolian-Russian) is labeled with green. The analyzed populations tended to be distributed with their languages in dimension 1, even though no clear boundaries could be detected. Specifically, the studied Shanghai_Han population and other Han populations were gathered in the central part, while ethnic minorities were scattered around them. The MDS analysis demonstrated the clustering tendency of the Han nationality and dispersion tendency of the other ethnic minorities. Besides, the two dimensional plots also suggested genetic similarities between the studied population and the other Han populations in mainland China.

The population structure analyses were performed on the basis of the genotypes for the same 21 STR loci among the studied Shanghai_Han and other previously published populations by the STRUCTURE program. Each of $K = 2–6$ with five runs was carried out; then the optimum $K$ was selected by STRUCTURE HARVESTER v.0.6.94 and the results are shown in Figure 2, suggesting that $K = 3$ was the most appropriate configuration. In the bar plot, one color represents each ancestry origin, and one bar represents each individual. One bar with several colors indicates an individual with admixed ancestry. From the bar plot, different patterns of ancestry components distribution were detected among the analyzed populations. Significant difference was reflected between the Fujian_She and the Xinjiang_Kyrgyz, which is in line with their geographic distance. Consistent with the phylogenetic and multidimensional scaling analyses, the results of the structure analyses showed that the tested Shanghai_Han population shares more similarities with Han populations from other regions of China than with other populations. The genetic components of the analyzed minorities differed from those of the Han populations. No additional subtle stratification was observed by further increasing the $K$ value.

3.3 Genetic distances and phylogenetic analyses

Nei's genetic distances, Cavalli-Sforza genetic distances, and Reynolds genetic distances were all analyzed based on 21 autosomal STR loci. The phylogenetic trees constructed

![FIGURE 1 MDS plot between the studied Shanghai_Han population and 23 reference populations. Populations from different language families are labeled with different colors. The studied Shanghai_Han population was labeled with a square and the other populations were labeled with circles. Note: “@” means “populations collected from”](image-url)
based on the genetic distances are shown in Figure 3. The studied population showed large genetic distances from the ethnic minority groups, for example, Nei’s genetic distance (0.043448) and Reynolds genetic distance (0.001820) with the Fujian_She and Cavalli-Sforza genetic distance (0.010305) with the Gansu_Yugu. In contrast, the studied population showed small genetic distances from the compared Han populations, for example, Nei’s genetic distance (0.008216) and Reynolds genetic distance (0.003100) with the Hunan_Han, Cavalli-Sforza genetic distance (0.001820) with the Shandong_Han. The calculated genetic distances between the ethnic minorities and the Han population were relatively large. Furthermore, the genetic distances among ethnic minorities were also relatively large, for example, Nei’s genetic distance (0.058813) between the Li and Tibetan group, Cavalli-Sforza genetic distance (0.016226) between the She and Yugu group, and Reynolds genetic distance (0.027740) between the She and Kyrgyz group.

The three employed measures all assume that differences between populations arise from genetic drift. However,
there are somewhat different assumptions. Nei's distance is formulated for the “infinite iso-alleles” model of mutation, in which each mutant forms a new allele. It is assumed that all loci have the same rate of neutral mutation and that the genetic variability in the population is initially at equilibrium between mutation and genetic drift, with the effective population size of each population remaining constant. Therefore, Nei's genetic distance is expected to increase linearly with time. However, the other two measures assume that frequency changes are results of genetic drift alone. The genetic distances under this assumption are expected to increase linearly with the sum over time of 1/N, where N is the effective population size. Thus if population size doubles, genetic drift will take place more slowly, and the genetic distance will be expected to increase only half as fast with respect to time. Reported simulation studies showed that the Cavalli-Sforza distance is more sensitive in distinguishing genetically similar populations and that the Reynolds genetic distance provides the highest sensitivity for highly divergent populations. It is also suggested that using the Cavalli-Sforza distance may provide less power for studies concerning human migration history (Libiger, Nievergelt, & Schork, 2009). In this study, the genetic relationship reflected by Nei's genetic distance is more similar to the relationship revealed by geographic distance. Nevertheless, the common reflections from the three different distances need more focuses. The main issue shown by three phylogenetic trees is that the Han populations tended to form one clade, while the other ethnic groups clustered together. The studied Shanghai_Han population shares more similarities with Han populations from other regions of China than with other populations.

4 CONCLUSION

In conclusion, the results suggested that the 21 non-CODIS STR loci were highly polymorphic in the tested Han population from Shanghai and hence could be utilized in forensic individual identification and parentage testing. These population data of the STR loci could be useful to enrich genetic information resources and provide reference data for population genetic studies in the future. Moreover, the interpopulation comparisons revealed population differentiation and assimilation of the Shanghai_Han and 23 other populations.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

AUTHOR CONTRIBUTIONS

K.S. and J.X. were involved in study design. C.S., and Y.L. were involved in experimental work. H.X. provide support. Z. Zhou, H.X., Y.Z., and Z.P. were involved in data collection and analysis. Z. Zhou, and K.S. were involved in writing of the initial manuscript. Z. Zhao and Q.T. were involved in revision. All authors reviewed the manuscript.

COMPLIANCE WITH ETHICAL STANDARDS

Samples were collected upon approval of Ethics Committee at Fudan University, P. R. China. This study was approved by the Ethics Committee of Fudan University, P. R. China.

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DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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**SUPPORTING INFORMATION**

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

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