| Title | Development of a method for pairwise kinship analysis based on chromosome sharing estimated from using high-density single nucleotide polymorphisms |
|-------|--------------------------------------------------------------------------------------------------|
| Author(s) | Morimoto, Chie |
| Citation | 京都大学 |
| Issue Date | 2019-03-25 |
| URL | https://doi.org/10.14989/doctor.k21698 |
| Right | |
| Type | Thesis or Dissertation |
| Textversion | ETD |
RESEARCH ARTICLE

Pairwise Kinship Analysis by the Index of Chromosome Sharing Using High-Density Single Nucleotide Polymorphisms

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Abstract

We developed a new approach for pairwise kinship analysis in forensic genetics based on chromosomal sharing between two individuals. Here, we defined “index of chromosome sharing” (ICS) calculated using 174,254 single nucleotide polymorphism (SNP) loci typed by SNP microarray and genetic length of the shared segments from the genotypes of two individuals. To investigate the expected ICS distributions from first- to fifth-degree relatives and unrelated pairs, we used computationally generated genotypes to consider the effect of linkage disequilibrium and recombination. The distributions were used for probabilistic evaluation of the pairwise kinship analysis, such as likelihood ratio (LR) or posterior probability, without allele frequencies and haplotype frequencies. Using our method, all actual sample pairs from volunteers showed significantly high LR values (i.e., >10⁸); therefore, we can distinguish distant relationships (up to the fifth-degree) from unrelated pairs based on LR. Moreover, we can determine accurate degrees of kinship in up to third-degree relationships with a probability of >80% using the criterion of posterior probability >0.90, even if the kinship of the pair is totally unpredictable. This approach greatly improves pairwise kinship analysis of distant relationships, specifically in cases involving identification of disaster victims or missing persons.

Introduction

Investigation of DNA polymorphisms, such as short tandem repeats (STRs) or single nucleotide polymorphisms (SNPs), can be used to identify familial relationships. In forensic genetics, unknown human remains, such as those used for disaster victim identification (DVI) or missing person identification (MPI), can be identified by kinship analysis between the unknown DNA sample and a reference DNA sample. In modern society, people have fewer children and
relatives. Since there are sometimes cases without reference profiles of first-degree relatives (i.e., parent-child and siblings), it is not uncommon to identify a person by using only a single reference sample from a more distant relative (i.e., grandchild, nephew, and first cousin).

In current forensic casework, DNA-typing systems of 15 STR loci are commonly used for kinship analysis. However, pairwise kinship analysis using these STR-typing systems is only feasible for the assessment of first-degree relatives [1]. SNPs have recently received attention for kinship determination in forensic fields [2–4]. We can currently analyze large numbers of SNP loci simultaneously by DNA microarray [5, 6] or next-generation sequencing [7, 8]. Analyzing genome-wide SNP data allows determination of how much heredity information is shared between two individuals and is useful for kinship determination, even for distant relationships. In previous studies, fifth-degree relative pairs could be readily distinguished from unrelated individuals by using genotype data from high-density SNPs [5, 6]. Kling et al [5] suggested that relationships as distant as second cousins (fifth-degree relationships) could be readily distinguished from both the unrelated and other relationships by calculating the posterior probabilities using genotype data from high-density SNP microarrays. Lareu et al [6] demonstrated the usefulness of genome-wide SNP genotyping to analyze claimed second cousins by counting the number of shared SNPs between two people. However, since considerable linkage disequilibrium (LD) between SNPs is ignored in these studies, these methods may be subject to false positives.

In this study, we developed a new approach for pairwise kinship analysis using 174,254 autosomal SNP loci in the HumanCore-12 BeadChip or HumanCore-24 BeadChip (Illumina, San Diego, CA, USA) and focused on chromosomal sharing between two individuals. In previous studies, rigorous identity by descent (IBD) segments on chromosomes were estimated using statistical methods (e.g., Hidden Markov Models) [9–12]. However, the purpose of these studies was IBD mapping and not kinship analysis in forensic genetics. Our approach uses “index of chromosome sharing” (ICS) between two individuals to determine the degree of kinship. The ICS value is simply calculated on the basis of identity by state (IBS) regions between two individuals and the genetic length [i.e., centi-Morgan (cM)] of these regions and results in probabilistic values [i.e., likelihood ratio (LR) or posterior probability]. Therefore, this approach is intuitive for non-experts of forensic genetics (e.g., police officers, judges, and juries).

To calculate LR or posterior probability, we proposed a method to compare the difference in the possibility of obtaining an ICS value in each relationship. Allele or haplotype frequencies are not used in this method; therefore, LD does not need to be considered for calculating LR or posterior probability. To increase calculation accuracy, the expected ICS distributions in each relationship need to be estimated. Therefore, we used computational genotypes of related and unrelated (UN) pairs for estimating the distributions. Related pairs included first- to fifth-degrees of collateral (C) relationships and first- to third-degrees of lineal (L) relationships. Computational genotypes were generated by using estimated SNP haplotypes and considering recombination between each SNP. These SNP haplotypes were estimated by the Shape-IT algorithm [13] using 1498 Japanese individuals to incorporate the effect of LD into computational genotypes.

We then validated the new method by investigating the accuracy of kinship determination. For the validation, we collected actual sample pairs from families of volunteers in each targeted relationship. In general, kinship analysis is performed to investigate whether an individual pair has a predicted relationship or is UN by calculating LR; therefore, we investigated whether relationships between up to fifth-degree relatives could be distinguished from UN pairs based on LR. To obtain more effective evidence, we also investigated the posterior probability from Bayes’ theorem, assuming all degrees of relationships. By using the posterior probability, we
can determine the degree of kinship from an ICS value, even if there is no predicted relationship between the pair.

**Results**

**Overview of a new approach for pairwise kinship analysis**

Fig 1 provides an overview of the method for pairwise kinship analysis proposed in this study. First, DNA is extracted from the samples of both the remains and the reference. Targeted relationships are first- to fifth-degrees of C relationships and first- to third-degrees of L relationships. Second, genotyping of 174,254 SNPs is performed by DNA microarray. Genotypes of samples delivering high-quality typing results (i.e., genotyping success rates > 0.99) are used for the following steps. Third, using the genotypes of two individuals from the 174,254 SNPs on the autosomal chromosomes, we calculate the ICS value. In the ICS calculation, we investigate the genetic length (cM) of all IBS segments between the two individuals, and then sum the genetic length of the IBS segments greater than a given threshold (Th). The most appropriate Th (i.e., C-Th for C relatives and L-Th for L relatives) has been determined by using computationally generated genotypes. Finally, we probabilistically evaluate the relationship of the pair by LR or posterior probability calculated by the probability density of the ICS distribution in each relationship. When the reference has the predicted relationship, LR is calculated to distinguish the relationship from the UN individuals. Alternatively, when there is no predicted relationship between the pair, posterior probability is calculated based on Bayes’ theorem to determine the degree of kinship.

![Flowchart of pairwise kinship analysis](image)

**Fig 1. Overview of the method for pairwise kinship analysis proposed in this study.** ICS is the index of chromosome sharing. C-Th and L-Th indicate the most appropriate thresholds for collateral relatives and for lineal relatives, respectively.

doi:10.1371/journal.pone.0160287.g001
The ICS between individual pairs

ICS between two individuals was calculated by the summation of the genetic length of the IBS segments greater than C-Th or L-Th. We denoted ICS for C relatives as \( ICS_C \) and ICS for L relatives as \( ICS_L \). To determine C-Th and L-Th, the summation of genetic length greater than \( Th \) was defined as the function of \( Th: ics(Th) \). We calculated the \( ics(Th) \) values from computationally generated genotypes from 249 families synthesized from the genotype data of a community-based prospective multi-omics cohort (the Nagahama Study), changing \( Th \) from 0 to 63 cM (S1 Fig). These families included five types of C relative pairs, three types of L relative pairs, and UN individuals. The five types of C relationships were sibling (C-1), uncle-nephew (C-2), first cousin (C-3), first cousin once removed (C-4), and second cousin (C-5). The three types of L relationships were parent-child (L-1), grandparent-grandchild (L-2), and great-grandparent-great-grandchild (L-3). Although relationships with the same degree of kinship are expected to have almost the same ICS values, each relationship may actually have a different ICS distribution, because the chromosomal inheritance pattern is different, even in cases with the same degree of kinship (i.e., C-2 and L-2). Therefore, we treated C and L groups separately.

By using the area under the curve (AUC) for the receiver operating characteristic (ROC) curve, we adopted 4 cM and 3 cM as C-Th and L-Th, respectively (S1 Table). There were large differences in both \( ics(4) \) (i.e., \( ICS_C \)) and \( ics(3) \) (i.e., \( ICS_L \)) between each relationship, and these \( ICS_C \) and \( ICS_L \) values became lower as the pair exhibited a more distant relationship between both C relatives (Fig 2A) and L relatives (Fig 2B).

To evaluate the accuracy of the estimated ICS (\( ICS_C \) or \( ICS_L \)) distribution from computationally generated pairs, we also investigated ICS values of actual individual pairs in each relationship, which were genotyped by the HumanCore-12 BeadChip or HumanCore-24 BeadChip. Genotyping success rates of all samples were > 0.99, which was recommended as

![Fig 2. The expected ICS variability for each relationship. Boxplots indicate the variability from the computationally generated genotypes. Triangular plots indicate the values from half of the actual samples. (A) ICS\(_C\) variability for collateral relatives using only IBS segments longer than 4 cM and (B) ICS\(_L\) variability for lineal relatives using only IBS segments longer than 3 cM.](https://doi.org/10.1371/journal.pone.0160287.g002)
good quality by the manufacturer. Using the $ICS$ values from half of the sample genotypes, we confirmed that there were no outliers by comparing these values (triangular plots in Fig 2) to those of the simulated pairs (boxplots in Fig 2).

For L-1, all loci had at least one shared allele, and the $ICS_L$ values of all pairs were the same as 3662.522, which was the maximum $ICS$ value when all SNPs were shared between a pair in simulated genotypes, because simulated genotypes did not consider uncalled SNPs and typing errors. Despite $ICS_L$ values of actual sample pairs being slightly lower than the maximum value due to typing errors in the DNA microarray, there was little difference between $ICS_L$ values from actual sample pairs and those from simulated pairs. Because the $ICS_L$ values in L-1 were completely separated from other relationships, even those with the same degree of kinship (C-1), we could distinguish L-1 from other relationships easily. Therefore, we removed L-1 from further analyses of L relatives.

Estimating the $ICS$ distribution

For kinship determination using $ICS$, we needed to know the expected distribution of $ICS$ values in each relationship. Using the $ICS$ values of simulated pairs, we estimated the $ICS$ distribution in each relationship, except for L-1. We proposed three models (i.e., normal distribution, truncated-normal distribution, and log-normal distribution) for distribution of the $ICS$ values due to the shape of $ICS$ values for simulated pairs. For example, Fig 3 shows the quantile-quantile (Q-Q) plots of the three models in C-5. The Q-Q plots indicated that the log-normal distribution was the best-fitted model, while the normal and the truncated-normal distributions differed from the observed $ICS_C$ values. In other relationship pairs (i.e., C-1 to C-4; L-2, and L-3), log-normal distributions were also fitted.

We also calculated the Akaike Information Criterion (AIC) values for each model in order to evaluate which model was the most appropriate. The model with the lowest AIC was considered the best-fit model. In C relatives, the AIC values from the normal distribution model, truncated-normal distribution model, and log-normal distribution model were 17992.85, 17992.78, and 17946.81, respectively. In L relatives, the AIC values from the three models were 9073.437, 9073.437, and 9068.232, respectively. The log-normal distribution model indicated the lowest AIC in both C relatives and L relatives. Therefore, we modeled the $ICS$ distribution

![Fig 3. Q-Q plots for the $ICS_C$ values calculated by the genotypes of computationally generated C-5. These plots are assumed as (A) the normal distribution, (B) the truncated-normal distribution, and (C) the log-normal distribution.](https://doi.org/10.1371/journal.pone.0160287.g003)
obtained from the simulated genotypes of C relatives and L relatives as a log-normal distribution. The estimated parameters (μ and σ) used in log-normal distributions for each relationship are shown in S2 Table.

In order to evaluate whether the modeled distribution of ICS values was appropriate, we compared the ICS values obtained from the SNP genotypes from half of the actual sample pairs (triangular plots in Fig 2) with the estimated distribution in each relationship by z test. All p-values were > 0.05; therefore, there was no significant difference between the estimated ICS distribution and the ICS values of the actual sample pairs. Thus, we considered that the estimated distribution properly reflected the actual ICS variability.

LR by comparing the ICS distributions of related and UN pairs

In general, kinship analysis is performed to investigate whether an individual pair has a predicted relationship or is UN by calculating LR. Using the estimated ICS distribution (i.e., log-normal distribution), we investigated the probability density obtained from an ICS value by hypothesizing a specific relationship between an individual pair. The difference in the probability density between two hypothesized relationships could be expressed as a LR. Using the rest of the actual sample pairs, we calculated the LR values based on the estimated ICS distribution. The minimum, median, and maximum LR values are shown in Table 1. Significantly high LR values were obtained from all related pairs, even C-5. Therefore, this suggested that it was possible to distinguish up to fifth-degree relationships for C relatives and third-degree relationships for L relatives from UN pairs using the method proposed in this study.

In order to investigate whether the UN pairs could be distinguished from distant relationships, we calculated the LR values assuming $H_1$ was that the pair was C-5 and $H_2$ was that the pair was UN using the $ICS_C$ values of UN. The three quantiles were $5.53 \times 10^{-7}$, $4.50 \times 10^{-5}$, and $1.10 \times 10^{-3}$. We also calculated LR values in L relatives (i.e., L-3) using the $ICS_L$ values of UN. The three quantiles were $1.25 \times 10^{-24}$, $8.16 \times 10^{-20}$, and $8.19 \times 10^{-16}$. These values suggested that we were able to discriminate UN from both C-5 and L-3.

Calculation of the posterior probabilities assuming all degrees of relationship

To determine the degree of kinship from an ICS value, we calculated the posterior probability from Bayes’ theorem assuming all degrees of relationship. The flat values of the prior

### Table 1. Three quantiles of LR values for the actual sample pairs obtained by comparing the true relationship with the UN.

| Collateral relatives | Relationship | N | Minimum     | Median     | Maximum     |
|----------------------|--------------|---|-------------|------------|-------------|
|                      |              |   | LR          |            |             |
|                      |              |   | Minimum     | Median     | Maximum     |
|                      | C-1          | 10| $1.76 \times 10^{52}$ | $6.62 \times 10^{55}$ | $7.16 \times 10^{57}$ |
|                      | C-2          | 8 | $2.66 \times 10^{39}$ | $2.87 \times 10^{43}$ | $1.38 \times 10^{47}$ |
|                      | C-3          | 7 | $1.37 \times 10^{24}$ | $1.93 \times 10^{27}$ | $6.82 \times 10^{33}$ |
|                      | C-4          | 5 | $2.76 \times 10^{9}$  | $2.28 \times 10^{16}$ | $1.05 \times 10^{18}$ |
|                      | C-5          | 3 | $1.63 \times 10^{8}$  | $2.74 \times 10^{10}$ | $6.86 \times 10^{18}$ |
|                      |              |   | LR          |            |             |
|                      |              |   | Minimum     | Median     | Maximum     |
|                      | Lineal relatives | | LR          |            |             |
|                      |              |   | Minimum     | Median     | Maximum     |
|                      | L-2          | 8 | $6.52 \times 10^{69}$ | $7.23 \times 10^{88}$ | $2.60 \times 10^{94}$ |
|                      | L-3          | 1 | $1.16 \times 10^{42}$ |             |             |

N indicates the number of pairs.

doi:10.1371/journal.pone.0160287.t001

PLOS ONE | DOI:10.1371/journal.pone.0160287 July 29, 2016
probabilities were used, because we assumed that all hypotheses were equally likely before obtaining the genotype data. Although there are a number of publications concerning the strength of evidence, judgment criteria have not been standardized. In this study, we adopted Hummel’s predicates [14] as the criteria of judgment for kinship determination. The predicates are as follows:

1. a relationship is “practically proven” if the posterior probability ≥ 0.998
2. a relationship is “highly likely” if the posterior probability is ≥ 0.99 and < 0.998
3. a relationship is “very likely” if the posterior probability is ≥ 0.95 and < 0.99
4. a relationship is “likely” if the posterior probability is ≥ 0.90 and < 0.95
5. a relationship is “undetermined” if the posterior probability is < 0.90

We first investigated the ICS range in each Hummel’s predicate using the estimated distribution for C relatives (Fig 4A) and L relatives (Fig 4B). The range is indicated by the bar under the distributions in each relationship and separated by specific colors. The deeper the color of the relationship, the higher the posterior probabilities. Table 2 shows the percentage of the distributions having posterior probabilities ≥ 0.998, ≥ 0.99, ≥ 0.95, and ≥ 0.90. For the C relatives, > 80% of C-1 and C-2 demonstrated that the posterior probabilities were ≥ 0.998. For C-3 and more distant relationships, it was difficult to obtain posterior probabilities ≥ 0.998, but > 80% of C-3 demonstrated posterior probabilities ≥ 0.90. However, it was difficult to distinguish C-4 from close relationships, because the percentages of these relationships demonstrating the high posterior probabilities (≥ 0.90) were 37.4%. Similarly, it was also difficult to distinguish C-5 from close relationships, because the percentages of these relationships demonstrating high posterior probabilities (≥ 0.90) were 56.9%. The percentage associated with C-5 was actually lower due to the lack of more distant relationships (e.g., sixth-degree relatives). For the L relatives, more than half of L-2 and L-3 demonstrated posterior probabilities of ≥ 0.998, and > 90% of pairs demonstrated posterior probabilities of ≥ 0.90. The identification rate of L-3 was actually lower due to the lack of more distant relationships (e.g., fourth-degree relatives). It may be difficult to determine more distant relationships not being considered in this study (e.g., sixth-degree C relationships or fourth-degree L relationships), but there are few opportunities to identify such relationships in forensic casework. In both C and L relatives, almost all UN pairs demonstrated posterior probabilities of ≥ 0.998; therefore, UN pairs were rarely predicted as certain relatives using our method.

We also calculated the posterior probabilities using half of the actual sample pairs (the same pairs that are shown in Table 1). The ICS values of the pairs were plotted in Fig 4, and probability densities, which were used for calculating the posterior probabilities, were obtained from the Y-axis of the graph in Fig 4. Table 3 shows the number of samples categorized by the range of posterior probabilities based on Hummel’s predicates. Detailed posterior probabilities are shown in S3 Table. For C relatives, almost all pairs of C-1, C-2, C-3, C-4, and UN supported true relationships (i.e., posterior probabilities were ≥ 0.90). However, three C-5 pairs demonstrated that the posterior probability of the true relationship was < 0.90 (i.e., categorized as “undetermined”). One of these three pairs demonstrated that the posterior probability of C-4 was 0.951 (i.e., falsely predicted as “very likely” C-4). For L relatives, all pairs of L-2, L-3, and UN demonstrated posterior probabilities of ≥ 0.90 (i.e., predicted as “likely” or higher); therefore, each relationship was determined correctly. These results suggested that this method has high discriminatory ability for comparing the closeness of up to third-degree relationships.
Fig 4. The range of ICS values in each Hummel’s predicate for kinship determination. ICS values for (A) collateral relatives and (B) lineal relatives. Solid lines indicate the estimated log-normal distributions of the $ICS_C$ and $ICS_L$ values. The colors in the bar under the distributions indicate
the range of the $ICS_C$ and $ICS_L$ values that can be obtained from the posterior probabilities corresponding to each Hummel’s predicate for each relationship. The plots indicate the $ICS_C$ and $ICS_L$ values calculated by the rest of the actual sample pairs. All actual L-1 pairs were almost equal to the maximum value of $ICS_L$ (i.e., 3662.522) and are indicated in red. * The distribution of UN is partially displayed because the density of the estimated distribution of UN is high.

doi:10.1371/journal.pone.0160287.g004

Discussion

We developed a new approach for pairwise kinship analysis by using the $ICS$ value, which was calculated on the basis of IBS regions between two individuals. Although $ICS$ value is not the total genetic length of actual chromosomal sharing (i.e., IBD segments), it demonstrates different distributions in each relationship. We can discriminate distant relationships up to the fifth-degree from UN individuals. Moreover, we can determine accurate degrees of kinship up to the third-degree in both C and L groups, even if the kinship of the pair is totally unpredictable. We also confirmed that the $ICS$ values calculated from actual samples had the same $ICS$ distributions obtained from computationally generated individuals, which was a different result from that reported in a previous study based on theoretical simulation [15]. Therefore, we have developed a realistic method for pairwise kinship analysis based on chromosomal sharing.

Here, we proposed a method for calculating LR and posterior probability by comparing the probability density of an $ICS$ value in each relationship, but not using the allele or haplotype frequencies. Sun et al [16] proposed a method of calculating posterior probability for the identification of distant relative pairs by using allele or haplotype frequencies and considering LD. In the near future, we will evaluate which method of probabilistic calculation is the most accurate for kinship analysis.

DNA microarrays used for this approach (i.e., HumanCore-12 BeadChip or HumanCore-24 BeadChip) can analyze 48 samples simultaneously at a low cost (≈$80 USD per sample). Moreover, all processes from DNA extraction to data analysis take only 3 days. Thus, this typing method is both cost- and time-effective for forensic casework, especially in cases such as DVI, where large numbers of samples need to be analyzed. Currently, pairwise kinship analysis using DNA-typing systems of 15 STR loci is only feasible for the assessment of first-degree

Table 2. Percentages of the posterior probabilities $\geq 0.998$, 0.99, 0.95, and 0.90 for the distributions of each relationship.

| Collateral relatives | Posterior probability* |
|----------------------|------------------------|
|                      | $\geq 0.998$ | $\geq 0.99$ | $\geq 0.95$ | $\geq 0.90$ |
| C-1                  | 83.0%        | 92.6%       | 97.2%       | 98.3%       |
| C-2                  | 80.6%        | 90.5%       | 95.7%       | 97.0%       |
| C-3                  | 0.0%         | 38.0%       | 71.6%       | 81.6%       |
| C-4                  | 0.0%         | 0.0%        | 11.5%       | 37.4%       |
| C-5                  | 6.66%        | 24.1%       | 46.4%       | 56.9%       |
| UN                   | 89.2%        | 94.8%       | 97.6%       | 98.4%       |

| Lineal relatives     | Posterior probability* |
|----------------------|------------------------|
|                      | $\geq 0.998$ | $\geq 0.99$ | $\geq 0.95$ | $\geq 0.90$ |
| L-2                  | 51.6%        | 73.2%       | 87.7%       | 91.9%       |
| L-3                  | 75.4%        | 83.7%       | 90.4%       | 92.7%       |
| UN                   | 99.99997%    | $\geq 99.99999%$ | $\approx 100%$ | $\approx 100%$ |

UN, unrelated.
* The flat values of the prior probabilities were used.

doi:10.1371/journal.pone.0160287.t002
relatives. While we can discriminate 69.8% of first-degree relatives (i.e., siblings) from UN, only 4.07% of second-degree relatives (i.e., half siblings) can be discriminated [17]. However, unknown human remains sometimes cannot be identified due to there being no reference profile for close relatives, such as first-degree relatives. In 2011, a tsunami following the Great East Japan Earthquake took the lives of approximately 16,000 people. Although identification of victims was performed using various methods (e.g., from information about clothing, fingerprints, dental records, and DNA analysis), >70 unidentified victims are still awaiting return to their families [18]. The new approach proposed in this study has the potential to properly identify these victims.

However, there are areas of our approach that need to be improved upon prior to application in practical forensic cases. First, additional studies are required to assess the versatility of microarray typing. In this study, we used 200 ng of DNA extracted from buccal swabs of the volunteers for the microarray; however, we often have access to smaller amounts of DNA and can obtain only degraded samples according to circumstances. We preliminarily confirmed that genotyping results with good quality (i.e., genotyping success rates > 0.99) were obtained using as little as 10 ng of DNA extracted from buccal swabs. We need to evaluate whether microarray typing can be applied to degraded forensic samples (e.g., nails, bones, or teeth).

Second, we currently have to choose from 174,254 SNPs in HumanCore-12 BeadChip or HumanCore-24 BeadChip (Illumina) when we apply this approach to actual casework. Any SNP panel can be used as long as a large-scale SNP genotype dataset is available for simulation. We expect that the length of IBS regions in autosomal chromosomes is less affected by increases or decreases in the number of SNPs. To enhance the versatility of the method, we will evaluate the effect of the set and number of SNPs on this approach.

This method is limited for use in Japanese populations, because other populations are expected to have different ICS distributions. If our approach is applied to a non-Japanese population, a large-scale SNP genotype dataset (i.e., >1000 individuals) for the population will be needed to generate distributions of ICS values. However, even if such a dataset for the
population is not available, our approach may be worth evaluating, because genotype imputation can be performed in a comparatively cross-ethnic way [19]. Additionally, use of different datasets for simulations may result in variations in the estimated ICS distribution and the most appropriate \( Th \) value (i.e., \( C-Th \) and \( L-Th \)). Further consideration will be needed to confirm the quality of \( C-Th \) and \( L-Th \).

Our approach is only useful for kinship analysis between two individuals and for applications to assess the minimal requirements depending on the number of DVI candidates. Additionally, we may need a new index other than \( ICS \) in cases where the references of two or more people are present. Furthermore, when there are many samples from both human remains and references, especially in DVI cases, it will be desirable to assess the results by comparing the remains of each individual with the entire reference, similar to the permanent method [20].

In conclusion, our approach using high-density SNP genotyping improved the accuracy for pairwise kinship analysis by using \( ICS \) values and performing probabilistic evaluation. Therefore, this approach can be applied to actual forensic casework involving personal identification for DVI and MPI cases.

**Methods**

**High-density SNP genotyping**

Buccal samples were collected from 67 Japanese individuals in five families by using a Buccal DNA Collector (Bode Technology, Lorton, VA, USA). These families contained nine types of relationships, including five types of C relationships, three types of L relationships, and UN individuals. Five types of C relationships included 20 sibling pairs (C-1), 17 uncle-nephew pairs (C-2), 14 first cousin pairs (C-3), 10 first cousin once removed pairs (C-4), and seven second cousin pairs (C-5). Three types of L relationships included 21 parent-child pairs (L-1), 17 grandparent-grandchild pairs (L-2), and two great-grandparent-great-grandchild pairs (L-3). We also collected samples from 15 UN pairs. These pairs were used regardless of sex, because we targeted only autosomal SNPs. All participants gave written informed consent, and this study was approved by the ethics committee of the Graduate School of Medicine of Kyoto University. Buccal-cell DNA was extracted using a QIAamp DNA Investigator kit (Qiagen, Venlo, Netherlands), and the extracted DNA was quantified using Quanti-iT PicoGreen dsDNA Assay kit (Invitrogen, Carlsbad, CA, USA). Each DNA sample normalized to a final DNA concentration of 50 ng/μL.

We genotyped all samples using the HumanCore-12 BeadChip or HumanCore-24 BeadChip according to manufacturer protocol. The genotyping platform adopted in this study was the Illumina Infinium HD Assay. SNP chips were scanned using iScan (Illumina) and analyzed using GenomeStudio (Illumina). Genotyping success rates of all samples were > 0.99, which is recommended by the manufacturer to ensure good quality. Half of the actual pairs were used to evaluate the estimated \( ICS \) distribution from simulated pairs, and the rest were used to validate the method proposed in this study.

HumanCore-12 BeadChip combines 12,602 InDels and 286,328 SNPs, of which 276,041 SNPs are from autosomal chromosomes and the remaining 10,287 SNPs include X and Y SNPs, mitochondrial SNPs, and non-informative SNPs (i.e., containing no information about their chromosomes from the manufacturer references). HumanCore-24 BeadChip is an update of the HumanCore-12 BeadChip and combines 12,663 InDels and 294,007 SNPs, of which 283,518 SNPs are from autosomal chromosomes and the remaining 10,489 SNPs include X and Y SNPs, mitochondrial SNPs, and non-informative SNPs. Results of InDels, X and Y SNPs, mitochondrial SNPs, and non-informative SNPs were not used in this study. We also
removed autosomal SNPs without information concerning genetic map position in SNAP version 2.2 [21].

Generating computer-based familial genotypes of high-density SNPs

We computationally synthesized a number of the familial genotypes shown in Fig 5. The family included five C relatives, three L relatives, and UN pairs. C relatives were C-1 (D-E), C-2 (D-I), C-3 (H-I), C-4 (H-L), and C-5 (K-L). L relatives were L-1 (A-E), L-2 (A-I), and L-3 (A-L). To synthesize realistic genotypes of the family, we generated founder haplotypes (i.e., A, B, C, F, G, and J in Fig 5) by considering the effect of LD and recombination. To generate these haplotypes, we used genotype data from the Nagahama Prospective Genome Cohort for Comprehensive Human Bioscience (the Nagahama Study), a community-based prospective multi-omics cohort study conducted by the Center for Genomic Medicine at Kyoto University. We used SNP data from 1616 individuals in the Nagahama Study, which were genotyped using the HumanOmni 2.5M-4 Beadchip (Illumina). Quality checks for this SNP data were performed using a series of quality control (QC) filters in a function of PLINK [22], which included individual genotyping success rate (> 0.95), SNP genotyping success rate (> 0.99), Hardy-Weinberg equilibrium ($p > 1 \times 10^{-6}$), and minor allele frequency (MAF) cutoffs (MAF > 0.01). Additionally, we did not use individually estimated first-degree relatives (i.e., parent-child and siblings) from PI-HAT to remove unexpected relationships from the genotype dataset.

After SNP data from 1498 individuals satisfied the QC, we estimated the haplotypes of these individuals for generating founder haplotypes in Fig 5. We used 174,254 autosomal SNPs as haplotype components. We did not select SNPs by MAF and genetic distances and used as many SNPs as possible to obtain rigorous IBS regions for all autosomal chromosomes for calculating ICS. These SNPs were included in both the HumanCore-12 BeadChip and the HumanCore-24 BeadChip, and genetic map positions (cM) of these SNPs were obtained from

![Fig 5. Pedigree consisted of twelve people.](https://doi.org/10.1371/journal.pone.0160287.g005)
SNAP version 2.2 [21]. Haplotypes of the 1498 individuals with the 174,254 SNPs were estimated using the Shape-IT algorithm [13].

Since there were six founders in one family, we synthesized 249 familial genotypes using the estimated haplotypes of 1494 individuals as founder haplotypes that were randomly selected from 1498 individuals. We then generated the haplotypes of descendants by taking into account recombination events. We produced recombined chromosomes of founders by using the recombination rates between adjacent SNPs. Because the genetic map distances between SNPs targeted in this study were small, the distances and recombination rates were approximately equivalent. Therefore, we regarded the recombination rates as 1/100 of cM values (e.g., if the genetic distance between two SNPs was 1 cM, the recombination rate was 0.01). We then chose one recombined chromosome at random and assigned the chromosome to the descendant. Similarly, the haplotypes of all members in Fig 5 were generated with occurrence of recombination events by generations. All programs used for simulations and statistical analyses described below were performed using the statistical software R version 3.1.2 [23] and are provided through GitHub (https://github.com/ChieMorimoto/Kinship-analysis-by-ICS).

Defining ICS between individual pairs

The ICS is a value based on IBS regions and reflects the total genetic length of chromosome sharing between two individuals. To calculate the ICS value, we first investigate the genetic length of all IBS regions, and then sum the genetic length of the IBS regions selected by a threshold for removing the effect of coincidental matches according to the following two subsections.

**Investigating genetic length of the shared segments based on identity by state.** We first order all 174,254 SNPs according to their chromosomal positions and then count the IBS states in each locus between two individuals. The IBS state has three possible outcomes: 0, 1, or 2. Uncalled SNPs in either individual are ignored in the following analysis. We choose “the shared segments” (i.e., the regions where 1 or 2 are continuously aligned). The genetic length of shared segments is expressed as the genetic distance, which is the difference between the genetic map position (cM) [21] of the first and last SNP of each shared segment. For example, assuming that the IBS state between a pair at ten SNPs is 2212010221, the shared segments are from the first SNP to the fourth SNP (i.e., 2212) and from the eighth SNP to the tenth SNP (i.e., 221). We do not regard isolated sharing, such as the sixth SNP, as a shared segment. The genetic length of “2212” represents the difference between the genetic map position of the first SNP and the fourth SNP. The genetic length of “221” similarly represents the difference between the genetic map position of the eighth SNP and the tenth SNP. We investigate the genetic length of all shared segments in all autosomal chromosomes.

**Summing the genetic length of the shared segments selected by a threshold.** Some of the shared segments obtained from the IBS state can be coincidental matches, because even UN pairs can share alleles by chance. For calculating ICS values, we have to exclude comparatively shorter shared segments in order to minimize the effect of coincidental matches. In contrast, if we exclude too many shared segments, we could miss the actual share of chromosomes.

We set a Th for the genetic length of the shared segments between two individuals and sum only the genetic length greater than the Th. The most appropriate Th value has been determined using computational genotypes of 174,254 SNPs. The summation under the most appropriate Th value is called the ICS, which is expected to become larger as an individual pair has a higher degree of kinship.

We used computationally synthesized genotypes of 174,254 SNPs to find the most appropriate Th value for discriminating each relationship. The summation of genetic length greater
than the $Th$ is defined as the function of $Th$: $ics(Th)$. We calculated the $ics(Th)$ values in each simulated pair, changing $Th$ from 0 to 63 cM, which was the genetic length of the shortest autosomal chromosome (i.e., chromosome 21). We then assessed the performance of discrimination between close-relative pairs by using the AUC. Although relationships with the same degree of kinship are expected to have almost the same $ICS$ values, each relationship may actually have a different $ICS$ distribution, because the chromosomal inheritance pattern is different, even in cases with the same degree of kinship (i.e., C-2 and L-2). Therefore, we treated C and L groups separately in this study, including for ROC analysis. The five AUC values were calculated by five comparisons of collateral relatives (i.e., C-1 vs. C-2, C-2 vs. C-3, C-3 vs. C-4, C-4 vs. C-5, and C-5 vs. UN) per $Th$. We adopted the $Th$ value that provided the maximum of the mean of five AUC values for discrimination of C relatives. Similarly, the three AUC values were calculated by three comparisons of L relatives (i.e., L-1 vs. L-2, L-2 vs. L-3, and L-3 vs. UN) per $Th$. We adopted the $Th$ value that provided the maximum of the mean of three AUC values for discrimination of L relatives. These adopted $Th$ values were the most appropriate for C relatives (i.e., C-$Th$) and for L relatives (i.e., L-$Th$). We termed the $ICS$ value under C-$Th$ as $ICS_C$ and the $ICS$ value under L-$Th$ as $ICS_L$.

### Estimating the $ICS$ distribution

For kinship determination, we need to know the expected variability of $ICS$ ($ICS_C$ or $ICS_L$) in each relationship. Therefore, we estimated the $ICS$ probability distribution. We proposed three models of $ICS$ distribution: normal distribution, truncated-normal distribution, and log-normal distribution. Truncated-normal distribution is the probability distribution of a normally distributed random variable whose value is bounded either below or above (or both). In this study, the random variable was $ICS$, and the limited range was from 0 to 3662.522, which is the maximum value of $ICS$ when all SNPs are shared between a pair. The maximum value is approximately the sum of the genetic map distances (cM) in all autosomal chromosomes. These three models in one relationship need a location parameter ($\mu$) and a scale parameter ($\sigma$). We estimated these parameters for nine types of relationships by maximum likelihood estimation using the $ICS$ obtained from 249 simulated familial genotypes. We then compared the fit of these models using a Q-Q plot and calculating the AIC:

$$AIC = 2k - 2(\log_{ML})$$

where $k$ is the number of parameters and $ML$ is the maximum likelihood in each model. The AIC is used to assess the relative goodness of fit of different models for the same dataset. We separated the C and L relatives for AIC calculation, and UN pairs were included in both C and L groups. The model with the lowest AIC was considered the best-fit model, which was adapted as the $ICS$ distribution.

To verify that the estimated $ICS$ distribution was fit to those of the actual sample pairs, a $z$ test was performed to compare the estimated distribution and $ICS$ values calculated using half of the actual sample pairs in each relationship. A $p < 0.05$ was considered statistically significant.

### Calculating LR by comparing the $ICS$ distributions between related and UN pairs

In general, kinship analysis is performed to investigate whether an individual pair has a predicted relationship or is UN by calculating a LR. Using a probability density of estimated $ICS$ distributions, we can investigate the possibility of obtaining an $ICS$ value by hypothesizing a specific relationship between an individual pair. The difference in probability density values
between two hypothesized relationships can be expressed as a LR. The LR is the ratio of the conditional probability of obtaining an ICS value of a pair under a certain relationship \( H_1 \) and the conditional probability of obtaining an ICS value of the pair under another relationship \( H_2 \). Here, the LR is defined as follows:

\[
\text{LR} = \frac{f(ICS|H_1)}{f(ICS|H_2)}
\]  

(1)

In this study, we calculated the LR values using ICS values obtained from the other half of the actual sample pairs in order to investigate our ability to discriminate a specific relationship from UN pairs. We hypothesized that \( H_1 \) represents a true relationship and \( H_2 \) represents UN individuals. For example, in a case of sibship analysis using an actual sibling pair, we first calculated the ICS value between the pair. We then obtained the probability density of the ICS value in the estimated distribution of a sibling (C-1) and the probability density of the ICS value in the estimated distribution of an UN individual. Finally, we calculated the ratio of the two densities as the LR value.

**Calculation of the posterior probabilities assuming all degrees of kinship**

To determine the degree of kinship from an ICS value in a case where there was no predicted relationship between the pair, we calculated the posterior probability from Bayes’ theorem, assuming all degrees of relationship in both C and L relatives. The posterior probability of a certain relationship \( H_i \) is defined as follows:

\[
\Pr(H_i|ICS) = \frac{f(ICS|H_i)\Pr(H_i)}{\sum_{j=1}^{n} f(ICS|H_j)\Pr(H_j)}
\]

(2)

where \( n \) denotes the number of assumed relationships and \( H_j (j = 1, 2, \ldots, i, \ldots n) \) denotes the hypothesis that an individual pair is the \( j \)th relationship. \( f(ICS|H_j) \) is the probability density of an ICS value obtained from the probability distribution in \( H_j \), and \( \Pr(H_j) \) is the prior probability under \( H_j \). The flat values of the prior probabilities were used because we assumed that all hypotheses were equally likely before obtaining the genotype data. Therefore, Eq 2 could be changed as follows:

\[
\Pr(H_i|ICS) = \frac{f(ICS|H_i)}{\sum_{j=1}^{n} f(ICS|H_j)}
\]

(3)

The posterior probabilities (i.e., \( \Pr(H_i|ICS) \)) were calculated in each assumed relationship of both C and L relatives.

Using actual sample pairs, which were also used for the LR calculation, we calculated the posterior probabilities of C and L relatives separately. In other words, when we targeted C relatives, the competing relationships were C-1, C-2, C-3, C-4, C-5, and UN, and when we targeted L relatives, the competing relationships were L-1, L-2, L-3, and UN.

Although there are a number of publications concerning the strength of evidence, judgment criteria have not been standardized. In this study, we adopted Hummel’s predicates [14] as the criteria for judging kinship determination, and also investigated the range of ICS values in each Hummel’s predicate using ICS probability distributions.
Supporting Information

S1 Fig. Boxplots of ics(Th) changing Th from 0 to 63 cM. Plots for (A) collateral relatives and (B) lineal relatives.
(DOCX)

S1 Table. Means of the AUC values changing Th.
(DOCX)

S2 Table. Estimated values of μ and σ for each relationship in the log-normal distribution.
(DOCX)

S3 Table. Posterior probabilities from actual sample pairs.
(XLSX)

Acknowledgments

We are grateful to Nagahama City Office and the Non-profit Organization Zero-ji Club. We also thank members of our lab for helpful discussions.

Author Contributions

Conceived and designed the experiments: CM KT. Performed the experiments: CM SF. Analyzed the data: CM TK SM CK YH. Contributed reagents/materials/analysis tools: RY FM. Wrote the paper: CM SM KT.

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Discrimination of relationships with the same degree of kinship using chromosomal sharing patterns estimated from high-density SNPs

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ARTICLE INFO

Keywords:
Kinship analysis
Single nucleotide polymorphisms
Same degree of kinship
Chromosomal sharing
DNA microarray

ABSTRACT

Distinguishing relationships with the same degree of kinship (e.g., uncle–nephew and grandfather–grandson) is generally difficult in forensic genetics by using the commonly employed short tandem repeat loci. In this study, we developed a new method for discerning such relationships between two individuals by examining the number of chromosomal shared segments estimated from high-density single nucleotide polymorphisms (SNPs).

We computationally generated second-degree kinships (i.e., uncle–nephew and grandfather–grandson) and third-degree kinships (i.e., first cousins and great-grandfather–great-grandson) for 174,254 autosomal SNPs considering the effect of linkage disequilibrium and recombination for each SNP. We investigated shared chromosomal segments between two individuals that were estimated based on identity by state regions. We then counted the number of segments in each pair.

Based on our results, the number of shared chromosomal segments in collateral relationships was larger than that in lineal relationships with both the second-degree and third-degree kinships. This was probably caused by differences involving chromosomal transitions and recombination between relationships. As we probabilistically evaluated the relationships between simulated pairs based on the number of shared segments using logistic regression, we could determine accurate relationships in > 90% of second-degree relatives and > 70% of third-degree relatives, using a probability criterion for the relationship ≥ 0.9. Furthermore, we could judge the true relationships of actual sample pairs from volunteers, as well as simulated data. Therefore, this method can be useful for discerning relationships between two individuals with the same degree of kinship.

1. Introduction

In forensic genetics, a pairwise kinship analysis is typically performed to investigate a relationship between two individuals by calculating the likelihood ratio [1]. The likelihood ratio is generally the ratio between the probability that certain DNA typing results would be obtained if the pair is related versus that obtained if the pair is unrelated. Meanwhile, when no predicted relationship exists between the pair (e.g., identification of bone remains, without personal belongings), all possible relationships should be compared simultaneously. We have previously proposed a method for pairwise kinship analysis using the “index of chromosome sharing” (ICS) calculated from high-density autosomal single nucleotide polymorphisms (SNPs) [2]. In the ICS calculation, we investigate the genetic length (centi-Morgan (cM)) of all identity by state (IBS) segments between the two individuals, and then sum the genetic length of the IBS segments longer than a given threshold (i.e., 4 cM), which is applied for removing coincidental matches. Because the ICS values reflect the total genetic length of shared chromosomal regions between two individuals, the values relate to the expected proportion of the shared genome (i.e., degree of kinship). Thus, the ICS values in sibling (first-degree relative), uncle–nephew (second-degree relative), first cousin (third-degree relative), first cousin once removed (fourth-degree relative), and second cousin (fifth-degree relative) were distributed as log-normal distributions, centering around the median values of 2805, 1919, 1018, 544, and 311, respectively. Using this approach, we could determine the degrees of kinship up to the third-degree, regardless of sex, even when the kinship of the pair was totally unpredictable.

However, we probably cannot distinguish relatives with the same degree of kinship (e.g., uncle–nephew and grandfather–grandson) using this approach, because these relatives have the same expected total genetic length of shared chromosomal regions. If possible, adding key individuals is effective for increasing information [3]. Otherwise, investigating markers on the sex chromosomes [4–6] might be useful to
discriminate such relationships. For example, Y chromosomal markers are useful for distinguishing paternal relatives from maternal relatives. However, it is impossible to discriminate within paternal relationships such as uncle–nephew and grandfather–grandson. Then, we can theoretically distinguish between second-degree relatives by using identity by descent (IBD) probabilities of two linked loci [7–9]. However, these studies were not verified by practical data, and the accuracy is unknown.

Chromosomal sharing patterns, including the number and the genetic length of the actual shared segments (i.e., IBD regions) on the chromosomes, are expected to differ between relatives, even for the same degree of kinship, due to differences in the frequency of meioses and chromosomal transitions from their common ancestors [10–14]. If two individuals have a collateral relationship, the shared chromosomal segments are expected to be larger in number and shorter in genetic length than those in lineal relationship with the same degree of kinship. This is because the number of recombination events from the common ancestors is larger in a collateral relationship than that in a lineal relationship with the same degree of kinship [11,13].

In this study, we further developed our previous approach to distinguish between relationships with the same degree of kinship by investigating the differences between chromosomal sharing patterns. We targeted the second and third-degree of kinship, including collateral and lineal relationships: uncle–nephew vs. grandfather–grandson as second-degree kinships, and first cousins vs. great-grandfather–great-grandson as third-degree kinships. We focused on the number of shared chromosomal segments between two individuals. The shared segments were estimated on the basis of IBS regions, the effect of coincidental matches being minimized. For a probabilistic prediction of the relationships between two individuals, we proposed a logistic regression model, using computationally generated genotypes. We also confirmed the validity of the proposed method employing actual samples that were genotyped using DNA microarray.

2. Methods

2.1. Generating computer-based familial genotypes of high-density SNPs

We utilized 249 computationally synthesized familial genotypes (Fig. 1) of 174,254 autosomal SNPs on the HumanCore BeadChip (Illumina, San Diego, CA, USA) used in our previous study [2].

The SNPs haplotypes of the founders (i.e., Nos. 1, 2, 3, 6, 7, and 10 in Fig. 1) were estimated using the Shape-IT algorithm [15] and 1498 Japanese individuals in the Nagahama Prospective Genome Cohort [16], to incorporate linkage disequilibrium into the computational genotypes. We assumed that the founders were unrelated (i.e., non-inbred pedigrees). We then generated the haplotypes of the descendants by taking into account recombination events between each SNP based on sex-averaged recombination rates [17]. The family included two types of second-degree relative pairs: uncle–nephew (Nos. 5–8 in Fig. 1) as a collateral relationship, and grandfather–grandson (Nos. 1–8 in Fig. 1) as a lineal relationship. It also included two types of third-degree relative pairs: first cousins (Nos. 8–9 in Fig. 1) as a collateral relationship, and great-grandfather–great-grandson (Nos. 1–11 in Fig. 1) as a lineal relationship.

In the SNP typing system based on DNA microarrays, uncalled events (i.e., genotypes that cannot be obtained) or genotyping errors (e.g., a true genotype, AB, is miscalled as AA) occur infrequently even in pristine DNA samples. In this study, we also included uncalled events and genotyping errors into the simulated genotypes for each probability. The values of the two probabilities were determined from the actual genotyping results of 67 pristine DNA samples typed on the HumanCore BeadChip (Illumina) in our previous study [2]. Detailed methods are presented in the Supplementary information and Supplementary Table 1. Uncalled events were randomly generated at all SNPs, with a probability of $5.97 \times 10^{-4}$. We assumed that genotyping errors occurred only in heterozygous genotypes through allele imbalance (i.e., one allele in the heterozygous genotype is undetected). Therefore, genotyping errors were introduced by converting heterozygous genotypes into randomly chosen homozygous genotypes, with a probability of $2.28 \times 10^{-4}$. All programs used for simulations were run using the statistical software R, version 3.3.2 [18].

2.2. Analyzing chromosomal sharing patterns between individual pairs

We investigated the differences in chromosomal sharing patterns between collateral relatives and lineal relatives with the same degree of kinship. We ordered all 174,254 SNPs according to their chromosomal positions and then counted the IBS state at each locus between two individuals. The IBS state has three possible outcomes: 0, 1, or 2. SNPs with uncalled events in either individual are ignored in the subsequent analysis. We then chose IBS regions (i.e., regions where 1 or 2 are continuously aligned), which could reflect chromosomal sharing. However, some IBS regions include coincidental matches, because even unrelated pairs can share SNPs by chance. Therefore, we have to exclude comparatively shorter IBS regions to minimize the effect of coincidental matches. We set a threshold for the genetic length of IBS regions between two individuals using receiver operating characteristic curve analysis. We identified IBS regions longer than the threshold (i.e., 4 cM) as apparent “shared segments.”

To analyze the total amounts of chromosomal sharing, ICS values in each pair were calculated by summing the genetic length of the shared segments in the same way as our previous study [2]. In this study, we also counted the number of the shared segments in each relationship. We then compared the ICS values and the number of shared segments between collateral relatives and lineal relatives with the same degree of kinship: uncle–nephew vs. grandfather–grandson and first cousins vs. great-grandfather–great-grandson.

2.3. Logistic regression analysis

We used a binary logistic regression analysis to determine whether the relationship of the pair is collateral or lineal because we assumed two outcomes (collateral or lineal). We modeled the probability that the pair has a collateral relationship as a function of the number of shared segments, using the following equation:

$$Pr(C|x) = \frac{1}{1 + \exp(\beta_0 + \beta_1 x)}$$

where $Pr(C|x)$ is the probability that the relationship of the pair is collateral, and $x$ is the number of shared segments. In this model, $\beta_0$ and $\beta_1$ are the logistic parameters. We estimated $\beta_0$ and $\beta_1$ for each degree of kinship. $Pr(L|x)$, the probability that the relationship of the pair is lineal, can be calculated by $Pr(L|x) = 1 - Pr(C|x)$.

We randomly selected 2/3 of the computationally generated
genotypes (including 166 collateral pairs and 166 lineal pairs for each degree of kinship) as training datasets to build the logistic models using the `glm` function in the R statistical language [18]. The remaining genotypes (including 83 collateral and 83 lineal pairs for each degree of kinship) were considered as test datasets. We calculated the $C_P$ and $L_P$ of the test datasets from Eq. (1) and evaluated the accuracy and false classification rate of the prediction model. Correct predictions essentially represent instances for which the true relationship is equal to the relationship with a higher probability. For example, when $C_P$ is 0.6 and $L_P$ is 0.4 in a collateral relative pair, the pair is correctly classified as a collateral relationship. However, these probabilities do not strongly support the collateral relationship in actual forensic casework. Therefore, we set three criteria of judgement for kinship determination by including stricter criteria referring to the classical Hummel’s predicates of paternity [19], as in our previous study [2]. The criteria were as follows:

1. a relationship is “supported” if the probability is $\geq 0.5$,
2. a relationship is “likely” if the probability is $\geq 0.9$, and
3. a relationship is “practically proven” if the probability is $\geq 0.998$.

When both $C_P$ and $L_P$ were lower than the criterion, we judged the relationship of the pair as “inconclusive” in the criterion of (2) and (3).

2.4. Validation using actual samples

To validate the prediction model, we used the SNP genotypes of actual second-degree relative pairs and third-degree relative pairs analyzed in our previous study [2]. These included 17 uncle–nephew pairs, 17 grandfather–grandson pairs, 14 first cousin pairs, and two great-grandfather–great-grandson pairs. These pairs were used regardless of sex because we targeted only autosomal SNPs. All participants provided written informed consent, and this study was approved by the Ethics Committee of the Graduate School of Medicine, Kyoto University. All samples were genotyped by using the HumanCore-12 BeadChip or HumanCore-24 BeadChip (Illumina), according to the manufacturer’s protocol. We checked the genotyping quality from the genotyping success rates by confirming that the rates were greater than 0.99, as recommended by the manufacturer to ensure good quality data.

From the genotypes of 174,254 SNPs, we investigated the number of shared segments between each pair, as well as for the computationally generated pairs. We then determined whether the pair has a collateral relationship or lineal relationship by using our prediction models, similar to the test datasets, and confirmed our prediction accuracy.

3. Results

3.1. Comparison of chromosomal sharing patterns between collateral and lineal relatives

In the simulation data, the distributions of $ICS$ values for the same
degree of kinship largely overlapped, as expected (Fig. 2). Therefore, we could not distinguish collateral relatives from lineal relatives for the same degree of kinship by using only ICS values. On the other hand, there were large differences in the number of shared segments between the uncle-nephew and grandfather-grandson pairs, despite the same degree of kinship (Fig. 3a). The minimum, median, and maximum values for the uncle-nephew pairs were 47, 63, and 79, respectively. These three quantiles were larger than the values for the grandfather-grandson pairs (29, 39, and 53, respectively). Similarly, the number of shared chromosomal segments in collateral relationships was larger than that in lineal relationships with a third-degree of kinship (Fig. 3b). The median values of the number of shared segments for first cousins and great-grandfather-great-grandson kinships were 51 and 37, respectively. From these results, the genetic length of each shared segment in collateral relationships is expected to be shorter than that in lineal relationships because the ICS values (the total genetic length of the shared segments) are approximately identical for the same degree of kinships. In fact, when we calculated the mean values and standard deviations (SDs) of the genetic length of shared segments in each pair, both values tended to be larger in lineal relationships than those in collateral relationships with the same degree of kinship. We summarized the means and SD values in each pair by using these medians among 249 pairs in each relationship (Table 1).

### Table 1

| Relationship                      | mean (cM) | SD (cM) |
|-----------------------------------|-----------|---------|
| Uncle-nephew                     | 30.5      | 28.4    |
| Grandfather-grandson             | 47.0      | 47.0    |
| First cousins                    | 19.9      | 19.6    |
| Great-grandfather–great-grandson | 26.7      | 29.9    |

3.2. Accuracy of the logistic regression model

By using the training datasets, we estimated $\beta_0$ and $\beta_1$ for the logistic regression model. In the second-degree relationship, $\beta_0$ and $\beta_1$ were $-52.5$ and $1.03$, respectively. In the third-degree relationship, $\beta_0$ and $\beta_1$ were $-24.0$ and $0.557$, respectively. Fig. 4 shows the logistic curves and the number of shared segments in the training datasets (boxplots) and indicates that the logistic models fitted the predicted relationship.

We evaluated the accuracy of the prediction models by using the test datasets (Table 2). About 80% of second-degree relatives had correctly predicted relationships, even when judged by the strictest criterion (i.e., Pr(Cix) or Pr(Lix) $\geq 0.998$). Third-degree relatives were predicted with over 70% accuracy, as the criterion was $\geq 0.5$ or $\geq 0.9$. 

![Fig. 3. Frequencies of the number of shared segments in 249 computationally synthesized familial genotypes of (a) second-degree relatives and (b) third-degree relatives.](image-url)
However, it was difficult to distinguish first cousins from great-grandparents–great-grandsons using the strictest criteria. However, false classification could be decreased because the criterion was stricter (Table 3). Meanwhile, inconclusive results increased (e.g., 13.9% of second-degree pairs and 77.1% of third-degree pairs, as the criterion was ≥0.998). To avoid incorrect determination in actual casework, we should select strict criteria, even if inconclusive results occur. In addition, when we tested the accuracy and false classification rates of the training datasets, few differences were observed (Supplementary Table 2).

### 3.3. Predicting relationships of actual samples

The genotyping success rates for all actual samples were > 0.99; therefore, all data could be used for predicting relationships. The prediction accuracy of actual samples was similar to that of test datasets (Table 4). When we adopted a probability of ≥0.5, all samples were judged as true relationships except for one first cousin pair. Because the number of shared segments in this pair was 43, Pr(C|x) and Pr(L|x) were 0.479 and 0.521, respectively. Although these probabilities slightly supported the hypothesis that the pair had a great-grandfather–great-grandson relationship, we could not judge whether either relationship was “likely” using our criteria. Furthermore, when we adopted a probability of ≥0.9 or ≥0.998, there was no false classification. These results suggest that this method has a high ability for discriminating relationships with the same degree of kinship, especially second-degree relationships.

### 4. Discussion

We had previously proposed an approach for pairwise kinship
analysis using ICS values calculated from high-density autosomal SNPs [2]. In this study, we further developed this approach to distinguish between collateral and lineal relationships with the same degree of kinship. This method investigates chromosomal sharing patterns in each relationship by counting the number of shared segments and evaluates the relationship of the pair probabilistically from logistic regression models. As expected, we determined that shared chromosomal segments in collateral relationships with the same degree of kinship are larger in number but smaller in genetic length than those in lineal relationships. Therefore, these results suggest that the number of recombination events from the common ancestors influences chromosomal sharing patterns. We also validated the utility of our method by using actual related samples from volunteers. Even when no predicted relationship exists, such as in disaster victim identification, we can determine the degree of kinship by using our previously published approach [2], and then judge whether the relationship is collateral or lineal using the method proposed in this study. Therefore, this method is useful for identifying remains in forensic cases. However, in fact, the final kinship determination of unidentified remains should be performed in combination with other information (e.g., clothing, fingerprints, skeleton morphological analysis, and dental records) [20].

Although uncalled events and genotyping errors rarely occurred in high-density SNP genotyping systems, the number of shared segments was seriously affected by the frequencies of these events. When we ignored events for computationally generated genotypes, a discrepancy between the number of shared segments in the simulation data versus the actual sample data was observed (data not shown). We estimated the probabilities of these events from the genotyping results of the actual samples analyzed using DNA microarrays (i.e., HumanCore BeadChip) and purposely added these events to synthesize realistic genotypes. Moreover, if another genotyping system is used, we should be able to estimate the probabilities depending on the quality of the system.

In this study, we focused on two types of relationships: collateral and lineal. In a certain degree of kinship, there is only one type of lineal relationship (e.g., grandfather–grandson as a second-degree of kinship). On the other hand, more than one type of collateral relationship exists in the same degree of kinship. For example, half-siblings represent collateral relationship with a second-degree of kinship. These relationships, which have not been addressed in this study, may have different chromosomal sharing patterns. Therefore, we need to evaluate chromosomal sharing patterns for other relationships, to discern more than two types of relationships.

Non-Japanese populations are expected to have different patterns of chromosomal sharing regions. If this approach is applied to another population, a large-scale SNP genotype dataset (i.e., > 1000 individuals) for the population will be needed to generate computer-based families [2] and estimate the parameters in the logistic model. Furthermore, we considered IBS regions longer than 4 CM as shared chromosomal segments. However, we understand that these IBS regions do not necessarily represent IBD regions. There are several methods for rigorously estimating IBD regions using genome-wide SNP data [21–25]. Therefore, these might improve the discrimination power between each of these relationships.

| Relationship               | n  | n ≥ 0.5 | n ≥ 0.9 | n ≥ 0.998 |
|----------------------------|----|---------|---------|-----------|
| Uncle–nephew               | 17 | 17      | 15      | 15        |
| Grandfather–grandson       | 17 | 17      | 14      | 13        |
| First cousin               | 14 | 13      | 12      | 5         |
| Great-grandfather–great-grandson | 2  | 2       | 0       | 0         |

* n indicates the number of sample pairs.

In conclusion, this method allowed us to distinguish between collateral relatives and lineal relatives with the same degree of kinship. Therefore, our approach can be applied to kinship determination involving personal identification. In the future, we plan to evaluate whether the microarray typing system can be applied to degraded forensic samples (e.g., nails, bones, or teeth).

Conflicts of interest

None.

Acknowledgments

We are grateful to the Nagahama Study Group. We also thank the members of our laboratory for helpful discussions. This work was supported by a Grant-in-Aid for JSPS Fellows (JSPS KAKENHI Grant Number 17J07871).

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.fsigen.2017.11.010.

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