

A cryptographic biometric authentication system based on genetic fingerprints

(Extended Version)

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Abstract

We specify a system for authentication and key derivation using genetic fingerprints which prevents the recovery of biometric information from data stored for verification. We present a detailed security analysis based on estimates of the entropy of the DNA data and formal security results. The scheme is shown to be robust and efficient by analysing the typical frequency and structure of errors in DNA measurements and selecting appropriate error correcting codes. As a result we obtain an authentication system that offers a security level equivalent to cryptographic keys with 73 bits and a FRR well below 1%.

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1 Introduction

Biometric authentication systems face various risks. In [15], Jain, Nandakumar and Nagar provide a systematic and thorough analysis of the vulnerabilities due to intrinsic failures and potential attacks by adversaries. One of the most serious threats is compromising of the templates database. An attacker with access to a reference template could try to impersonate a legitimate user by reconstructing the biometric trait and by creating a physical spoof [1, 7, 12]. Therefore, compromising the database can have disastrous impact on the whole authentication system. Furthermore, the potential disclosure of digitally stored biometric data raises serious concerns about privacy and data protection [13]. However, the effectiveness of access control mechanisms is inherently limited, e.g. against internal attacks or in the presence of software vulnerabilities.

The inherent fuzziness of biometric measurements rules out a simple verification of the presented template against a reference value computed with a one-way hash function, as is common for password-based authentications. In order to overcome these limitations, several schemes have been proposed which deploy error correcting codes to provide error-tolerant biometric authentication while, at the same time, preventing the recovery of biometric information from the reference data stored for verification by cryptographic techniques. These properties minimize the risk of unauthorized access to biometric templates and thereby render biometric authentication more secure and privacy friendly. Most of the schemes also use the biometric data to conceal or derive a cryptographic key¹ and are therefore also referred to as *biometric cryptosystem* or *biometric encryption*.

However, even if the stored reference data have been hashed with a one-way function, they enable an attacker to launch an exhaustive search by systematically executing the verification algorithm on a large set of candidate templates. In order to render such an approach infeasible, the biometric templates must contain sufficient entropy. Furthermore, the template must have enough entropy to ensure the secrecy and unpredictability of the concealed or derived cryptographic key.

After a detailed discussion of these aspects, Plaga [20] concluded that single conventional biometric features that are nowadays in common use do not offer the amount of entropy required for biometric cryptosystems. On the other hand, the human DNA represents a potential biometric feature which contains several megabyte of discriminatory information and hence could leverage the implementation of a secure biometric cryptosystem. Therefore, we decided to base our design on DNA data. Although genetic fingerprints do not allow real-time authentication with current technology, there are many other application scenarios where such a system could be useful. In fact, identification by genetic fingerprints is already routinely deployed for forensic and criminalistic purposes.

Our system is based on a scheme of Juels and Wattenberg [17]. By generalizing their security analysis to the case of a non-uniform distribution of the templates, we will show that the reference data stored for verification does not reveal the template. Furthermore, we will show that our system is reliable and secure against potential masquerades by estimating the false rejection rate and the false acceptance rate.

2 Previous Work

Many proposals have been made for biometric authentication in which the need to store templates for verification has been eliminated by combining error correction techniques and cryptographic methods.

¹Accordingly, in [15] the schemes are classified as *key binding* or *key generation* systems.

In [17], Juels and Wattenberg published a *fuzzy commitment scheme* which can be used as a basis for biometric cryptosystems. Their construction was based on the Hamming metric (as shown in [10], it is even optimal for that metric) and is thus appropriate for any biometrics where the impact of measurement errors on the template is regional. Juels and Wattenberg provided a strict security analysis only for the case that the biometric templates are uniformly distributed, but in Section 3.2 we will generalize their result to arbitrary distributions. Our authentication system is based on their construction.

A different approach has been taken by Juels and Sudan in [16]. Their construction was based on secret sharing techniques, making it tolerant against any reordering as well as limited deletions and insertions of the substrings defining the template. This property makes the scheme particularly interesting for biometric features like fingerprints or iris scans where such errors are commonly induced by the measurement. In [10] Dodis, Reyzin and Smith proposed an optimization of the scheme of Juels and Sudan and provided a stringent security analysis of both the original scheme and the optimization. Furthermore, a general model for biometric authentication with key derivation was defined that covers many of the published biometric cryptosystems.

In [8] and [11], specific systems were proposed for iris scans. The construction of [11] was based on the scheme of Juels and Wattenberg and also addresses general limitations of biometric key derivation (in particular, the irrevocability and limited secrecy of biometric information) by introducing hardware tokens as a third authentication factor. Both publications estimated the security of the scheme based on estimates of the entropy of iris codes. However, the analysis of the entropy achievable with current biometric measurement techniques carried out in [20] indicated that the estimates of [8] and [11] were far too optimistic.

In [3], Bohannon et al. addressed various cryptographic approaches to privacy of forensic DNA databases and provided a “solution framework” for this problem, strongly related to the present one. In particular, we will design and analyze the biological and cryptographic features of one of their suggested solutions in detail.

In [14], Itakura et al. discuss the applicability of DNA data for identification systems. They empirically verify that the genotypes at different loci are uncorrelated and give estimates for matching probabilities between distinct persons. Finally, they specify a cryptosystem using keys constructed from the genetic fingerprints. However, they neglect correlations between the maternal and the paternal part of the DNA (see Section 4.4.2 for a discussion) and do not consider potential measurement errors and error correction.

3 General Cryptographic Scheme

This section reviews how a biometric template f is cryptographically protected in the scheme of Juels and Wattenberg (section 3.1) and then quantifies its security in general (section 3.2).

3.1 The Scheme of Juels and Wattenberg

In [17], Juels and Wattenberg proposed a very simple biometric encryption scheme based on any binary (not necessarily linear) error correcting code. In the following, we will consider the generalization of the scheme to symbol based codes, i.e. codes over arbitrary finite fields. (The elements of the finite field are referred to as *symbols*.) This generalization is very useful for biometrics, because for short messages symbol based codes are far more efficient than binary codes.

Let $\mathcal{C} \subset \mathbf{F}_q^n$ be an $[n, k, 2t + 1]$ error correcting code with encoding function $G : \mathbf{F}_q^k \leftarrow \mathcal{C}$, where \mathbf{F}_q denotes the finite field with q elements. The encoding function transforms messages consisting of k symbols into n symbol code words ($n > k$), that can be retransformed into the messages even if up to t symbols of the received codeword are corrupted due to errors.

During enrolment, a secret key $s \in \mathbf{F}_q^k$ (the “message”) is randomly selected and

$$y = G(s) - f \quad (1)$$

is stored in the database. Here f is the biometric template that is obtained during enrolment. Furthermore, the secret key s is hashed with a cryptographic hash function h and $h(s)$ is stored in a database. For authentication, the template \tilde{f} presented by a user is added to the value y stored in the database. The result is $G(s) + \tilde{f} - f$ and if the hamming distance between f and \tilde{f} is at most t , $G(s)$ and hence s can be recovered. If the hash value of the recovered s matches the one stored in the database, the user is authenticated.

3.2 Security Analysis

For our security analysis we make use of the following notations. Let $\Pr(X)$ denote the probability of an event X and let $\mathbf{E}_{a \leftarrow A} [f(a)]$ be the expectation of the function value of a random variable A . The *min-entropy* of a random variable A is given by

$$\mathbf{H}_\infty(A) := -\log_2(\max_a(\Pr(A = a))),$$

and the *average min-entropy* of A given B is defined as

$$\tilde{\mathbf{H}}_\infty(A|B) = -\log_2\left(\mathbf{E}_{b \leftarrow B} \left[\max_a(\Pr(A = a | B = b)) \right]\right).$$

We use the term *B reveals u bits of A* to indicate that $u = \mathbf{H}_\infty(A) - \tilde{\mathbf{H}}_\infty(A|B)$.

Now let S, F and Y denote the random variables for s, f and y , respectively. The distribution F of the templates refers to any fixed population and can be arbitrary, i.e. we do not assume a uniform distribution of the templates. S is uniformly distributed and the distribution of Y is induced by those of S and F .

We assume that the hash value $h(s)$ does not reveal any information about the secret key s .² Consequently, we restrict our analysis to attackers who, in order to compute the template f or the secret key s of a user, take as input only the corresponding value y defined by Equation 1. The following result limits the success probability of such attackers.

Theorem 1. *Any algorithm that takes as input a random output y of the scheme and tries to output the corresponding s has at most an average success probability of $2^{-\tilde{\mathbf{H}}_\infty(S|Y)}$. Any algorithm that takes as input a random output y of the scheme and tries to output the corresponding f has at most an average success probability of $2^{-\tilde{\mathbf{H}}_\infty(F|Y)}$.*

For fixed y the probability that the output of the algorithm equals the secret key is at most

$$\max_s(\Pr(S = s | Y = y)) = 2^{-\mathbf{H}_\infty(S|Y=y)}.$$

Taking expectations on both sides yields the first statement.

The second statement follows analogously.

The following result shows that it is equally difficult to determine s from y as it is to determine f from y .

²This assumption would be fulfilled if we model the hash function h as a random oracle ([2])

Lemma 2. $\tilde{\mathbf{H}}_\infty(S|Y) = \tilde{\mathbf{H}}_\infty(F|Y)$.

For fixed y , the key s is uniquely determined by f and vice versa. Therefore, for a given y the most likely s corresponds to a unique, and hence equally likely f , and thus

$$\max_s (\Pr(S = s | Y = y)) = \max_f (\Pr(F = f | Y = y)).$$

Taking expectations over y on both sides yields the claim.

We now turn to the security of the authentication. For biometric authentications this is usually measured by the *False Acceptance Rate (FAR)*, which can theoretically be modelled as the probability that unauthorized persons are accepted as authorized, i.e. are authenticated as a legitimate user. The probability is taken over a random choice of the enrolled users and the impostor from the considered population.

For the following result, we assume that the hash function maps all $s \in \mathbf{F}_q^k$ to distinct hash values, and consequently, that the authentication is only successful if the correct secret key is recovered.³

Theorem 3. *The FAR, i.e. the probability that a random impostor is accepted as one of m randomly selected users, is limited by $m2^{-\tilde{\mathbf{H}}_\infty(F|Y)}$.*

For fixed y let $R_y(f)$ be the unique secret key s that is recovered during authentication from template f , e.g. the unique s with $|y + f - G(s)| \leq t$.⁴ In case there is no such s , i.e. if the distance of $y + f$ to the next code word is greater than t , let $R_y(f) = \emptyset$.

By assumption, an impostor is authenticated as a user U_i enrolled with (f_i, s_i, y_i) , if and only if his own template f satisfies $R_{y_i}(f) = s_i$. Thus, for given y_i the probability that a random impostor is accepted as user U_i is given by

$$\Pr(S = R_{y_i}(f) | Y = y_i),$$

which is at most $2^{-\mathbf{H}_\infty(S|Y=y_i)}$. Consequently, for a given set of users U_1, \dots, U_m the probability that a random impostor is accepted as one of the users is limited by

$$\sum_{i=1}^m 2^{-\mathbf{H}_\infty(S|Y=y_i)}.$$

Now, the result is obtained by taking expectations over the y_i (i.e. over the random selection and enrollment of the users) and applying [Lemma 2](#).

Our analysis has shown that the security of the scheme can be measured by $\tilde{\mathbf{H}}_\infty(F|Y)$, which can be determined from the entropy of the template and the number of bits of f revealed by y . In [section 4.4](#) we will estimate the entropy of templates derived from genetic fingerprints. Subsequently, we will analyse the number of bits of f revealed by y .

In [\[17\]](#), Juels and Wattenberg show that if the templates are uniformly distributed in \mathbf{F}_2^n , y reveals only $n - k$ bits of information about f and no information about s . However, biometric templates are usually not uniformly distributed. In [section 5.3](#) of [\[17\]](#) Juels and Wattenberg argue that “a good security analysis” of the scheme for a non-uniform distribution \mathcal{D} “will, in general, require detailed knowledge of \mathcal{D} ”. Fortunately, this presumption is not true: The following theorem generalizes

³This assumption can be justified by selecting a hash function with an output length considerably greater than the bit length of the secret keys.

⁴The uniqueness follows from the precondition that G is the encoding function of a $[n, k, 2t + 1]$ error correcting code.

the result of Juels and Wattenberg by giving an upper bound for the number of bits of s revealed by y for arbitrary distributions of the templates.⁵

We consider the generalized scheme over an arbitrary finite field \mathbf{F}_q . The result for the original scheme is implied by the case $q = 2$.

Theorem 4. *For any distribution F of the templates, at most $(n - k) \log_2 q$ bits of f are revealed by y , i.e.*

$$\tilde{\mathbf{H}}_\infty(F|Y) \geq \mathbf{H}_\infty(F) + (k - n) \log_2 q.$$

By definition, we have

$$2^{-\tilde{\mathbf{H}}_\infty(S|Y)} = \sum_{y \in \mathbf{F}_q^n} \Pr(Y = y) \cdot \max_s (\Pr(S = s | Y = y)). \quad (2)$$

Using the Bayes' theorem and [Equation 1](#) we obtain

$$\begin{aligned} \Pr(Y = y) \cdot \Pr(S = s | Y = y) &= \Pr(Y = y | S = s) \cdot \Pr(S = s) \\ &= q^{-k} \Pr(Y = y | S = s) \\ &= q^{-k} \Pr(F = G(s) - y | S = s). \end{aligned} \quad (3)$$

The distributions of F and S are statistically independent. Therefore, we can omit the condition $S = s$ on the right hand side of [Equation 3](#). Consequently, we get

$$\begin{aligned} \max_s (\Pr(Y = y) \cdot \Pr(S = s | Y = y)) &= q^{-k} \max_s (\Pr(F = G(s) - y)) \\ &\leq q^{-k} \max_z (\Pr(F = z)) \\ &= q^{-k} 2^{-\mathbf{H}_\infty(F)}. \end{aligned} \quad (4)$$

[Equation 2](#) and [Equation 4](#) yield

$$\begin{aligned} 2^{-\tilde{\mathbf{H}}_\infty(S|Y)} &\leq \sum_{y \in \mathbf{F}_q^n} q^{-k} 2^{-\mathbf{H}_\infty(F)} \\ &= q^{n-k} 2^{-\mathbf{H}_\infty(F)}, \end{aligned}$$

and with [2](#) we obtain the desired result.

4 DNA as Biometric Feature

In this section, we summarize some basic properties of short tandem repeats ([section 4.1](#)), review the analytical tools for DNA typing ([section 4.2](#)), analyse its reliability ([section 4.3](#)) and the information content of the resulting data ([section 4.4](#)).

4.1 Short Tandem Repeats

The most common DNA variations used for the identification of individuals, in particular in forensic applications, are *Short Tandem Repeats (STR)* – arrays of 5 to 50 copies (repeats) of the same pattern (the *motif*) of 2 to 6 base pairs. Two properties make STR particularly eligible for identification purposes:

⁵In [\[10\]](#), a much weaker result has been shown for a variant of the scheme.

- The number of repeats of the motif is highly variable among individuals, even in small populations.
- Forensic STR are typically located in the non-coding regions of the DNA and (at the time of writing) no biological functions of these STR loci are known.⁶

The human genome contains several 100,000 STR loci, i.e. physical positions in the DNA sequence where an STR is present. Today, approximately 20 STR loci are in practical forensic use,⁷ and some more can be considered as candidates. In order to optimize their suitability for forensic applications these loci have been selected to maximize the variability of the genotype and to minimize the likelihood of any association between the genotype and a biological function, or between the genotypes of different loci.

As with any other DNA polymorphism, an individual variant of an STR is called *allele*. Alleles are denoted by the number of repeats of the motif. In some cases, one or more motifs may not be complete within the STR, in which case the allele is denoted by a decimal number, where the digit after the decimal point equals the number of base pairs modulo the length of a complete repeat. For instance, if the motif is AGT the allele AGTGTAGT is denoted as 2.2.

The genotype of a locus comprises both the maternal and the paternal allele. If these two alleles are identical the genotype is called *homozygous*, and if they are different it is called *heterozygous*. However, without additional information, one cannot determine which allele resides on the paternal or the maternal chromosome, i.e. allele combinations (A, B) and (B, A) are indistinguishable. Therefore, genotypes are denoted by the allele numbers in ascending order, i.e. as (A, B) with $A \leq B$.⁸ The range of possible genotypes differs from one STR locus to another. Typically, there are 10-20 different alleles known per locus. However, these alleles are not of equal frequency in a given population. In [section 4.4](#) we give estimations of the entropy per locus.

For a given set of loci, the combined genotypes at these loci are called an *STR profile*. If the set of loci is large enough to allow reliable distinction between individuals, the profile is also referred to as a *genetic fingerprint*.

4.2 Measurement of STR Data

The measurement of an STR profile is conducted by specialized laboratories using commercially available STR kits. We briefly sketch the methods deployed for the measurement of STR data. Further details on this procedure are described in [\[6\]](#), p. 313 onwards.

In order to determine an STR profile, the DNA sequence of each STR locus is tagged by nearby unique sequences, called primers, and amplified using polymerase chain reaction (PCR, see [\[6\]](#), p.63 onwards, or section 2.1.7 of [\[18\]](#) for details). The number of repeats in an STR allele is then determined by electrophoresis of the amplification product, which measures the weight and thus indirectly the length of the molecules. From the electrophoretic drift, a fluorescence image is obtained which is scanned electronically to calculate an *electropherogram*, an x-y-plot of density versus molecular weight (see [Figure 1](#)). Peaks in this x-y-plot are interpreted as occurrences of alleles and by comparing the plot with a template plot (the *allelic ladder*) that contains the peaks for all relevant alleles (i.e. all alleles present in a population), individual alleles are identified.

⁶However, correlations between STR genotype and ethnical, regional or familial affiliation exist.

⁷Of these, 13 are the CODIS (Combined DNA Index System) core loci, which are the basis of the forensic system used by the FBI.

⁸A homozygous genotype is sometimes denoted by the single allele, i.e. as (A) .

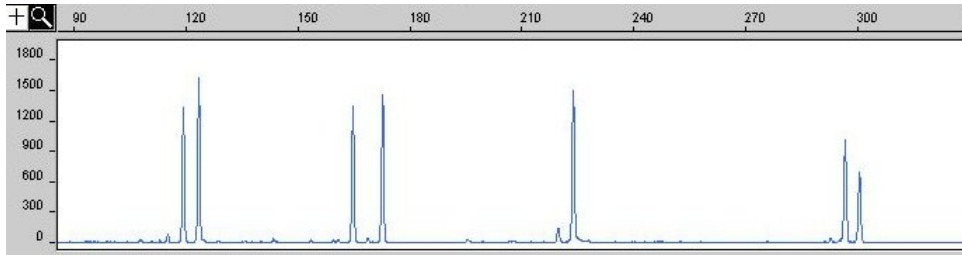


Figure 1: Part of an electropherogram, displayed with the GeneScan[®] software by Promega. The horizontal axis measures the size of the amplified DNA fragment; the vertical axis measures the intensity of the fluorescence which correlates with the number of DNA fragments of that size. Usually, electropherograms contain four such plots (taken at different colours), each of which corresponds to a different set of loci

At the time of writing, an STR profile can be determined in approximately 4 hours. Since this time is sufficient for typical forensic applications, manufacturers of STR kits have presumably made only limited efforts for further optimization.

4.3 Reliability of STR measurements

The reliability of STR measurements is assessed by comparing the results reported under different conditions, by different laboratories, or using different commercial kits.

4.3.1 Types of Errors in DNA analysis

Like any measurement, DNA analysis is not free of errors. Several types of error can be distinguished:

- *PCR primer mismatch.* The polymerase chain reaction may fail to amplify an STR fragment because the primers fail to bind to the corresponding strands of the sample DNA. This can be due to mutations in the DNA of the proband or due to degradations of the sample. We refer to [9] and [7] for details.
- *Experimental inaccuracy.* If the experimental conditions of the electrophoresis (e.g. temperature and salt concentration) are not within the required ranges, the fragments may not align as expected.
- *Misinterpretation of the electropherogram.* The electropherogram may contain peaks which do not correspond to actual STR alleles. Such artifacts may then be wrongly counted as alleles, or the peaks of true alleles may wrongly be disregarded as artefacts.
- *Human errors* like an exchange or contamination of samples, shifted application of the allelic ladder, or typing errors when recording the results.

The latter two sources of errors can be minimized by automated processing whenever possible, e.g. through the evaluation of the electropherogram by special software.

STR measurement errors are commonly classified into three groups:

1. *Allelic drop-outs.* An allele of a heterozygous genotype is missing, e.g. genotype (7, 9) is measured as (7, 7).

2. *Allelic drop-in*. In a homozygous genotype, an additional allele is erroneously included, e.g. genotype (10, 10) is measured as (10, 12).
3. *Allelic shift*. An allele is measured with a wrong repeat number, e.g. genotype (10, 12) is measured as (10, 12.2).

4.3.2 Estimation of Error Rates

The German DNA Profiling Group (GEDNAP) regularly performs ring experiments to assess the quality of laboratories performing forensic STR analysis. A central laboratory prepares several sets of identical samples. These samples are sent to all participating laboratories, who send back their results for all or a subset of samples at their discretion. The organising laboratory verifies the correctness of the results by comparison to their own, carefully conducted measurements. We refer to [21] for details.

In 2005, GEDNAP 30 and 31 were conducted with more than 175 laboratories, reporting approximately 19,000 measurements. An analysis of the (anonymized) data provided to us by GEDNAP showed that few laboratories account for most of the errors, whereas the vast majority of the laboratories have low error rates.⁹ Assuming a minimum quality standard for biometric measurements, we evaluated the error rates with the results of the four worst laboratories disregarded. [subsubsection 4.3.2](#) summarizes the results for each type of error.¹⁰ For allelic shifts we distinguished between homozygous and heterozygous genotypes; this distinction is useful for the selection of an eligible encoding of the template (see [section 5.1](#)).

Type of Error	Number	Frequency
Allelic drop-ins	7	0.04%
Allelic drop-outs	8	0.04%
Allelic shifts (het.)	19	0.10%
Allelic shifts (hom.)	2	0.01%
All errors	36	0.19%

Table 1: Errors reported in GEDNAP 30 and 31

The results in [subsubsection 4.3.2](#) show that error rates below 0.2% are achieved when a minimum quality standard for the laboratory is assumed. Furthermore, none of the good laboratories measured more than one locus incorrectly which indicates that measurement errors occur independently for different loci.¹¹

4.4 Entropy of STR Data

As shown in [section 3.2](#) the security of the authentication depends on the entropy of the templates. We now estimate the entropy of an STR profile based on the 28 loci which include all loci used in the GEDNAP 30 and 31 ring experiments ([21]).

4.4.1 Assumed Probability Distribution

The entropy of a biometric template is generally limited by the size of the population from which the enrolled individuals are chosen. For instance, for the German

⁹GEDNAP members explain the bad performance of some laboratories by their poor equipment and the lack of proper quality management.

¹⁰The classification of errors is not always unambiguous. However, the error rates based on different interpretations differ only marginally.

¹¹This finding is a bit surprising as STR measurements are usually performed for many loci simultaneously using commercial multiplex STR kits.

population the entropy could not exceed $\log_2(8 \cdot 10^7) \approx 26$. In terms of security, this bound corresponds to a brute force attack that exhaustively searches through the STR profiles of all individuals within the population. For sufficiently large populations however, a comprehensive list of templates is usually not available and consequently, a brute force attack on a large scale biometric application has to be based on a much wider search space. In particular, the relevant search space and probability distribution of the templates is implied by the statistical data available on biometric sub-features. In other words, we must consider the distribution of STR profiles that can be extrapolated from statistical data under the assumption that the population is infinite.¹²

As pointed out by Bohannon et. al. [3], an analysis based on this idealized probability distribution may not be adequate in the presence of a large scale (e.g. nation wide) DNA database. Furthermore, the potential search space for attackers being related by blood to enrolled users is considerably smaller.

4.4.2 Entropy of a single locus

The probability of the occurrence of an allele in a given population is estimated from the corresponding allele frequency, as observed in scientific experiments. For many populations, allele frequencies are readily available from the literature.

In order to estimate the genotype probabilities for a given locus from individual allele probabilities, we assume that the population in question is in so-called *Hardy-Weinberg equilibrium*. This assumption is fulfilled if the population is sufficiently large and panmictic, i.e. the mating behaviour is random, and if there is no migration or selection (see [6], p. 484, or [18], p. 65–67). While the assumption of the Hardy-Weinberg equilibrium is clearly an idealization, it is widely accepted as a method to obtain good approximations for DNA profile distributions for the large populations of industrialized countries.

If a population is in Hardy-Weinberg equilibrium, the frequency of the homozygous genotype (A, A) is P_A^2 , and the frequency of the heterozygous genotype (A, B) is $2P_AP_B$, where P_A and P_B are the frequencies of alleles A and B , respectively. We refer to [6] or [18] for details. Thus, the min-entropy of the genotype G at a locus is given by

$$\mathbf{H}_\infty(G) = -\log_2 \left(\max \left(\max_A (P_A^2), \max_{A,B} (2P_AP_B) \right) \right).$$

4.4.3 Entropy of a Set of Loci

The frequencies of the compound genotypes at all loci used in a profile can be calculated from the genotype frequencies of the individual loci under the assumption that the genotype distributions of distinct loci are statistically independent. In this case, the frequency of a compound genotype is the product of the genotype frequencies at the individual loci (multiplication rule) and, consequently, the entropy of an STR profile P is the sum of the entropies of the contributing single loci genotypes G_l . Consequently, we obtain

$$\mathbf{H}_\infty(P) = \sum_l \mathbf{H}_\infty(G_l),$$

where indices l in the sums refer to the contributing loci.

The assumption of statistical independence of the genotype frequencies at different loci is an idealization, the applicability of which depends on a sufficiently

¹²This approach is also used for assessing the assurance of evidences based on genetic fingerprints, e.g. in court cases.

high level of homogeneity of the relevant population, and on an appropriate choice of the loci. The assumption of sufficient homogeneity is widely considered valid for populations of large western countries (see [18], pp. 78 for a discussion). The second condition is fulfilled for the loci commonly used in forensics because a main objective for their establishment was the minimization of potential dependencies.

4.4.4 Experimental Estimation of the Entropy

In Table 2 we have estimated the entropy for the 18 loci used by GEDNAP 30 and 31 (ACTBP2 to VWA) and 10 additional loci for which sufficient statistical data is available. In order to match the population distribution in a potential application we have based these estimations on allele frequency data from German and Austrian populations, respectively, which have been obtained from a database of the Institute of Forensic Medicine at the University of Düsseldorf¹³. The intervals denote the 68.25%-confidence interval, i.e. the standard deviation of the estimated values of the min-entropy. We refer to [4] for details.

Locus	Entropy	Locus	Entropy
ACTBP2	6.22 ± 0.06	Penta E	4.05 ± 0.21
CSF1PO	2.56 ± 0.05	TH01	2.82 ± 0.03
D13S317	2.56 ± 0.07	TPOX	1.77 ± 0.03
D16S539	2.53 ± 0.11	VWA	3.07 ± 0.02
D18S51	4.39 ± 0.05	CD4	2.12 ± 0.04
D19S433	2.64 ± 0.11	D12S391	4.34 ± 0.08
D21S11	3.27 ± 0.04	D1S80	2.57 ± 0.03
D2S1338	4.06 ± 0.15	D8S1132	4.10 ± 0.11
D3S1358	3.05 ± 0.03	F13A1	2.18 ± 0.03
D5S818	2.04 ± 0.04	F13B	2.30 ± 0.03
D7S820	3.25 ± 0.06	FES/FPS	2.02 ± 0.02
D8S1179	2.87 ± 0.11	HLA-DQ α	2.97 ± 0.04
FGA	3.89 ± 0.03	HPRTB	2.43 ± 0.08
Penta D	3.16 ± 0.17	LIPOL	2.09 ± 0.05

Table 2: Estimated entropy of several STR loci for the population of Germany and Austria.

Summing up yields a min-entropy of 85 for an STR profile based on these 28 loci.

5 An Authentication System Based on STR Data

We specify a biometric encryption scheme based on the scheme of Jules and Wattemberg (Section 3.1) using templates obtained from Short Tandem Repeats (STR) in human DNA. Furthermore, we analyse its properties on the basis of our previous results and justify our design decisions.

5.1 Encoding and Error Correction

As described in Section 4.1 an STR profile is composed by the genotypes of the individual loci, each of which given by an ordered pair (a_i, b_i) of (not necessarily integer) numbers representing the alleles on both chromosomes at this locus.

¹³www.uni-duesseldorf.de/WWW/MedFak/Serology/dna.html

We construct the templates from STR profiles comprising the 28 loci listed in [Table 2](#). An appropriate encoding of the STR profiles to a template should minimize the impact of measurement errors to the template with respect to the Hamming metric. Based on the characterization of errors in DNA fingerprinting in [Section 4.3](#), our selection of the encoding function was led by the following considerations:

- *Independent encoding for each locus:* Each error in an STR measurement only affects the genotype of a single locus. Therefore, we encode each locus independently and concatenate the result using a fixed order of the loci.
- *Encoding of homozygous genotypes:* In order to prevent bit insertions or deletions resulting from allelic drop-ins or drop-outs, we must encode homozygous genotypes by the same number of bits as heterozygous genotypes. There are two options to accomplish this:
 1. The allele number is doubled in the encoding, i.e. a genotype (A, A) is encoded as $a||a$, where a is an encoding of A and $||$ denotes concatenation. With this encoding, at least half of the bits in the encoding remain correct for allelic drop-ins and drop-outs.
 2. The allele number is encoded and concatenated with a constant, e.g. genotype (A, A) is encoded as $a||0$. With this encoding, at least half of the bits in the encoding remain correct for allelic shifts.

Our analysis in [Section 4.3](#) shows that allelic shifts of homozygous genotypes are very rare compared to allelic drop-ins or drop-outs. Therefore, we decided to use the first encoding method.

- *Constant number of bits per allele:* In order to enable the application of symbol based error correcting codes (see [Section 5.2](#)), we use the same number of bits for each locus. For the 28 loci considered in [Section 4.4](#) the cardinality of the domains of allele numbers is limited by 64. Therefore, we decide to represent each allele by 6 bits. The actual coding of alleles into integers can be done by any numbering of the known alleles.

Using the 28 loci listed in [Table 2](#) and 12 bits per locus, we obtain templates with 336 bits.

5.2 Error Correction

Generally we can choose between symbol-based codes and binary codes. Both have their advantages and disadvantages:

- **Symbol-based codes:** Good symbol-based codes only introduce minimal redundancy even for small bit lengths. In particular, Reed-Solomon codes match the Singleton bound (see [\[19\]](#)) by which $2t$ symbols redundancy are required to correct t symbol errors. Hence, we would only need 2 symbols (i.e. 12 bits) redundancy. On the other hand, symbol-based codes are unable to reckon the severity of an error, i.e. each type of error (see [Section 4.3](#)) is treated as a symbol error regardless of the number of affected bits.
- **Binary codes:** In contrast, binary codes require much more redundancy. For bit lengths around 330 the best general construction are BCH-Codes over F_{2^9} , which can correct t bit errors using $9t$ bits redundancy (see [\[19\]](#)). In order to tolerate one wrong allele, we would need 54 bits redundancy.

Since we are mainly interested in minimising the redundancy (and, hence, the loss of entropy), we decided to choose a Reed-Solomon code. Because each allele is coded with 6 bits, the natural choice is $q = 64$, i.e. a Reed-Solomon code over \mathbf{F}_{2^6} . Considering the low error probabilities of good laboratories in the STR measurements, we suggest to select $t = 1$, i.e. a code that is able to correct 1 symbol error (coding one allele). Consequently, we decided in favour of a $[56, 54, 2]$ -Reed-Solomon over \mathbf{F}_{2^6} and obtained the following parameters: $q = 64$, $n = 56$ and $k = 54$. This code introduces 2 symbols (i.e. 12 bits) redundancy and can correct 1 symbol error.

5.3 Security

As shown in [Section 4.4](#) the min-entropy of an STR profile based on the 28 loci listed in [Table 2](#) (and hence of our templates) is approximately 85. From [Section 3.2](#) we know that the scheme of Juels and Wattenberg leaks at most $(n - k) \log_2 q$ bits on the biometric template f and $n \log_2 q - \mathbf{H}_\infty(F)$ bits of information on the secret key s . Using [Theorem 4](#) and $q = 64$, $n = 56$ and $k = 54$ we can estimate

$$\tilde{\mathbf{H}}_\infty(F|Y) \geq 73.$$

As has been shown in [Theorem 1](#) and [Lemma 2](#) this implies that an attacker who tries to determine the secret key s or the template f from the reference data y has at most a success probability of 2^{-73} .

Using [Theorem 3](#), we obtain

$$FAR \leq m2^{-73},$$

where m is the number of enrolled individuals.

5.4 False Rejection Rate (FRR)

Our determination of the FRR is based on the error rates of STR measurements as determined empirically in [Section 4.3](#). Since this database is rather small, we can only obtain a rough estimate for the FRR.¹⁴ As suggested by the results of the GED-NAP ring experiments, we assume that measurement errors occur independently for different loci.

In our setting, the BioKey system can correct up to one symbol error. With our basic encoding (see [Section 5.1](#)) most measurement errors only result in one wrong symbol (*single error*); the only exception is an allelic shift in the measurement of a homozygous genotype which results in two wrong symbols (*double error*). Therefore, we can estimate $FRR = p_d + p_s$, where p_d is the probability that at least one double error occurs in 28 independent measurements and p_s is the probability that more than one single error occurs.

From [Table 4.3.2](#) we can infer that double errors only occurred twice in approximately 19,000 experiments and hence we estimate $p_d \approx 28 \cdot 2/19,000 \approx 0.003$. On the other hand, $p_s = 1 - (1 - p)^{28} - 28p(1 - p)^{27}$, where p is the probability of a single error in a single locus measurement. From [Table 4.3.2](#) we conclude that $p \approx 34/19,000$ for good laboratories. This yields $p_s \approx 0,001$ and hence $FRR \approx 0.4\%$.

¹⁴As the FRR is not a measure of security but only of user comfort, this situation is still satisfactory.

5.5 Reference Implementation

A reference implementation has been developed as a proof-of-concept for the “BioKey system” constructed theoretically in the previous subsection. This reference implementation is a small demonstration program developed with MatLab© which provides the following functionalities:

- Flexible parameterization of the scheme by entering n and the error correction capability $t = (n - k)/2$.
- Generation of a random codeword $c = G(s)$, where s is the secret key, and computation of its hash value.
- Input of a reference template f or generation of a random template.
- Calculation of the public string y .
- Input of a template f' for verification or generation of a random template.
- Reconstruction of the code word $G(s)$ and verification of correctness by its hash value.

6 Summary and Conclusion

In this paper, we have examined the feasibility of a biometric authentication system based on genetic fingerprints that does not store genetic data in clear. The system allows the authentication of users, based on information encoded in their genome, and prevents a disclosure of the biometric information from the reference data stored in databases.

Our system is based on the biometric cryptosystem of Juels and Wattenberg. We have been able to generalize the security analysis of this scheme to the case of non-uniformly distributed templates, and to prove an upper bound of the FAR.

Our system uses a genetic fingerprint template obtained from short tandem repeats (STRs) on 28 loci in the DNA with an entropy of about 85 bits. The error rate of the measurement of the genetic information has been determined experimentally using the data of existing quality experiments for genetic labs. Error correcting codes have been applied to effectively reduce the false rejection rate to the low value of 0.4%. It has been found that approximately 70 bits of discriminatory information can be extracted using this method. This amount of information makes an exhaustive search through the template space infeasible and provides a very low FAR. By adding more suitable loci the security of the system could even be increased.

Further research activities, as for example regarding performance aspects and the unpredictability of allele combinations, are necessary and actually performed by the forensic community. Furthermore, an analysis of potential threats arising from correlations of templates among relatives could be useful.

Comparing our results with previous publications concerning schemes based on other biometric information like iris images, fingerprints, etc., our proposed system shows a significant improvement of the security due to the higher entropy of the DNA data compared to the limited information content of other biometric features.

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