Novel Hairpin-Shaped Primer Assay To Study the Association of the −44 Single-Nucleotide Polymorphism of the DEFB1 Gene with Early-Onset Periodontal Disease

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A powerful, cost-effective new method for studying single-nucleotide polymorphisms (SNPs) is described. This method is based on the use of hairpin-shaped primers (HP), which give a sensitive and specific PCR amplification of each specific allele, without the use of costly fluorophore-labeled probes and any post-PCR manipulation. The amplification is monitored in real-time using SYBR Green I dye and takes only 

The completion of the Human Genome Project has led to researchers becoming increasingly interested in studying single-nucleotide polymorphisms (SNPs). SNPs are the most abundant form of genetic variation between individuals; some have been observed to cause several diseases (e.g., sickle cell anemia [28]) and have been proven to be useful tools for identifying genes involved in common polygenic and multifactorial diseases [8]. SNPs have also proven to be useful tools for understanding the pathogenesis of several diseases (e.g., [4]).

Most of the methods available for SNP detection, such as techniques based on mass spectrometry [20], high-pressure liquid chromatography [23], microarray analysis [6], Taqman probes [16], or molecular beacons [25], are generally costly, time consuming, and/or difficult to automate, hindering their implementation in most laboratories.

The recently developed hairpin-shaped primer (HP) assay is a feasible alternative method for SNP detection, which is easy to implement in a wide variety of laboratories [10]. This method offers an improved specificity over conventional amplification refractory mutation system (ARMS) assays [22] because it uses primers that carry a 5′-end nucleotide tail complementary to the 3′ end of the SNP-detecting primer, creating a hairpin structure. This prevents nonspecific priming. The 3′ end of the HP hybridizes to the SNP residue. Two HPs are designed, one complementary to each allele, together with a common linear primer. Two PCRs are performed in parallel, each with one of the two HPs, using the same DNA template, and the amplification is monitored in real-time, by fluorescence increase, using the free fluorophore SYBR green I. An HP that is fully complementary to the target DNA will yield a more efficient amplification than a mismatched reaction, leading to an earlier threshold cycle (Ct). The greater the difference between the Cts of the mismatched and matched primer-template reactions (∆Ct) for each HP, the more robust the assay is. The technique has been developed and proven to be effective in prokaryotic contexts [10]. This study presents the first report of the application of a HP assay to SNP genotyping in a human disease.

Early-onset periodontal disease (EOP) is a complex multifactorial disorder triggered by colonization of the oral epithelia by bacteria and yeasts, which leads to a gingival inflammation, loss of alveolar bone, and tooth loss [27]. The importance of the protective role played by β-defensins against pathogenic microorganisms in oral epithelia has been discussed by several authors [5, 15, 18]. Moreover, a recent observation by Putsep et al. on patients suffering from morbus Kostmann [24] highlights the role of antimicrobial peptides in periodontal disease. Epidemiological studies clearly demonstrate that genetic factors play a significant role in EOP [9, 21], underscoring the importance of applying genetics approaches to study EOP. The C→G transversion at position −44 of the DEFB1 gene (which encodes human β-defensin 1) has been associated with Candida sp. carriage in the mouth [12] and with the risk of human immunodeficiency virus type 1 infection in the Caucasian population [1].

In order to establish if the −44 SNP is associated with EOP, we studied its distribution in EOP patients and healthy controls from different ethnic backgrounds.

MATERIALS AND METHODS

DNA samples. Anonymous, historically acquired DNA from two cohorts coming from different ethnic backgrounds was analyzed. DNA samples from 76 Caucasian EOP patients and 44 African-American EOP patients were included.

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RESULTS

For the HP assay, the three possible genotypes were identified as follows: (i) Allele C homozygous samples gave an earlier \( C_t \) with the C HP; (ii) allele G homozygous samples gave an earlier \( C_t \) with the G HP; (iii) heterozygous samples gave similar \( C_t \)s in both reactions. Figure 1 demonstrates that the HP assay allows a clear discrimination of the three different genotypes, using human DNA samples.

Samples were analyzed in sets of up to 44 in 96-well plates, together with the three artificial templates and a no-template negative control, in less than 2 h. The HP assay was able to identify C/C, C/G, and G/G genotypes with average \( \Delta C_t \)s of 11.6, 0.3, and 7.7, respectively. Results are summarized in Table 1.

Genotypes were confirmed by direct PCR sequencing of the region containing the C→G transversion at position −44 of the \( DEFB1 \) gene. Five samples for each genotype were selected for this purpose. Sequencing data agrees with the HP assay in all cases (data not shown).

Genotype and allele frequencies for the four populations studied are shown in Table 2. All the cohorts agree with the Hardy-Weinberg law, further confirming the reliability of the assay. No significant differences were observed in the allele or genotype frequencies at position −44 of the \( DEFB1 \) gene among patients suffering from EOP versus healthy controls in the two ethnic populations studied. The C-allele frequency for the Caucasian (23%) and African American (5%) healthy controls agrees with previously reported data by Jurevic et al. (13).

DISCUSSION

We have developed a useful assay for medium-scale association studies of human populations based on an ARMS principle. The HP assay uses unlabeled hairpin-shaped primers, designed following the conditions and the parameters recommended for the design of molecular beacons (26). Kaboev et al. (14) reported that HPs increase the specificity and the sensitivity of PCRs by strongly diminishing the formation of primer dimers and nonspecific products. The superiority of

![FIG. 1. HP assay for detection of the −44 \( DEFB1 \) SNP. Real-time PCR results using chromosomal DNA extracted from blood from two homozygous (C/C and G/G) patients and a heterozygous (C/G) patient. Amplifications using the C-allele HP are represented by closed triangles (▲), and the G-allele HP is represented by open circles (○).](http://cvi.asm.org/)

| Genotype | C-Allele HP \( C_t \) (CI) | G-Allele HP \( C_t \) (CI) | \( \Delta C_t \) |
|----------|--------------------------|--------------------------|-----------------|
| C/C      | 17.4 (17.3–17.5)         | 29.0 (28.8–29.2)         | 11.6            |
| C/G      | 17.7 (17.5–17.9)         | 18.0 (17.9–18.1)         | 0.3             |
| G/G      | 26.4 (25.7–27.1)         | 18.7 (18.2–19.3)         | 7.7             |

*CI, 95% confidence intervals; \( \Delta C_t \), mismatched primer − \( C_t \), matched primer.*
these primers over conventional linear primers for SNP detection has previously been demonstrated (10). However, this is the first report describing the use of HPs to increase the specificity of an ARMS assay for human DNA samples.

The HP assay was designed using the conditions recommended by Hazbón and Alland (10) and required no further modifications, illustrating the robustness of the technique. A secondary mutation was introduced in the HPs to avoid an undesired secondary structure. Secondary mutations have also been reported to improve the discriminatory power of the assay (3, 10, 22, 29).

The use of the SYBR Green I chemistry for the detection of the amplicons in a real-time PCR instrument, instead of expensive fluorophore-labeled probes, decreases the cost of the assay to an estimated $0.27 per genotype. Recently, new techniques based on the analysis of the melting profile of the amplicon have been developed for single-tube genotyping, using SYBR Green I chemistry in a real-time instrument. However, most of them require post-PCR manipulation (17) (increasing the risk of cross contamination) or a time-consuming data handling process (11). Germer and Higuchi (7) proposed a protocol for SNP frequency studies, based on ARMS, followed by a melting profile of the amplicons, but it requires a demanding optimization process, which our technique does not.

Sequencing data confirmed the results obtained with our HP assay. Using a 96-well format, we were able to test the 264 samples in only 6 PCR runs (each of which lasts 2 h). These results confirm the potential of the HP assay for medium- to large-scale association studies of complex human diseases.

We applied this technique to study whether there was an association between the C→G transversion at position –44 in the DEFB1 gene and EOP. The results do not support a direct association of this SNP with EOP. It is possible that other SNPs in the DEFB1 gene could be associated with EOP. However, there are no reported associations of other polymorphisms in the DEFB1 gene with an increased risk of infection, with the exception of a nonsynonymous mutation, associated with chronic obstructive pulmonary disease, that has been reported only in a Japanese population (19). It is also possible that human beta-defensin 1 is not active against periodontal bacteria and does not play a role in this disease. Supporting this hypothesis, it has been demonstrated that Treponema denticola is resistant to human beta-defensin 1 (2). Unfortunately, no data are available regarding the biological activity of this peptide against Actinobacillus actinomycetemcomitans and Porphyromonas gingivalis, two key oral pathogens implicated in periodontal disease.

In conclusion, we have designed and optimized a simple, reliable assay for examining SNP frequencies that provides several major advantages over traditional SNP-detecting techniques. We have successfully applied this technique to examine, for the first time, the associations between the SNP at position –44 in the DEFB1 gene and EOP.

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