Usefulness of Nucleic Acids (DNA/RNA) from Buccal Cells Isolated from Mouthwashes using a Modified Method

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Abstract

Human buccal cells are an easy source of nucleic acids (DNA/RNA) other than human blood. We have extracted buccal cells from mouthwashes of 10 individuals and tested its usefulness for genotyping, telomere length determination and gene expression. Genomic DNA was extracted using a salt precipitation method whereas total RNA was extracted using Hipura total RNA extraction kit. The quantity and quality of DNA and RNA was found to be satisfactory. PCR amplification was achieved using locus specific primers and genotyping was performed for minisatellites, microsatellites and Single nucleotide polymorphisms (SNPs). Telomere length was determined from the genomic DNA of buccal cells obtained from mouthwashes. Telomere length obtained from the DNA isolated from buccal cells and peripheral blood mononuclear cells was compared. Real time quantitative PCR was successfully performed for GADD45A, G6PD, B-Actin and B2M. In conclusion, our method of DNA /RNA extraction can be considered as an alternative method of sample collection for forensic medicine.

Keywords: Forensic science; Buccal cells; DNA; RNA; Genotyping; Gene expression; Telomere length

Introduction

Isolation of Nucleic Acids from human peripheral blood samples serves as the most important source in studies pertaining to epidemiology, forensics and biomedical/molecular genetics. However, genotyping is sometimes difficult if the blood sample from an individual is not available. In that circumstance, the most preferred source of nucleic acids is buccal cells from where nucleic acids (DNA/RNA) can be extracted easily. Several studies have been undertaken using buccal cells as a source of nucleic acids. DNA obtained from buccal cells has been used for genetic studies [1-10] and cancer associated changes in oral cavity [11]. Total RNA obtained from buccal cells and its usefulness for gene expression studies is comparatively novel. Similarly telomere length determination from DNA from buccal cells is relatively new to population genetic and forensic medicine.

For epidemiological and forensic studies the ideal method of choice for genomic DNA or RNA should be rapid, non-invasive and with a good yield and quality of nucleic acids. Most of the methods use organic solvents for the procedure which are difficult to handle and are hazardous. We have collected buccal cells which is suitable for large scale population monitoring programme, genetic and genotoxicity studies. In the present study, we have introduced a simple non-enzymatic salt precipitation method for DNA extraction which can be useful for genotyping as well as telomere length determination.

This protocol gave high yield and good quality DNA which was be used for PCR amplification, quick to collect, requires no abstinence from food or drink. This process is devoid of using any enzymatic or phenol/chloroform steps for DNA isolation. We have used salt solution as mouthwashes in order to collect buccal cells for our study which can be easily handled for large scale population studies. Similarly attempts have been made to extract RNA from buccal cells which can be useful for gene expression study using total RNA extracted from buccal cells.

Materials and Methods

Collection of buccal cells from mouthwashes

We have introduced saline rinse method to collect the buccal cells from random donors. In this procedure, buccal cells were collected from individuals by rinsing the mouth in the morning with freshly prepared 0.9% of saline (NaCl) solution. Individuals were asked to rinse mouth with freshly prepared 15 ml (0.9% NaCl) saline vigorously at least for a minute. Then it was expectorated into a 50 ml collection tube. A total of 10 volunteers were participated in this study where mouth wash samples were collected from them in the morning with the above procedure.

Isolation of nucleic acids

Extraction of DNA: Genomic DNA was isolated from mouthwash buccal samples of 10 individuals. The mouthwashes collected in 15 ml centrifuge tubes were directly centrifuged for 5 minutes at 2500 rpm. Cell pellet from each sample tube was collected and the supernatant was discarded. The pellet was resuspended in 5ml of low salt solution (TKM1: 10 mM Tris-HCl, pH 7.6, 10 mM MgCl₂, 2 mM EDTA, pH 8, and 10 mM KCl) and 125 µl of NP-40 (Non-iodet P40, Roche Diagnostics GmbH, Germany) and mixed thoroughly. It was then centrifuged at 800g (2200 rpm) for 10 minutes (Remi RBC laboratory centrifuge, Mumbai, India). The pellet was resuspended in 5 ml of the TKM1 buffer and centrifuged for 10 minutes at room temperature at 2200 rpm. The pellet was resuspended in 0.8 ml of high salt buffer, TCM2 (10 mM Tris-HCl (pH 7.6), 10 mM MgCl₂, 2 mM EDTA (pH 8), and 10 mM KCl, 0.4M NaCl) and 125µl of 10% SDS was added to it. The whole suspension was mixed thoroughly by pipetting back and forth several times and was incubated at 55°C until clear solution appears. To this solution 0.3 ml of saturated salt solution (6 M NaCl)

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was added and mixed well in order to precipitate the proteins. The tubes were centrifuged at 12,000 rpm for 20 minutes at 4°C. The supernatant (having nucleic acid) was distributed equally into two 2 ml centrifuge tubes and the pellet was discarded. Twice the volume of chilled absolute ethanol was added to the supernatant and mixed well to precipitate the DNA. Precipitated genomic DNA was pelleted by centrifuging at 12,000 rpm at 4°C for 20 minutes. The pellet was washed three times using 200 µl chilled 70% ethanol and centrifuging it at 12,000 rpm at 4°C for 15 minutes each. The pellet containing genomic DNA was dried by keeping it at 55°C for about 20 minutes. The dried pellet was resuspended in 75 µl sterile distilled water.

DNA was quantified by taking absorbance (A260 and A280) using 'Picodrop Microlitre Spectrophotometer. Purity and the concentration of isolated DNA were calculated. The ratio of absorbance at 260 and 280 nm was also obtained to determine purity of DNA and it was observed to be between 1.8 and 2.0. DNA was also subjected to electrophoresis using 1% agarose for qualitative analysis.

Amplified fragment length analysis (Amp-FLP)

Amplified Length polymorphism was carried out for three minisatelites and three microsatellites. Genomic DNA obtained from buccal cells was PCR amplified using locus specific primers of four microsatellites (D10S1412, D3S2459, Leptin and APOB tetranucleotide repeats) and three minisatelites (APOB VNTR, D1S80, D19S20). PCR amplification was achieved using 25 ng of genomic DNA and the amplimers were subjected to restriction fragment length analysis using restriction endonucleases. The restriction enzymes used for XRCC3, MTHFR (C-T), and MTHFR (A-G) were NcoI, HinfI, TaqI for overnight digestion. The digested products were run on 6% PAGE gel images of three minisatellites (APOB VNTR, D1S80 and D19S20), single nucleotide polymorphisms. The silver stained polyacrylamide gel Electrophoresis (PAGE) respectively for mini- and microsatellites. The annealing temperatures for the microsatellites: D10S1412, APOB tetra, Leptin and D3S2459 were 58°C, 56°C, 56°C, respectively. The melting and extension step for all the PCR reactions were 95°C and 72°C, respectively. A total volume of 25 µl PCR reaction was set up for both microsatellites and minisatelites. The PCR cocktail mixture contained 10 mM of dNTPs, 0.5 U Taq polymerase, 10 pmol of PCR primers each for all the loci studied.

Single Nucleotide Polymorphism (SNP) analysis

The genomic DNA isolated from buccal cells using our method was also checked if they could be used for SNP analysis. Three polymorphic loci were studied are XRCC3 (Thr/Met), MTHFR (C-T) and MTHFR (A-G). PCR amplification of genomic DNA was performed using locus specific primers and then subjected to restriction fragment length analysis using restriction endonucleases. The restriction enzymes used for XRCC3, MTHFR (C-T), and MTHFR (A-G) were NcoI, HinfI, and TaqI respectively. The restriction digestion mixture contained 5µl of PCR product, 1 Unit of restriction enzyme each and kept at 37°C for overnight digestion. The digested products were run on 6% PAGE followed by silver staining.

Determination of Telomere Length using real time PCR

Telomere length was determined using DNA obtained from buccal cells for 10 individuals using SYBR green based real time PCR approach. This method measures telomere length (T) with respect to a single copy gene (36B4). Telomere length in buccal cells and blood cells (leukocyte telomere length) was also compared in these individuals. For that purpose geneomic DNA was also extracted from peripheral blood mononuclear cells of these individuals. Genomic DNA (25 ng) was amplified using a telomeric specific primer and a single copy gene primer as described by Cawthon et al. [12]. This method measures the factor by which the ratio of telomere repeat copy number to single gene copy number differs between a sample and that of a reference DNA sample. PCR amplification was achieved using telomere (T) and single copy gene, 36B4 (encodes acidic ribosomal phosphoprotein) primers(S) which serves as a quantitative control. The mean telomere repeat gene sequence (T) to a reference single copy gene (S) is represented as T/S ratio which is calculated to determine the telomere length.

The PCR mixture contained 5pmoles of each of the primers, 100 µM of each dNTPs and 0.3 X SYBR green dye and 0.5 Units of Faststart taq DNA polymerase (Roche Diagnostics, GmbH, Germany). PCR reactions were performed in 20 ml reaction volumes using 25 ng DNA sample per reaction. The PCR reactions were performed using telomere and single copy gene primers in the same 96 well plate. The PCR thermal conditions for relative telomere length assay consisted of a initial denaturation of 5 minutes at 95°C, followed by a total of 40 cycles at 95°C for 30 seconds, 58°C for 30 seconds, and 72°C for 30 seconds and fluorescence acquisition.

Analysis of gene expression

Total RNA isolation, cDNA Preparation and quantitative real time PCR: Total RNA was also isolated from the cell pellet collected from the mouthwashes of these individuals using HipurA total RNA miniprep purification spin kit (Himedia Pvt ltd, Mumbai, India). The ratio of absorbance at 260 and 280 nm was also obtained to determine purity. Total RNA was quantified by taking absorbance (A260 and A280) using 'Picodrop Microlitre Spectrophotometer, (Version 3.01). The concentration was calculated. Amount of RNA obtained was between 0.30µg to 1.94 µg per mouth wash collected.

cDNA was synthesized from 0.5 µg total RNA using Transcriptor high fidelity cDNA synthesis kit (Roche diagnostics, GmbH, Germany). PCR amplification of the synthesized cDNA was carried out using LC480 (Roche diagnostics, GmbH, Germany). G6PD, GADD45A, B-Actin & B2M genes were amplified. All the genes gave a single specific product. This was checked by analyzing the melting curve as well as on a 6% silver stained polyacrylamide gel.

Results

High molecular weight DNA was isolated from the mouthwashes of 10 random and healthy volunteers using non-invasive, non-enzymatic and simple cost effective procedure. The average DNA obtained from buccal cells after mouth washes among these individuals studied were in the range of 3.5 µg-146 µg. The quality of DNA was good where the ratio of 260 nm and 280 nm was found to range from 1.8 to 2.0. Similarly the average RNA obtained from buccal cells of these individuals was ranging from 0.30 to 1.94 µg with a ratio of 260 nm to 280 nm to be around 2.0.

The DNA obtained from buccal cells was successfully used for locus specific PCR amplification of minisatelites, microsatellites and single nucleotide polymorphisms. The silver stained polyacrylamide gel images of three minisatelites (APOB VNTR, D1S80 and D19S20), four microsatellites (Leptin, ApoB, D10S1412 and D3S2459) and three Single nucleotide polymorphisms (XRCC3(C-T), MTHFR (C-T), and MTHFR (A-G)) were shown in Figures 1-3. These polymorphic loci were amplified with 100% efficiency. For Single nucleotide polymorphisms, DNA was treated with restriction enzymes such as NcoI, HinfI, TaqI for XRCC3(C-T), MTHFR (C-T), and MTHFR (A-G) SNPS respectively. All the amplimers were successfully digested using restriction endonucleases and was run on native polyacrylamide gel.
Telomere length is associated with respect to age, stress, other age related diseases such as hypertension and diabetes. We have measured telomere length using DNA obtained from buccal cells of mouth washes. For that purpose, relative quantitation was performed using a SYBR green based real time PCR approach. Telomere specific and single copy gene specific primers were used to amplify DNA. For each DNA sample the factor by which the sample differed from a reference DNA sample from its ratio of telomere repeat copy number to single gene copy number was measured. The mean telomere repeat gene sequence (T) to a reference single copy gene (S) is represented as T/S ratio which is calculated to determine the telomere length. Beta-globin gene was used as a positive control. The mean telomere length calculated was 1.21 ± 0.08. It was also compared with the telomere length measured from the DNA extracted from blood lymphocytes. The telomere length obtained from the DNA of both the sources did not show any significant difference in telomere length (Figure 4). The buccal cell and lymphocyte was isolated from the same individuals for comparison so that there was no error due to inter individual variation.

Gene expression study was carried out with the total RNA obtained from buccal cells. Total RNA was isolated and converted to cDNA using transcriptor high fidelity kit (Roche Diagnostics, GmBH, Germany). Gene expression analysis was carried out for cell cycle growth arrest gene (GADD45A). Three housekeeping genes (B2M, B-actin, and G6PD) were also taken into consideration to check the quality. Relative
The yield and quality was similar to other studies conducted by different investigators [7,13]. The quality of DNA obtained for all the samples showed high purity. All the samples were PCR amplified with 100% efficiency. The DNA obtained from buccal cells was successfully used for genetic polymorphism studies. It was successful for typing minisatellites, microsatellites and short tandem repeat polymorphism.

Telomeres are ends of eukaryotic chromosomes which are biomarkers of aging. The length of telomere decreases with increased oxidative stress. Telomere length was measured in buccal cells as well as peripheral blood mono-nuclear cells for a comparison. As shown in Figure 5, the telomere length is comparable. Hence buccal cells can be used for telomere length determination in forensic medicine as well as population genetics. Moreover, processing of buccal cells for nucleic acid isolation is a simple, inexpensive, non-invasive and could be used for large scale population monitoring programme.

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