Improved protocol for extracting genomic DNA from frozen formalin-fixed tissue resulting in high-quality whole mtDNA

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Received October 30, 2015
Accepted January 15, 2015
Published August 26, 2016
DOI http://dx.doi.org/10.4238/gmr.15037972

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ABSTRACT. Formalin fixation and paraffin embedding is widely used for convenient and long-term storage of tumor tissue and precious sources to perform genetic studies. However, DNA fragmentation is one of the major flaws of genomic DNA isolation from formalin fixation tissues, which limits its further usage. Here, we present an improved method for isolating high-quality genomic DNA from formalin fixation tissue. We obtained high-quality genomic DNA of more than 20 kb
from samples frozen for more than 2 years. Furthermore, to verify DNA quality, the whole mitochondrial DNA (mtDNA) genomes from the normal and tumor tissue of the same patient were successfully amplified with two overlapping PCR fragments comprising more than 8379 bp in length for each fragment. In addition, the whole genomes were sequenced with a 48-well based primer panel in order to avoid potential sequencing errors from artificial recombination, which was further confirmed with an mtDNA phylogenetic strategy. Our improved DNA extraction method from formalin fixation tissue and sequencing strategy for entire mtDNA genomes will generate unambiguous sequence analysis results for clinical samples.

**Key words:** Formalin-fixed tissue; DNA isolation; Entire mtDNA genome

**INTRODUCTION**

Formalin fixation is commonly used for preservation of pathological tissue and biological specimens. Formalin-fixed paraffin-embedded (FFPE) tissue contains a rich source of genetic material that can be used for medical research (Lipson et al., 2012; Chen et al., 2013) and biological disciplines (Santos et al., 2009) in downstream DNA-based experiments, such as copy number analysis (Gilbert et al., 2007; Jacobs et al., 2007), real-time quantitative polymerase chain reaction (RT-PCR) (Gjerdrum et al., 2004), single nucleotide polymorphism (SNP) bead arrays (Oosting et al., 2007), microarray comparative genomic hybridization (Vékony et al., 2007), and genome-wide sequencing (Aviel-Ronen et al., 2006; Schumer et al., 2013). However, formalin may cause the fragmentation of DNA (Douglas and Rogers, 1998), which reduces the quality and quantity of DNA (Poljak et al., 2000). Isolation of sufficient amounts of intact DNA from FFPE tissue is a challenge due to crosslinking between DNA and protein caused by formalin fixation, which decreases the quality of most DNA molecules by breaking them into 200 bp or less fragments (Gilbert et al., 2007). A number of approaches have been published to obtain higher quality DNA from FFPE tissue; however, very high-quality DNA has never been isolated (Farrugia et al., 2010; Okello et al., 2010). Therefore, in view of the invaluable resources contained in FFPE tissue on studying pathogenesis of cancer and a variety of other diseases, exploring an effective method to preserve and purify higher quality genomic DNA from FFPE tissue is imperative.

One of the critical steps for obtaining high-quality genomic DNA is to break up the crosslinks between DNA and protein during the extraction process (Srinivasan et al., 2002; Gilbert et al., 2007). Fortunately, the formation of these crosslinking is reversible in aqueous solution (Dubeau et al., 1986; Fang et al., 2002). Thus, the possibility of obtaining high-quality genomic DNA from FFPE tissue was presented by performing a pre-extraction procedure to remove paraffin from fixed tissues (Santos et al., 2009). Previous study has confirmed the superiority of genomic DNA after pretreatment of the FFPE tissues (Fang et al., 2002).

Mitochondria, the power source of the cell, are the major cellular component performing oxidative phosphorylation to supply energy to cells, and participate in complex apoptotic signaling pathways (Lee et al., 2004). Mitochondria also have their own genomic DNA and, since they lack the protection of histones, mitochondrial DNA (mtDNA) are more vulnerable
to oxidative damage. They also have poor DNA mismatch repair mechanisms (Croteau and Bohr, 1997), resulting in genome variations in mtDNA that are 10 to 200 times more frequent than those in nuclear genomic DNA (Larsen et al., 2005). Previous study indicates that mtDNA mutations are associated with various diseases (Schon et al., 2012), such as Leber hereditary optic neuropathy (Wallace et al., 1988), Leigh syndrome (de Vries et al., 1993), mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes (Kirby et al., 2004). The somatic mutations detected in different tumor tissues may also relate to the process of cancer (Schon et al., 2012). As DNA fragmentation is the major shortcoming of FFPE tissues (Gilbert et al., 2007), the role of mtDNA mutations in cancer has been difficult to understand, partly due to the artificial recombination that occurs during processing of small PCR segments (Fendt et al., 2009). Considering the fact that formalin fixation is one of the most commonly used preservation methods for tumor tissues, it is important to be able to isolate intact genomic DNA and sequence the entire mtDNA genome in order to avoid artificial recombination.

In this study, we developed an improved protocol to extract higher quality genomic DNA from frozen formalin fixation colorectal cancer tissues that were frozen for more than 2 years. This protocol uses gradient ethanol to remove the formalin by reversing the crosslinks with genomic DNA in aqueous solution, and takes advantage of the good solubility of formaldehyde in water and ethanol (Fang et al., 2002). We were able to obtain 30 to 50 μg high-quality total genomic DNA for each sample, the quality of which was verified by absorbance measurements and gel electrophoresis. Furthermore, two overlapping PCR fragments comprising over 8379 bp in length were successfully amplified (Fendt et al., 2009) and high-quality genomic mtDNA was obtained according to our improved sequencing strategy.

**MATERIAL AND METHODS**

**Tissue specimens**

To perform the protocol for isolating genomic DNA from formalin fixation tissues, a total of 6 tissues from 2 colorectal cancer patients were collected randomly from our biological specimens’ bank, including the primary colorectal cancerous tissue, two symmetrical adjacent cancerous tissues, and distant normal tissue, which were at least 5 cm away from the edge of the cancerous tissue. The tumor tissues were collected during surgery with consent and personal information was omitted during the process of data analysis and reporting. All procedures were supervised and approved by the Human Tissue Research Committee of the First People’s Hospital of Yunnan Province. The study was performed according to the Declaration of Helsinki. The tumor tissues were verified with hematoxylin and eosin staining and the para-cancerous tissues were obtained from the symmetrical side of the cancer. The primary colorectal cancerous tissues and corresponding para-cancerous normal tissues were marked with C and CP, respectively (Wang et al., 2011), fixed with an excess volume of 10% (v/v) formalin, and then stored at -80°C.

**DNA extraction**

Approximately 30 mg frozen tissue was cut into small pieces and collected into a 1.5-mL centrifuge tube. To reverse the crosslinking between DNA and protein, the formalin fixation tissues were initially soaked in 100% ethanol for 10 min then centrifuged at 12,000
rpm for 10 min. This was followed by successive washes with gradient ethanol (90, 85, 80, 70, and 50%) and sterile water. After the final rehydration, the tissue was cut and ground with an electric mill (Tiangen Biotech Co., Ltd., Beijing, China). The suspension was digested with 10 µg Proteinase K and 100 µL SDS buffer (10%) for at least 16 h at 56°C, and then mixed with 700 µL Tris-saturated phenol for at least 12 h. This was followed with the standard phenol/chloroform method (Gross-Bellard et al., 1973) and the final genomic DNA was dissolved in 100 µL TE buffer. The quality and integrity of genomic DNA was tested by gel electrophoresis on 1% agarose gels and compared to a 23-kbp genomic DNA standard. The quantity and quality of the genomic DNA was measured with a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA) by calculating the ratio of absorbance readings at 260 and 280 nm (A260/280).

**PCR amplification, sequencing, and quality control**

As the quality and quantity of genomic DNA extracted from FFPE tissues is poor (Gilbert et al. 2007), long fragments of PCR for mtDNA genomes were amplified and sequenced as two fragments comprising more than 8379 bp in length for each fragment (Fendt et al., 2009). The entire mtDNA genomes of the two colorectal cancer patients were sequenced with 48 internal sequencing primers. The sequencing reactions were performed with the BigDye Terminator v3.1 cycle sequencing kit (Applied Biosystems, Foster City, CA, USA) on a 3700 DNA Analyzer (Applied Biosystems). Original sequences were edited and aligned with the DNASTAR software package version 7.0 (DNASTar Inc., Madison, WI, USA). Mutations in the mtDNA genome were scored according to the revised Cambridge Reference Sequence (rCRS) (Andrews et al., 1999). The phylogenetic status of the mtDNA genomes was determined according to the reconstructed East Asian mtDNA phylogeny (Kong et al., 2006).

To avoid sequencing errors prevalent in the medical field (Bandelt et al., 2007), strict quality control measures were followed in our procedure upon extracting genomic DNA and generating the mtDNA genome sequences. First, each sample was given a unique sample identification (ID) number and all samples were amplified and sequenced according to their sample IDs, which were verified at each step. Second, the entire mtDNA genome was amplified as two segments of more than 8379 bp each (Fendt et al., 2009), which can efficiently reduce the risk of artificial recombination caused by sample crossover and help to avoid the amplification of pseudo-mitochondrial genes from the nuclear genome. Third, a total of 48 internal primers were used for sequencing the entire mtDNA genome. These sequencing reactions were performed on the same 96-well plate, which can effectively reduce the potential for artificial recombination during the sequencing process. Lastly, variations were recorded and inspected by different experienced analysts and compared with the rCRS (Andrews et al., 1999). All suspicious variations were identified by a phylogenetic approach (Bandelt et al., 2007).

**RESULTS**

In order to isolate intact genomic DNA from frozen tissue for further genetic analysis, frozen tissue was pretreated to remove paraffin. This was done by first soaking the tissue in gradient ethanol and deionized water then cutting and grinding the tissue. The genomic DNA was further purified with a standard phenol/chloroform method. The quantity and quality of
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the genomic DNA were evaluated with traditional agarose gel electrophoresis analysis and NanoDrop, and universal and long fragment PCR were performed and sequenced with our improved strategy. In addition, phylogenetic analysis for the whole mtDNA genomes of the two colorectal cancer patients was performed by comparing them to published mtDNA genomes (Wang et al., 2011).

To evaluate the integrity of the genomic DNA isolated from frozen formalin fixation samples, the genomic DNAs was analyzed by 1% agarose gel electrophoresis, as shown in Figure S1a. The intact genomic DNA, which was approximately 23 kbp and extracted from the frozen formalin fixation tissues, was the same size as that isolated from the leukocytes of fresh blood (Fang et al., 2002). Our results indicate that the improved method can effectively isolate intact genomic DNA from frozen formalin fixation tissues (Figure S1a).

The quality and quantity of genomic DNA extracted from the frozen formalin fixation samples were evaluated. The total DNA was diluted 10-fold and the concentration of the diluted DNA was measured with a NanoDrop-1000. We obtained a high yield of genomic DNA from the 4 tissues, varying from 32.8 to 65.4 μg (Table S1). The purity of the genomic DNA was assessed by the ratios of A$_{260/280}$. As shown in Table S1, two genomic DNA samples had ratios of A$_{260/280}$ between 1.8 and 2.0 and the other two samples had ratios of A$_{260/280}$ slightly higher than 2.0. A$_{260/280}$ values between 1.8 and 2.0 indicate an almost protein-free DNA samples while a ratio of A$_{260/280}$ higher than 2.0 indicates the presence of contaminating RNA, which absorbs strongly at 260 nm.

The quality of the genomic DNA isolated from the frozen formalin fixation tissues was also evaluated in terms of its successful PCR amplification and sequencing results (Fang et al., 2002; Santos et al., 2009). The whole mtDNA control region (around 1589 bp) was amplified with 1 pair of primers (Zhao et al., 2009), and the whole mtDNA genome (around 16,569 bp) was amplified with two covering fragments, which were 8379 and 8604 bp long (Fendt et al., 2009) (Figure S1b and S1c). We successfully obtained all PCR fragments with 50 ng genomic DNA, which was isolated from the frozen formalin fixation colorectal cancer tissue. The entire mtDNA genome was sequenced with 48 internal primers, which were arranged on the upper and lower parts of a 96-well plate for the two patients, respectively. This strategy avoids possible artificial recombination during the sequencing process. The primer sequences of the 48 internal primers were taken from the literature (Wang et al., 2008; Fendt et al., 2009). The mtDNA genomic sequences of the two patients were both allocated to haplogroup F1 for sharing a string of diagnostic variations (Figure 1). To further confirm the reliability of the two genomes, 12 whole mtDNA genomes were retrieved from more than 22,514 whole mtDNA genomes. As shown in Figure 1, the two whole mtDNA genomes have six and three private variations, which only appeared in the two new sequenced mtDNA genomes by comparing reported sequences of the same haplogroup. These were identified by comparing the genomes sharing much more basal variations with the sequence obtained in this study. In detail, patient 7B has four control region variations (including variations 151, 309+C, 315+C, and 16193+C) and two coding region variations (containing variations 9272 and 15908). Patient 12B has two control region variations (including variations 143 and 315+C) and one coding region deletion (4314d). Furthermore, by screening these mutations among the 22,514 whole mtDNA genomes, we found that mutation 9272 was prevalent among African mtDNA haplogroup L3 and European mtDNA haplogroup J. Mutation 15908 was one of defining variations of mtDNA haplogroup M33a and M74b, and it also appeared in individuals of F4a. The adenine deletion was widely distributed among individuals of mtDNA haplogroup F1e2, and the
control region variations 143, 151, 309+C, 315+C, and 16193+C were hotspot mutations in mtDNA genomes, which imply that these mutations occurred randomly.

**DISCUSSION**

FFPE tissue is a broadly used method to preserve tumor tissue because it allows for stable storage conditions for the specimen, which is a valuable biological resource for studying genetic-based disease (Sengüven et al., 2014). However, the crosslinks between DNA and protein result in the degradation of genomic DNA, limiting its usage in genetic studies (Gilbert et al., 2007). In this study, the method for isolating genomic DNA from formalin fixation tissues was improved to achieve higher quality and quantity of DNA. This was primarily done by displacing the formalin from the tissues with a decreasing ethanol gradient and deionized water. We obtained genomic DNA that was 23 kbp (Figure S1a), which was identical in size
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To DNA isolated from the leukocytes of fresh blood. In addition, the yield (A$_{260/280}$) of genomic DNA isolated from frozen formalin fixation tissue was better than that from blood leukocytes (Table S1). Our results indicate that removing the formalin from the formalin fixation tissues before isolating genomic DNA is a very important step (Fang et al., 2002).

The quality and quantity of the genomic DNA were evaluated by amplifying and sequencing fragments of 1589, 8379, and 8604 bp (Figure S1b and S1c). The latter two fragments covered the whole mtDNA genome. Our phylogenetic analysis supported the F1 haplogroup status of the two patients, F1g and F1d1. By comparing with 12 individuals sharing the identical mtDNA haplogroup status, which were retrieved from 22,514 whole mtDNA genomes, we found that most of the private variations were detected in the mtDNA control region. Therein, we found that heterogeneity was the main character of the variations, especially the C-tract around positions 310 and 16189, which varied in length. These two variations were also detected in patients with breast cancer, lung cancer, and type 2 diabetes (Alhomidi et al., 2013; Chen et al., 2016). Our results confirmed the reliability of the two whole mtDNA genomes of the patients with colorectal cancer, which implied that our sequencing strategy could reduce the risk of artificial recombination efficiently (Fendt et al., 2009; Bi et al., 2011). The complex purification process, which may enhance the opportunity for DNA contamination, was also verified based on the sequencing results of the non-recombinant whole mtDNA genomes. No other mutations belonging to different haplogroups were detected, thus further confirming the high quality of our protocol.

By comparing our DNA isolation strategy with those described in previous studies (Lin et al., 2009; Santos et al., 2009; Farrugia et al., 2010; Okello et al., 2010), our results indicate that the method of stocking frozen formalin fixation tissues for long periods of time is the major factor affecting the quality of genomic DNA. The pretreatment of the formalin fixation tissues by gradual dehydration can effectively reduce the crosslinking between histone proteins and DNA, even though formaldehyde cannot be completely removed (Fang et al., 2002). However, the proper ratios of A$_{260/280}$ and successful amplification of DNA fragments more than 8379 bp in length support that triturating the pretreated tissues and digesting the suspensions with Proteinase K can effectively reduce crosslinking. The improved sequencing strategy and the analysis based on mtDNA phylogeny can generate high-quality sequences. In addition, this strategy can rapidly identify private mutations in patients with colorectal cancer, as described in previous studies (Bi et al., 2011). However, the pretreatment of tissue was time consuming, and the complex process may enhance the chances of DNA contamination.

In this study, we describe an effective protocol for isolating high-quality genomic DNA from frozen formalin fixation tissue. Amplifying whole mtDNA genomes with long PCR fragments, sequencing the whole genomes with our improved strategy, and analyzing the variation based on mtDNA phylogeny can generate higher quality sequences and identify private mutations rapidly. This will benefit future studies on investigating variation in mtDNA genomes in clinical fields.

Conflicts of interest

The authors declare no conflict of interest.

ACKNOWLEDGMENTS

Research supported by the Foundation for Science and Technology Planning...
REFERENCES


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Supplementary material

**Figure S1.** a. Represents the electrophoretogram of total genomic DNA; b. shows the PCR fragment of the mtDNA control region covering 1569 bp; and c. denotes the PCR fragments of 8379 bp (A) and 8604 bp (B), which covered the whole mtDNA genome. In (a) and (c), the molecular weight marker was 23 kbp and in (b), the molecular weight marker was the DL2000 DNA marker.

**Table S1.** A<sub>A260/A280</sub> ratios of genomic DNA isolated from two formalin-fixed tissues.