Objective: To date, there are only few systematic reports on the quality of DNA extracted from routine diagnostic cytologic specimens. It was the aim of the present study to evaluate the ability of 50% ethanol/2% carbowax (Saccomanno fixative) to preserve bronchial secretions with high quality genomic DNA as well as to compare different DNA extraction methods.

Methods: DNA was extracted from 45 bronchial aspirates by four different extraction protocols. Beside DNA yield, DNA quality with regard to purity, integrity, and PCR success rate were investigated.

Results: No fragmentation of sample DNA due to the fixative was detected. It was preserved as high molecular weight DNA. DNA yield, purity, and integrity were dependent on the DNA extraction method to some extend. Irrespective of the DNA extraction method the PCR success rate for amplification of β-globin gene fragments (268, 536, and 989 bp) was 100%.

Conclusions: A fixative containing 50% ethanol/2% carbowax preserves high quality DNA which is well suited for PCR-based assays regardless of the extraction protocol used. The selection of the DNA extraction protocol has to be adjusted to the circumstances of application.

Keywords: Molecular cytology, fixative, PCR, bronchi, DNA extraction

1. Introduction

The application of molecular biological methods to bronchial aspirates for the diagnostic detection and the investigation of cancer cells is a challenge. Bronchial aspirates contain only a low total amount of DNA as compared with tumor tissue obtained by surgery. In addition, usually only a small portion of total genomic DNA actually represents tumor cell DNA. Thus, obtaining high quality genomic DNA is critical for molecular cytologic studies on bronchial aspirates. It is well known from studies using tissue specimens that the quality of genomic DNA is mainly influenced by the fixative and the method of DNA extraction applied [2,3]. However, regarding routine diagnostic cytologic specimens there are only few systematic reports on this topic [1,5,9].

This study investigates the ability of Saccomanno fixative (50% ethanol/2% polyethylene glycol [carbowax]) to preserve high quality genomic DNA since this fixative is often used in routine diagnostic cytopathology of the respiratory tract. In addition, the study compares phenol–chloroform DNA extraction to the following alternative methods: (a) NucleoSpin Tissue (Macherey-Nagel, Düren, Germany); (b) Puregene DNA Isolation Kit (Gentra Systems, Minneapolis, MN); (c) QIAamp DNA Mini Kit (Qiagen, Hilden, Germany).
samples; (b) no purulent and clotty material; (c) evaluated as tumor cell negative, as unevenly distributed clusters of hyperdiploid tumor cells might distort the results. Thus, a homogeneous distribution of nuclear DNA in each split sample was ensured. The age of the patients ranged between 32 and 80 (median age, 62). Twenty-one of the individuals were females, and 24 were males.

After bronchoscopy, the specimens were immediately transferred to 50-ml conical tubes containing 20 ml of Saccomanno fixative. On arrival at the laboratory, all samples were primarily processed for routine diagnostic purposes [6]. They were centrifuged at 670 \( \times g \) for 5 min. A part of the cell pellet was used to prepare conventional smears. They were subsequently stained according to Papanicolaou.

For this study conventional smears were reviewed to examine cellular preservation and bacterial growth as well as to determine the degree of inflammation. A mild, moderate, or severe inflammation was assigned when inflammatory cells (neutrophilic granulocytes, lymphocytes) were scanty, numerous, or predominating respectively.

Residual material was then centrifuged again, the supernatant was decanted, the cell pellet was resuspended in 800 \( \mu l \) of Saccomanno fixative and split into four 1.5 ml tubes. The sample DNA was then extracted using the following methods (see Appendix for detailed protocols): (a) phenol–chloroform extraction; (b) NucleoSpin Tissue; (c) Puregene DNA Isolation Kit; (d) QiAamp DNA Mini Kit.

The DNA concentration of each split sample was quantified using UV-spectrophotometry at 260 nm. The DNA yield per aliquot was calculated by multiplying the DNA concentration by the final volume of DNA extract. DNA purity was estimated as ratio of \( \frac{A_{260}}{A_{280}} \).

The integrity of genomic DNA was assessed by electrophoresis of DNA extracts on 0.3% agarose gels. DNA was visualized by ethidium bromide staining. Each gel was documented with the BioDocAnalyze (Biometra, Göttingen, Germany). The semi-automatic image analysis software of BioDocAnalyze allows measuring the relative migration (RF-value) of sample-DNA and 1 kb DNA ladder (1 kb DNA Extension Ladder, Invitrogen, USA). Based on the resulting RF-values it was possible to calculate the size of the DNA fragments.

The adequacy of the DNA extracts for PCR-based assays was assessed by amplifying three different sized fragments of the \( \beta \)-globin gene (268, 536, and 989 bp) as described by Greer et al. [3]. Amplicons were separated on 1.5% agarose gels by electrophoresis and visualized by ethidium bromide staining.

For statistical analysis the differences between methods of DNA extraction in DNA concentration, DNA purity, and main DNA fragment size were tested with the Wilcoxon matched-pair signed-rank test for paired observations.

### 3. Results

#### 3.1. Light microscopy

Light microscopic review of conventional cytologic smears showed an excellent cellular preservation and no relevant bacterial growth in all cases. The latter was restricted mostly to some oral squamous epithelia. Mild, moderate, and severe bronchitis was present in 10, 43, and 2 cases, respectively.

#### 3.2. Saccomanno fixative

Table 1 shows that Saccomanno fixative preserves DNA of bronchial aspirates very well and allows preparation of high molecular-weight DNA. This is

<table>
<thead>
<tr>
<th>DNA extraction method</th>
<th>DNA yield ( [\mu g] )</th>
<th>DNA purity ( \frac{A_{260}}{A_{280}} )</th>
<th>Fragment size ( [kb] )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>median (min; max)</td>
<td>mean ( \pm SD )</td>
<td>median (min; max)</td>
</tr>
<tr>
<td>Phenol–chloroform</td>
<td>2.76 (0.78; 55.60)</td>
<td>1.69 ( \pm 0.16^* )</td>
<td>( &gt;40 ) (( &gt;40 ); ( &gt;40 ))</td>
</tr>
<tr>
<td>NucleoSpin Tissue</td>
<td>1.44 (0.18; 29.28)*</td>
<td>1.70 ( \pm 0.18^* )</td>
<td>26 (13; ( &gt;40 ))</td>
</tr>
<tr>
<td>Puregene</td>
<td>1.53 (0.24; 30.90)</td>
<td>1.77 ( \pm 0.16 )</td>
<td>( &gt;40 ) (17; ( &gt;40 ))</td>
</tr>
<tr>
<td>QiAamp DNA Mini Kit</td>
<td>1.14 (0.24; 18.90)*</td>
<td>1.83 ( \pm 0.12 )</td>
<td>23 (13; ( &gt;40 ))</td>
</tr>
</tbody>
</table>

*Difference not significant, Wilcoxon’s rank sum test.
documented in Fig. 1 exemplary for four different cases investigated. It was not possible to resolve the DNA extracted with phenol–chloroform on a 0.3% agarose gel. Thus, conventional agarose gel electrophoresis utilized in this study was not able to detect any fragmentation of genomic DNA attributable to the fixative.

3.3. DNA yields

DNA yield was significantly dependent on the extraction method used (Table 1). Phenol–chloroform (P–C) produced the best DNA yield followed by the extraction kits Puregene (Pur), NucleoSpin Tissue (NST) and QIAamp DNA Mini Kit (QIA) ($p < 0.0001$, P–C versus NST, Pur, and QIA; $p < 0.0001$, Pur versus NST and QIA; $p = n.s.$, NST versus QIA). Irrespective of the DNA extraction method, the DNA yield of bronchial secretions varied considerably. No significant correlation was found between degree of inflammation and total DNA yield (data not shown).

3.4. DNA purity

DNA purity estimated as ratio of $A_{260}/A_{280}$ was sufficient irrespective of DNA extraction method used (Table 1). The best purity was achieved with QIAamp DNA Mini Kit ($p < 0.005$) followed by Puregene, NucleoSpin Tissue and phenol–chloroform ($p < 0.0001$, QIA versus NST and P–C; $p < 0.005$, QIA versus Pur; $p < 0.01$, Pur versus P–C; $p < 0.05$, Pur versus NST).

3.5. DNA fragment size

The DNA fragment size was clearly dependent on the extraction method used (Table 1). The differences between extraction protocols in fragment size were statistically highly significant ($p < 0.0001$). Without exception DNA extracted by phenol–chloroform had such high molecular weight that it did not migrate into the 0.3% agarose gel (Fig. 1). DNA of high molecular weight with a median fragment size above 40 kb was also extracted using Puregene. In addition, smaller fragments were detected (minimal fragment size 17 kb). DNA isolated using the silica membrane based extraction kits NucleoSpin Tissue and QIAamp DNA Mini Kit was significantly more fragmentised producing a median fragment length of 26 and 23 kb, respectively. The minimal fragment size was 13 kb.
3.6. PCR amplification of \(\beta\)-globin gene fragments

The extracts of each of the four split samples of 15 bronchial aspirates were successfully amplified for the three different sized \(\beta\)-globin gene fragments resulting in a total of 180 successful consecutive PCR amplifications. Since no differences were observed between the four DNA isolation methods until then, the PCR-series was stopped.

4. Discussion

Numerous previous studies used tissue specimens to investigate the influence of different fixatives and DNA extraction methods on the quality of genomic DNA [2, 3]. On the other hand, there are only few such studies that applied routine diagnostic cytologic specimens [1,5,9]. In this study we tested four different protocols of DNA extraction from bronchial secretions and evaluated the extracted DNA with regard to yield, purity, integrity, and adequacy for PCR-based assays. All bronchial aspirates had been fixed with Saccomanno fixative immediately after bronchoscopy as it excellently preserves cell morphology and effectively reduces bacterial growth [4,7]. We showed that this fixative also preserves sample DNA as high molecular weight DNA. There was no fragmentation of genomic DNA attributable to the fixative when investigated using conventional agarose gels.

Garcia-Closas and co-workers [1] evaluated mouthwash samples fixed by 14.3 wt% alcohol and also found high molecular weight DNA (>23 kb) in all cases. Another study analysed DNA samples fixed in 14.3 wt% alcohol by pulse-field gel electrophoresis [9]. The majority of the recovered DNA was between 35 and 63 kb (range, >15–97 kb). The occurrence of DNA fragments below 40 kb in both studies may be linked to some DNA damage resulting from the use of a fixative with only 14.3 wt% alcohol instead of 50% ethanol used in this study.

All four isolation methods tested extracted a DNA well suitable to amplify even large PCR-products. This was demonstrated by the amplification of three different sized \(\beta\)-globin gene fragments (268, 536, and 989 bp). Kerstens et al. [5] used the same fixative (50% ethanol/2% carbowax) as we did for the fixation of cervix brushes and found it sufficient for reliable HPV screening with LiPA-PCR. Out of several fixatives tested by Greer et al. [2,3] 95% ethanol belonged to those most successful in subsequent PCR amplifications producing DNA amplification fragments of up to 1327 bp in length. However, alcohol of high concentration, as was used for tissue fixation in previous studies, coagulates mucoproteins and causes shrinkage of cells [7]. It is therefore not suitable for cytologic specimens. The usage of 14.3 wt% alcohol for the fixation of cytologic specimens proved a PCR success rate almost as high as found in this study and was only slightly less efficient with large (989 bp) PCR products (94% versus 100%) [1]. However, the low alcohol content suppressed bacterial growth incompletely resulting in considerable contribution of bacterial DNA to total DNA content [1]. In this study none of the specimens showed relevant bacterial growth as evaluated by light microscopy.

In contrast to several other fixatives like formalin, alcohol fixation over a period of several days, e.g., during mailing, does not decrease the PCR success rate [2,3].

Irrespective of DNA extraction method applied, the DNA yield of bronchial secretions varied considerably. This variation was not correlated to the degree of inflammation seen by light microscopic evaluation. It should be taken into consideration when planning a study using bronchial aspirates, that a considerable subset of the specimens may not be sufficient due to a low amount of total DNA.

This study showed that DNA yield, purity, and fragment size were significantly dependent on the DNA extraction protocol used. Phenol–chloroform extraction achieved the highest DNA yield and extracted the largest DNA fragments. DNA purity was slightly lower than that in other extraction methods, but still acceptable. Phenol–chloroform extraction has the advantage that the components can be adjusted to the considerably varying amounts of genomic DNA present in bronchial aspirates. However, the procedure is relatively time consuming and applies toxic chemicals. Remaining phenol does interfere with UV-spectrophotometry. Thus it is necessary to additionally check the absorption at 230 nm. The Puregene DNA extraction kit purifies the DNA by precipitating the proteins of the cellular lysate using ammonium acetate. Subsequently the DNA is precipitated with isopropanol. This method produced the second best results with regard to DNA yield, purity and fragment size which, in half of the cases, lies above 40 kb. On the one hand, this protocol is as flexible as phenol–chloroform extraction concerning variable DNA amounts present, on the other hand, a relatively
time consuming procedure. But it is not toxic. NucleoSpin Tissue and QIAamp DNA Mini Kit work on the same principle. The DNA is bound onto silica membranes, is purified and then eluted. The protocols are quick and easy to perform even for inexperienced laboratory staff. In addition, no toxic chemicals have to be handled. But both methods produced a lower DNA yield than phenol–chloroform extraction and Puregene. In addition, the DNA was significantly more fragmented. This is supported by experiments of other authors who also found that DNA prepared by binding to silica-based matrices tend to be of smaller size than DNA extracted with phenol–chloroform or the Puregene kit [1,9]. In this study the smallest DNA fragment extracted sized 13 kb and the median DNA fragment length was 26 (NucleoSpin Tissue) and 23 kb (QIAamp Kit) respectively. DNA extracts isolated by NucleoSpin Tissue came up with a slightly higher DNA fragment size, whereas DNA extracted by the QIAamp DNA Mini Kit showed the highest DNA purity of all protocols tested. As a consequence, both extraction kits are well suited if sufficient genomic DNA is present and a quick and easy DNA extraction method is desired.

In conclusion, fixation of bronchial aspirates by Saccomanno fixative provides genomic DNA well preserved for PCR-based methods in molecular cytology. Since the DNA content of bronchial aspirates varies considerably, extraction methods that can be adjusted to this appear favourable. However, the method of preference depends on the circumstances of application.

Appendix. DNA extraction protocols

Phenol–chloroform DNA extraction [8]

Place 200 μl resuspended cells, 200 μl lysis buffer (10 mM Tris-HCl pH 8.0, 0.1 M EDTA pH 8.0, 0.5% SDS (w/v)) and 20 μl proteinases K (20 mg/ml) into a 1.5 ml reaction tube and incubate at 65°C for 30 min. Add 20 μg/ml DNase-free pancreatic RNase and incubate for 30 min at 37°C. Add an equal volume of phenol–chloroform (1 : 1) and vortex for 10 s. Centrifuge the probe at 15,000 × g for 2 min. Transfer the aqueous layer to a new tube, add an equal volume of chloroform and vortex for 10 s. Centrifuge the probe at 15,000 × g for 2 min. Transfer the aqueous layer to a new tube. Add 0.1× volume of 3 M NaOAc (pH 5.2) and 2× volume of 100% EtOH, vortex the probe and store it overnight at −20°C. Centrifuge the probe at 15,000 × g for 10 min. Wash the DNA pellet with 70% EtOH. Dry and resuspend the DNA in 50 μl 10 mM Tris-HCl pH 8.0.

QIAamp DNA Mini Kit (according to manufacturer’s instructions)

Place 200 μl resuspended cells, 20 μl RNase A (10 mg/ml), 20 μl proteinases K (20 mg/ml) and 200 μl buffer AL in a 1.5 ml reaction tube, vortex for 5 s and incubate at 56°C for 10 min. Add 200 μl 100% EtOH. Vortex the mixture for 5 s and apply it into the QIAamp spin column in a 2 ml collection tube. Centrifuge the collection tube at 6000 × g for 1 min and add 500 μl buffer AW1. Centrifuge the spin column in a new collection tube at maximum speed for 1 min. Add 500 μl buffer AW2 and change the collection tube. Centrifuge at maximum speed for 1 min, discard the flow-trough and centrifuge once more. Put the spin column onto a clean 1.5 ml reaction tube, add 50 μl buffer AE and incubate for 5 min at room temperature. Elute the DNA by centrifugation at 6000 × g for 1 min.

NucleoSpin Tissue (according to manufacturer’s instructions)

Place 200 μl resuspended cells, 25 μl proteinases K (20 mg/ml) and 200 μl buffer B3 in a 1.5 ml reaction tube, vortex for 15 s and incubate at 70°C for 10–30 min. Digest RNA with 20 μl RNase A (10 mg/ml) for 30 min at 37°C. Add 210 μl 100% EtOH. Vortex the mixture for 5 s and apply it onto the NucleoSpin column in a 2 ml collection tube. Centrifuge the collection tube at 6000 × g for 1 min. Add 500 μl buffer BW and centrifuge the spin column in a new collection tube at maximum speed for 1 min. Add 600 μl buffer B5 and change the collection tube. Centrifuge at maximum speed for 1 min, discard the flow-trough and centrifuge once more. Place the spin column onto a clean 1.5 ml reaction tube, add 50 μl prewarmed (70°C) buffer BE and incubate for 5 min at room temperature. Elute the DNA by centrifugation at 6000 × g for 1 min.

4.1. Puregene (according to manufacturer’s instructions)

Place 200 μl resuspended cells in a 1.5 ml reaction tube and centrifuge for 5 min at 600 × g. Discard the supernatant and resolve the cell pellet in 50 μl. Add 250 μl cell-lysis-buffer and 25 μl of proteinases K (20 mg/ml). Vortex for 15 s and incubate at 65°C for 30 min. Digest RNA with 30 μl RNase A-solution (4 mg/ml) for 15 min at 37°C. Add 100 μl protein-precipitation-solution, vortex for 20 s and incubate for 5 min at 5°C. Centrifuge at 13,000 × g for 3 min. The
precipitated proteins should form a tight pellet, if not repeat this step. Transfer the supernatant in a 1.5 ml reaction tube and add 300 µl isopropanol. Mix by inverting the tube gently 50 times and incubate for 5 min at room temperature. Centrifuge at 13,700 \( \times g \) for 5 min. Discard the supernatant and wash the DNA pellet with 70% EtOH. Dry pellet and rehydrate DNA in 50 µl DNA-hydration-solution.

References


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