Developing a rapid, efficient and low cost method for rapid DNA extraction from arthropods

Desenvolvimento de um método rápido, eficiente e de baixo custo para extração de DNA de artrópodos

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ABSTRACT

Here, it is presented a rapid and efficient method to obtain good quality DNA from small samples of arthropod tissues generating low quantities of hazardous wastes. This new method was compared with another homemade protocol using phenol and other two commercial kits. The quality of DNA obtained was checked by spectrophotometer and evaluated by an AFLP assay. Low shearing DNA was obtained from all samples and the best readings were observed to DNA recollected with the new method. The AFLP assay indicated that DNA obtained with all methods were suitable for use in molecular biology techniques sensitive to contaminants. However, homemade protocols were more efficient in recollect DNA than commercial kits, without lose any quality of samples. Also, they were less time and fund consuming, with costs ten times cheaper than commercial kits. The quicker, less pollutant and cheaper protocol was the one described here (USD 0.52 per sample).

Key words: arthropod DNA, high quality DNA, molecular biology, AFLP.

INTRODUCTION

The analysis of genetic variation using deoxyribonucleic acids (DNA) polymorphisms has become an important tool to access evolutionary history, genetic diversity, identification of cell, bacteria, and virus strains, as well as some ecology issues and genetic improvement programs. The first step in DNA analyses is the extraction of nucleic acids from tissues or cells. Several methods for DNA extraction have been described for different groups and biological material (MOELLER et al., 1992; LODHI et al., 1994; YEATES et al., 1998; LEFORD & DOUGLAS, 1999; SAMBROOK & RUSSELL, 2001; BACKER et al., 2001; ROHLAND et al., 2004; GRACHEV et al., 2006; LEE & PRYS-JONES, 2008; DE OLIVEIRA et al., 2009) and the most important

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Received 07.30.10   Approved 06.21.11   Returned by the author 07.27.11
concern on these methods have been the quantity and quality of DNA. The quality of samples depends on biological material as well as on protocol followed (REINEKE et al., 1998; CHEN et al., 2008).

Homemade DNA extraction protocols are cheap and very efficient to recollect DNA from samples, but are usually time-consuming; commercial kits are generally very fast but too expensive and generate large quantities of contaminant wastes, such as beads, filters, columns and microcentrifuge tubes. Also, homemade DNA extraction protocols are less efficient in the removal of proteins/carbohydrates and other organic contaminants from samples or extraction solutions used, thus causing enzymatic inhibition in further molecular analysis. In these cases the use DNA clean-up steps are required, which can be time and fund consuming. An ideal protocol should optimize DNA yield, minimize degradation, and be efficient in terms of cost, time, labor, and supplies (CHEN et al., 2010).

REINEKE et al. (1998) evaluated several DNA extraction protocols using insect tissue samples and determined the quality of DNA recollected by Amplified Fragment Length Polymorphism (AFLP). The study also investigated the effect of additional clean-up steps on the improvement of DNA quality, concluding that the best protocols to obtain good quality DNA for AFLP are phenol or phenol-chloroform based methods, but for these protocols additional purification steps are recommend. Cationic detergent cetyltrimethyl ammonium bromide (CTAB)-based methods, as described in MOELLER et al. (1992), demonstrated to provide good quality DNA samples, using less hazardous chemicals, by combining its use with some proteinase and ribonuclease (RNAse), which can cause more fund consuming.

Here, it is described a method to obtain good quality DNA from small samples of freshly and frozen insect tissues. The method is CTAB-based and does not use phenol, a hazardous chemical and a contaminant in DNA samples. Sodium Chloride (NaCl) has been used to remove polysaccharides (FANG et al., 1992), and polyvinyl pyrrolidone (PVP) to purge polyphenols from arthropods diet (MALIYAKAL, 1992). To improve DNA quality it was used cross-linked polyvinyl pyrrolidone (PVPP) to remove alkaloids, terpenoids, glycosides, phenols, phenazines polyketides and peptides, as well as protein and carbohydrate (HAAF et al., 1985). The main objective of this research was to develop a quick, cheap and less hazardous alternative to protocols available on literature and market.

**MATERIAL AND METHODS**

Species and tissue samples

Two flies, two butterflies, one moth, two spiders and three isopods species were used as biological material for test the protocol’s efficiency to extract DNA from tissue samples. The species used were, respectively, *Anastrepha fraterculus*, *Drosophila immigrans*, *Dryas iulia alcionea*, *Heliconius erato phyllis*, *Grapholita moesta*, *Paratrechalea azul*, *Paratrechalea ornata*, *Benthana cairensis*, *Balloniscus glaber*, *Balloniscus sellowi*. All specimens were collected from nature and weighted. The tissue sources used for DNA extraction were whole body (flies, isopods and moth), head and thorax (butterflies) and cephalothorax (spiders). Each sample was frozen in liquid nitrogen and immediately crushed into a fine powder in 1.5ml sterile microcentrifuge tubes.

The DNA extraction protocol

Crushed tissues were mixed with PVPP (Sigma, P6755) in a proportion of 100mg of PVPP per 1g of grinded tissue, 600μL of extraction buffer (50mM Tris pH 8.0, 20mM EDTA pH 8.0, 1.1M NaCl, 0.4M LiCl, 1% CTAB, 2% PVP40 (Sigma), 0.5% Tween 20, 0.2% β-mercaptoethanol) in 1.5ml microcentrifuge tubes and mixed thoroughly. The tubes were incubated at 60°C for 25 minutes and inverted each 10 minutes for good solution of the crushed tissues with the buffer. The tubes were cooled at room temperature and 600μL of chloroform: isoamyl alcohol mixture (24:1) was added. The tubes were then mixed gently by inversion to form an emulsion during 4 minutes. After that the tubes were centrifuged (Eppendorf Centrifuge 5415R) at 10.600g for 5 minutes at room temperature. The aqueous phase present in each tube (~500μL) was transferred to a new 1.5ml microcentrifuge tube gently, to avoid DNA shearing. To each tube was added 250μL of 5M NaCl and 750μL of cold (-20°C) isopropanol. The solution was kept in the freezer at -20°C for 20 minutes to improve precipitation nucleic acids. Samples were then centrifuged at 10.600g (Eppendorf Centrifuge 5415R) for 10 minutes at room temperature and the aqueous poured off. DNA pellets were washed with cold (4°C) 76% ethanol (1000μL). After the washing step, the samples were spun quickly and the excess of ethanol removed with a micropipette. Washed DNA pellets were dried by leaving the tubes uncovered in a 37°C stove for 20 minutes. DNA samples were dissolved in 50μL TE (10mM Tris HCl pH 8.0, 0.1mM EDTA pH 8.0) and treated with 1μL of RNAase A (10mg mL⁻¹) at 50°C for 20 minutes.
Quality and quantification tests

Integrity of the DNA samples was analyzed in 0.8% ethidium bromide-stained agarose gels, using 1Kb Ladder (Fermentas) as control for DNA shearing. Visual patterns of DNA samples were photographed under ultraviolet light and scanned with Stratagene EagleEye II still video system. The DNA samples were quantified using a spectrophotometer (GeneQuant PRO, Amersham Biosciences) in the absorbance spectrum for nucleic acids (260 nm). To access DNA purity, 230 nm and 280 nm readings were recorded and compared with 260 nm readings. The 230 nm reading can indicate the presence of organic impurities and buffer contaminants, once phenols, carbohydrates, peptides, urea, thiocyanates, β-mercaptoethanol, humic acids and other buffer components absorb light at this wavelength (YEATES et al., 1998; SAMBROOK & RUSSELL, 2001; HANSEN et al., 2007). The 280 nm reading indicate the presence of proteins originated from the cells where the nucleic acids were extracted (YEATES et al., 1998; HANSEN et al., 2007), once tyrosine, tryptophan and phenylalanine amino acids are absorbed at this wavelength (SAMBROOK & RUSSELL, 2001). It was considered as having good DNA quality, the samples which exhibited 260/230 and 260/280 nm rates higher than 1.7. Background correction was done using the 320 nm reading to compensate the turbidity.

AFLP assay and evaluation of method’s efficiency against other DNA extraction protocols

The quality of some DNA samples was also validated by an AFLP assay (VOS et al., 1995), a sensitive multi-step molecular technique based on the detection of genomic restriction fragments by polymerase chain reaction (PCR), which requires high quality DNA samples, free of organic compounds, proteins and RNA. During the AFLP assay, it was compared the AFLP profiles of the DNA samples obtained with protocol presented here with DNA samples obtained with another three protocols. The first one was a homemade sodium dodecyl sulfate (SDS)/phenol-chloroform based protocol, modified from RIED (2002). The other two protocols were guanidine hydrochloride/mini-column based protocols from commercial kits available on market for animal genomic DNA isolation: a silica membrane purification method from Macherey-Nagel GmbH & Co. KG (NucleoSpin® Tissue, Ref. n° 740952.10), and a resin membrane purification method from Norgen Biotek Corp. (Genomic DNA Isolation Kit, Ref n° 24700). Additional information on DNA shearing and spectrophotometer absorbance were also provided for the DNA samples. All DNA samples used in this comparative test were obtained from Heliconius erato (Lepidoptera) specimens (numbered from H01 to H08) collected from a natural population near Porto Alegre, southern Brazil (lat -30.06681, log -51.122582), and were prepared as described for the other species. The selective amplification step of the AFLP assays were performed using four set of primers EcoR1+3 and MseI+3 nucleotides (set 1, EcoR1+ACC / MseI+CGC; set 2, EcoR1+AGG / MseI+CCG; set 3, EcoR1+ATA / MseI+CCT; and set 4, EcoR1+ACT / MseI+CCC) and visualized in silver stained 6% polyacrylamide denaturing gels according to CRESTE et al. (2001). It was considered as having good quality DNA the samples which produced clear AFLP patterns.

RESULTS AND DISCUSSION

The DNA extraction protocol

After the protocol application, DNA was obtained from all arthropod species and samples exhibited a transparent to very light color. In the 0.8% ethidium bromide-stained agarose gels the DNA samples revealed good quality (Figure 1). The DNA-shearing effect was not prominent for any sample and only low smearing fragments were observed. This low shearing effect is expected, since the tissue samples were mechanically grinded and subjected to homogenization procedures. The RNase treatment was efficient to eliminate RNA from the samples, once no RNA was observed in any sample. The table 1 shows the results for the protocol’s efficiency test with the different species. The yield of DNA recollected from the tissue samples shown a significant correlation between the weight of tissue sample used and the quantity of DNA recollected was observed (r=0.842; P<0.001). All samples tested shown high quality DNA after extraction procedure (Table 1). No significant amounts of proteins were detected in the samples, showing that protein removal was very efficient using the protocol presented here, without the necessity of using phenol or any other proteolytic method. Organic compounds presented in DNA samples also were inexpressive, dispensing the need of purification step after the extraction.

AFLP assay and evaluation of method’s efficiency against other DNA extraction protocols

As observed for the other arthropod species, all DNA samples used for the AFLP assay revealed good quality. DNA samples obtained with the four methods compared shown similar results, but the yield of DNA recollected with the commercial kits was lower than homemade protocols. Wavelength readings
and wavelength rates also were very similar, excepting the samples H06 and H08, which were obtained with the commercial kits; those samples revealed some excess of organic impurities or contaminants (Table 1). No expressive amounts of proteins were observed in any sample.

The figure 2 shows the AFLPs patterns obtained for the DNA samples used in the AFLP assay. All four set of primers tested exhibited a great number of loci with good visualization. Overall, it was observed 251 distinguished loci, being 240 polymorphic. Regarding the DNA quality for molecular application, all four methods provided good quality, proving to be suitable for enzyme digestion, linkage and PCR, once AFLP has those three steps and it is very sensitive to poor quality DNA samples (REINEKE et al., 1998; CHEN et al., 2008).

The results obtained with the AFLP assay indicate that the use of different methods may produce good quality DNA. Homemade protocols provide an alternative to low budget projects, but must be optimized to produce good results. Commercial kits are very convenient, but also too expensive. These statements become clear when time/fund consumption and toxic wastes produced are concerned (Table 2). The homemade protocol based on SDS/phenol-chloroform extraction showed similar runtime and results to the CTAB/PVPP/chloroform method presented here, and also was very inexpensive as well. Moreover, the use of phenol demands an additional step of organic solvent cleaning which generates higher amounts of contaminated wastes when compared with the method developed here. On the other hand, the commercial kits showed lower efficiency in recollect DNA from tissue samples when compared with the homemade protocols, without gain any advantages on samples purity. Despite having less time consuming, they were ten times more expensive than the homemade protocols. They did not generate hazardous wastes containing phenol and chloroform, dispensing the need of a fume hood to be applied, but the use of guanidine hydrochloride solutions also produces noxious wastes and some care on the disposal of residues should be taken as well. Scoring the benefits and costs of all protocols compared, the results indicate that the method presented here is the best choice to obtain good quality DNA.
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High yields of clean DNA were obtained from arthropod tissues by using the DNA extraction procedure presented here. The strategy of using NaCl to remove polysaccharides and PVP to purge polyphenols from arthropods’ diet proved to be very efficient. The PVPP proved to be a proficient tool to isolate high quality DNA from tissue samples, especially by removing proteins and organic contaminants, without the need of further purification steps. It was also essential to keep apart cellular debris from the aqueous phase containing nucleic acid solution, by forming a compact and consistent interface.

Although the DNA extraction protocol developed here had shown promising results, some care must be taken to maximize the yield and quality of the DNA samples. First, the tissue sample must be completely dried before the grind procedure starts. The formation of ice crystals can prejudice the grinding procedure. Second, excess or lack of grinded tissue can generate samples with excess of proteins and organic compounds or residual buffer components, respectively. Although, as shown in the AFLP assay, the residual buffer components presented in the DNA samples did not interfere in the AFLP technique. Excess of tissue can cause pigmentation of DNA samples after extraction, but also increase the amount of DNA and other substances. The pigmentation does not affect the quality of DNA suitable to sensitive molecular techniques, but to obtain best results it is recommend increasing the dilution factor of DNA samples (75-200 μl of TE). According to the observations best results are achieved when 0.10-0.06g of grinded tissue are used.

Table 1 - Analysis of concentration, wavelength absorbance readings and rates, and extraction efficiency for the biological samples used in the efficiency test and in the AFLP assay.

<table>
<thead>
<tr>
<th>Species/Sample 1</th>
<th>DNA concentration (ng μl⁻¹)</th>
<th>Wavelength absorbance (A)²</th>
<th>Wavelength rates³</th>
<th>Extraction efficiency⁴</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>230μm</td>
<td>260μm</td>
<td>280μm</td>
</tr>
<tr>
<td>Efficiency test</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A. fraterculus (n=1)</td>
<td>640.0</td>
<td>0.078</td>
<td>0.137</td>
<td>0.076</td>
</tr>
<tr>
<td>A. fraterculus (n=2)</td>
<td>1545.0</td>
<td>0.189</td>
<td>0.336</td>
<td>0.188</td>
</tr>
<tr>
<td>B. cairensis (n=1)</td>
<td>955.0</td>
<td>0.119</td>
<td>0.204</td>
<td>0.120</td>
</tr>
<tr>
<td>B. glaber (n=1)</td>
<td>877.0</td>
<td>0.106</td>
<td>0.183</td>
<td>0.107</td>
</tr>
<tr>
<td>B. sellowii (n=1)</td>
<td>690.0</td>
<td>0.083</td>
<td>0.142</td>
<td>0.081</td>
</tr>
<tr>
<td>D. iulia (n=1)</td>
<td>1050.0</td>
<td>0.133</td>
<td>0.233</td>
<td>0.135</td>
</tr>
<tr>
<td>D. immigrans (n=1)</td>
<td>185.0</td>
<td>0.219</td>
<td>0.377</td>
<td>0.195</td>
</tr>
<tr>
<td>D. immigrans (n=20)</td>
<td>5580.0</td>
<td>0.263</td>
<td>0.570</td>
<td>0.284</td>
</tr>
<tr>
<td>G. molesta (n=1)</td>
<td>148.0</td>
<td>0.188</td>
<td>0.323</td>
<td>0.189</td>
</tr>
<tr>
<td>G. molesta (n=4)</td>
<td>785.0</td>
<td>0.095</td>
<td>0.164</td>
<td>0.093</td>
</tr>
<tr>
<td>H. erato (n=1)</td>
<td>885.0</td>
<td>0.108</td>
<td>0.186</td>
<td>0.109</td>
</tr>
<tr>
<td>P. azul (n=1)</td>
<td>2150.0</td>
<td>0.258</td>
<td>0.442</td>
<td>0.241</td>
</tr>
<tr>
<td>P. ornata (n=1)</td>
<td>630.0</td>
<td>0.078</td>
<td>0.134</td>
<td>0.073</td>
</tr>
<tr>
<td>AFLP assay</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H. erato H01 (n=1)A</td>
<td>649.0</td>
<td>0.873</td>
<td>1.492</td>
<td>0.929</td>
</tr>
<tr>
<td>H. erato H02 (n=1)A</td>
<td>905.5</td>
<td>1.229</td>
<td>1.998</td>
<td>1.111</td>
</tr>
<tr>
<td>H. erato H03 (n=1)B</td>
<td>753.0</td>
<td>0.803</td>
<td>1.532</td>
<td>0.797</td>
</tr>
<tr>
<td>H. erato H04 (n=1)B</td>
<td>1020.0</td>
<td>1.381</td>
<td>2.154</td>
<td>1.215</td>
</tr>
<tr>
<td>H. erato H05 (n=1)C</td>
<td>129.5</td>
<td>0.132</td>
<td>0.259</td>
<td>0.135</td>
</tr>
<tr>
<td>H. erato H06 (n=1)C</td>
<td>92.0</td>
<td>0.199</td>
<td>0.191</td>
<td>0.111</td>
</tr>
<tr>
<td>H. erato H07 (n=1)D</td>
<td>107.0</td>
<td>0.124</td>
<td>0.214</td>
<td>0.121</td>
</tr>
<tr>
<td>H. erato H08 (n=1)D</td>
<td>55.5</td>
<td>0.210</td>
<td>0.130</td>
<td>0.087</td>
</tr>
</tbody>
</table>

1 Numbers in brackets indicate the quantity of specimens used.
2 Absorbance were measured by the optical density of the sample in relation to ultrapure distilled water.
3 Wavelength rate higher than 1.7 indicates pure samples and lower than 1.7 indicate samples with significant levels of buffer/organic components (260/230μm) and proteins (260/230μm).
4 In grams of DNA / grams of tissue.
5 Mega e Revers’ method (CTAB/PVPP/chloroform - described here).
6 Ried’s method (SDS/phenol-chloroform).
7 Kit 1 method (Macherey-Nagel GmbH & Co. KG, NucleoSpin® Tissue - Ref. n° 740952.10).
8 Kit 2 method (Norgen Biotek Corp., Genomic DNA Isolation Kit - Ref. n° 24700).
It was concluded that the protocol presented here is a good method to obtain high quality DNA without the necessity of further steps of proteolysis and purification. The protocol has an approximate cost of USD 0.52 per sample, including all chemicals and plastics, which is very inexpensive if...
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Ciência Rural, v.41, n.9, set, 2011.

compared with the commercial kits. Also, hazardous wastes at the end of extraction procedure totaled one microcentrifuge tube with cellular debris and 600 μl of chloroform and isoamyl alcohol mixture per sample. Thus, researchers from Agrarian and Biological Sciences may study arthropod DNA at low costs, generating lower quantities of toxic waste when compared with other methods and commercial kits available on market.

ACKNOWLEDGMENTS

The researchers are grateful to Alessandra Russi, Felippe Gomes, Gisele Passaia, Ivan Ceriotti, Rafaela Nalin and Vanessa Buffon for providing laboratory support, and Aline Quadros, Lisandra Robe, Luis E. Schmidt and Wilson Morandi Jr. for providing specimens for DNA extraction. This study was funded by CNpq, EMBRAPA Uva e Vinho and FAPERGS.

REFERENCES


Table 2 - A comparative of time, costs and hazardous wastes generated per sample with the four methods used to recollect DNA from fresh arthropod tissues.

<table>
<thead>
<tr>
<th>Method</th>
<th>Time to finish the protocol</th>
<th>Cost per sample (USD)</th>
<th>Hazardous wastes per sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mega &amp; Revers^A</td>
<td>110 min.</td>
<td>0.52</td>
<td>One microtube contaminated with chloroform; ≈600ul of chloroform contaminated residues</td>
</tr>
<tr>
<td>Ried’s^B</td>
<td>125 min.</td>
<td>0.67</td>
<td>Two microtube contaminated with phenol and chloroform; ≈1.2 ml phenol and chloroform contaminated residues</td>
</tr>
<tr>
<td>Kit 1^C</td>
<td>110 min.</td>
<td>5.10</td>
<td>One microtub, two collection tubes and one column contaminated with guanidine hydrochloride; ≈1.7ml of guanidine hydrochloride contaminated residues</td>
</tr>
<tr>
<td>Kit 2^D</td>
<td>85 min.</td>
<td>4.03</td>
<td>One microtube, one collection tube and one column contaminated with guanidine hydrochloride; ≈1.3ml of guanidine hydrochloride contaminated residues</td>
</tr>
</tbody>
</table>

1 Grinding tissue time and solution preparation time not considered.
2 Sample incubation time of one hour.
3 Calculated on February 2011, from lowest Brazilian market prices.
A CTAB/PVPP/chloroform - described here.
B SDS/phenol-chloroform.
C Macherey-Nagel GmbH & Co. KG, NucleoSpin® Tissue (Ref. nº 740952.10).
D Norgen Biotek Corp., Genomic DNA Isolation Kit (Ref. nº 24700).


