METHOD ARTICLE

A DNA extraction protocol for improved DNA yield from individual mosquitoes [version 1; referees: 3 approved]

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Abstract

Typical DNA extraction protocols from commercially available kits provide an adequate amount of DNA from a single individual mosquito sufficient for PCR-based assays. However, next-generation sequencing applications and high-throughput SNP genotyping assays exposed the limitation of DNA quantity one usually gets from a single individual mosquito. Whole genome amplification could alleviate the issue but it also creates bias in genome representation. While trying to find alternative DNA extraction protocols for improved DNA yield, we found that a combination of the tissue lysis protocol from Life Technologies and the DNA extraction protocol from Qiagen yielded a higher DNA amount than the protocol using the Qiagen or Life Technologies kit only. We have not rigorously tested all the possible combinations of extraction protocols; we also only tested this on mosquito samples. Therefore, our finding should be noted as a suggestion for improving people’s own DNA extraction protocols and not as an advertisement of a commercially available product.

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Competing interests: No competing interests were disclosed.

**Introduction**

DNA extraction for *Anopheles* mosquitoes is typically done using commercially available products (Brown et al., 2011; Demirci et al., 2012; Horton et al., 2010; Main et al., 2015; Norris et al., 2015; Weetman et al., 2012). They work well enough to provide a sufficient amount of DNA for PCR-based assays from a single mosquito. However, next-generation sequencing applications and high-throughput SNP genotyping assays exposed the limitation of DNA quantity from a single mosquito using typical extraction protocols (Marsden et al., 2011). Whole genome amplification could alleviate the issue but it also creates bias in genome representation (see Table 1 and results section below).

While trying to find alternative DNA extraction protocols for improved DNA yield but without compromising the automation option, we found that a combination of tissue lysis protocol from Life Technologies and DNA extraction protocol from Qiagen yielded higher DNA amount than the protocol using Qiagen or Life Technologies kit only.

**Method**

**Samples**

Adult mosquitoes from a single generation of Pimperena and Mopti-NIH colonies obtained from Malaria Research and Reference Reagent Resources Center (Manassas, VA) were used for this study. Prior to extraction, the whole adult mosquito samples are individually preserved either by freezing at -20°C or storing in 80% ethanol; for the latter case, samples are rehydrated in water for 1 hour prior to DNA extraction.

**Tissue lysis using Life Technologies MagMAX protocol**

We used a protocol adapted from the manufacturer’s mouse tail protocol and Whatman FTA card protocol. For each sample contained in a 1.5mL tube, 10µL or 40 µL of proteinase K (Qiagen, 20mg/mL concentration) were added with 90µL or 60µL ATL buffer (Qiagen, Vallencia, CA), respectively. A 3mm diameter steel bead was added to each of the remaining samples and homogenized using Qiagen Tissulyser (Qiagen, Valencia, CA) for 30sec at 30Hz. DNA lysis buffer was added to each tube. Each tube was vortexed briefly (<10s) and centrifuged at 15,000rpm using an Eppendorf micro-centrifuge. This created white precipitate, which was mixed by pipetting up and down several times before transferring to a 2.0mL deep well plate for DNA extraction. 90µL of lystate was used for DNA extraction using the Biosprint 96 instrument. The other 90µL of lystate from the same sample was used for DNA extraction using the MagMAX™ Express-96 Magnetic Particle Processor.

**DNA extraction using Life Technologies MagMAX protocol**

As noted earlier, we used a protocol adapted from the manufacturer’s mouse tail protocol and Whatman FTA card protocol. 120µL of 100% isopropanol was added to each lyste. The plate containing lystate and isopropanol was gently mixed (220rpm) using a shaker for 3 minutes. 20µL of DNA binding bead mix (16µL binding beads and 4µL of PCR-grade water) was added to each sample and shook for 3 minutes at 220rpm. We used the “4412021 DW Blood” protocol on the MagMAX instrument, which washes lystate once with wash buffer 1 and twice with wash buffer 2 (Life Technologies).

**Table 1.** Whole genome sequencing quality comparison between original and whole-genome-amplified (WGA) DNA. Increase or decrease in comparison with sequence from original DNA is marked in up or down arrows in parentheses.

<table>
<thead>
<tr>
<th>Sample</th>
<th>DNA type</th>
<th>Total reads</th>
<th>% Unmapped</th>
<th>Mean coverage</th>
<th>Insert size</th>
<th>Whole</th>
<th>X</th>
<th>2L</th>
<th>2R</th>
<th>3L</th>
<th>3R</th>
<th>UNKN</th>
<th>chrM</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>A. gambia</em> 1</td>
<td>WGA</td>
<td>63,903,121</td>
<td>(↑)</td>
<td>1.22% (↓)</td>
<td>318 (↑)</td>
<td>27.02</td>
<td>26.5</td>
<td>20.86</td>
<td>47.8</td>
<td>18.6</td>
<td>21.38</td>
<td>15.39</td>
<td>57.23 (↓)</td>
</tr>
<tr>
<td>original</td>
<td></td>
<td>17,720,693</td>
<td>1.76%</td>
<td>185</td>
<td>7.19</td>
<td>8.15</td>
<td>7.29</td>
<td>6.71</td>
<td>6.9</td>
<td>6.91</td>
<td>8.12</td>
<td>897.11</td>
<td></td>
</tr>
<tr>
<td><em>A. gambia</em> 2</td>
<td>WGA</td>
<td>42,663,199</td>
<td>(↑)</td>
<td>17.46% (↓)</td>
<td>92 (↓)</td>
<td>11.06</td>
<td>21.4</td>
<td>8.21</td>
<td>6.28</td>
<td>11.6</td>
<td>7.87</td>
<td>13.64</td>
<td>38.48 (↓)</td>
</tr>
<tr>
<td>original</td>
<td></td>
<td>1,893,678</td>
<td>33.37%</td>
<td>141</td>
<td>0.38</td>
<td>0.43</td>
<td>0.39</td>
<td>0.37</td>
<td>0.37</td>
<td>0.36</td>
<td>0.36</td>
<td>214.62</td>
<td></td>
</tr>
<tr>
<td><em>A. gambia</em> 3</td>
<td>WGA</td>
<td>34,384,169</td>
<td>76.98% (↑)</td>
<td>462 (↑)</td>
<td>3.75 (↑)</td>
<td>4.48</td>
<td>3.79</td>
<td>3.87</td>
<td>3.19</td>
<td>3.4</td>
<td>2.06</td>
<td>10.61 (↓)</td>
<td></td>
</tr>
<tr>
<td>original</td>
<td></td>
<td>11,877,569</td>
<td>21.71%</td>
<td>261</td>
<td>3.30</td>
<td>3.62</td>
<td>3.32</td>
<td>3.23</td>
<td>3.16</td>
<td>3.18</td>
<td>2.77</td>
<td>132.53</td>
<td></td>
</tr>
<tr>
<td><em>A. arabiensis</em> 1</td>
<td>WGA</td>
<td>30,334,217</td>
<td>8.07% (↑)</td>
<td>464 (↑)</td>
<td>13.47 (↑)</td>
<td>16.93</td>
<td>12.99</td>
<td>15</td>
<td>10.4</td>
<td>12.11</td>
<td>7.03</td>
<td>5.7 (↓)</td>
<td></td>
</tr>
<tr>
<td>original</td>
<td></td>
<td>14,357,720</td>
<td>4.21%</td>
<td>489</td>
<td>6.20</td>
<td>9.8</td>
<td>5.4</td>
<td>5.2</td>
<td>5.3</td>
<td>5.4</td>
<td>10.2</td>
<td>11.9</td>
<td></td>
</tr>
</tbody>
</table>
Initial heated elution volume was 75µL (elution buffer 1) and then when prompted, 75µL of elution buffer 2 was added to complete the DNA elution step.

### DNA extraction using Qiagen Biosprint protocol

100µL of 100% isopropanol, 100µL of AL buffer (Qiagen) and 15µL of MagAttract Suspension (Qiagen) was added to each lysate. We used the “BS96 DNA Tissue” protocol on the BioSprint 96 instrument, which washes lysate twice with AW1 buffer, twice with AW2 buffer (Qiagen), and once with water with added tween 20 (Sigma) at a final concentration of 0.02%. The DNA was eluted in 150µL AE buffer (Qiagen).

### DNA quantification & analysis

DNA yield was measured using a Qubit high sensitivity, double stranded DNA kit (Life Technologies), using 1µL of input DNA. R statistics software version 3.0.0 was used to calculate mean and standard deviation and to perform Wilcoxon rank sum test with α of 0.05 after multiple comparison.

### Whole genome sequencing

5µL of original input DNA was used to amplify the whole genome using Qiagen Repli-g kit. We followed the manufacturer’s protocol. We followed the library protocol provided in Norris et al. (2015). Genomic DNA libraries were sequenced using Illumina HiSeq2500 platform with paired-end 150 bp reads at the QB3 Vincent J Coates Genomics Sequencing Laboratory at UC Berkeley. Adaptor sequences and poor quality sequences were trimmed from the raw Illumina fastq files using the Trimmmomatic software version 0.30 (Bolger et al., 2014) using default options. Reads were aligned to the A. gambiae reference genome AgamP3 (Giraldo-Calderon et al., 2015) using BWA-MEM version 0.7.5 (Li, 2013).

### Results & Discussion

**Dataset 1. Raw data for ‘A DNA extraction protocol for improved DNA yield from individual mosquitoes’**

http://dx.doi.org/10.5256/f1000research.7413.d107517

Sample: An arbitrary sample identification code; Group: Particular protocol id listed in Table 3 in the main paper; Read 1: DNA concentration reading in ng/µL using Qubit instrument; Read 2: DNA concentration reading in ng/µL using Qubit instrument using different standard curve; Read 3: DNA concentration reading in ng/µL using Qubit instrument using different standard curve; DNA concentration: Average DNA concentration of Read 1–3 as consensus DNA quantification; Overall mean: The mean of DNA concentrations for each group; std: The standard deviation of DNA concentrations for each group; %>(0.375ng/µL): The percentage of samples yielded greater than 0.375ng/µL DNA.

In our attempt to sequence whole genomes from field-collected individual mosquitoes, about 50% of specimens failed to pass the DNA quantity required (>30ng) for whole genome sequencing. These DNA samples were extracted using our established DNA extraction protocols using Qiagen kits and instruments (Lee et al., 2009; Main et al., 2015; Marsden et al., 2011; Norris et al., 2015; Slotman et al., 2006). The requirement of high genomic DNA content is not new. In the past, people have circumvented the problem by conducting whole genome amplification (Lee et al., 2013; Marsden et al., 2011; Weetman et al., 2012). We sequenced the whole genomes using original DNA in parallel with whole-genome amplified DNA to test if we can use whole genome amplification to bypass the DNA quantity issue.

Whole genome amplified DNA provided a higher number of reads than the original DNA with less DNA input (Table 1). However, comparison revealed that the particular whole genome amplification kit we used is not suitable for retrieving certain sections of genomes such as mitochondrial genome. This is indicated in the lower depth of coverage in mitochondrial genome in whole genome amplified material while the rest of chromosomes had higher depths relative to the library from original DNA. More importantly, the sequence generated from whole genome amplified samples produced number of inconsistent genotype calls (Table 2). This inconsistency became more apparent in mitochondrial sequences where heterozygous calls were produced where the genome sequence from the same original DNA had no such calls. These biases are likely introduced by the random primers used in the whole genome amplification kit. This result prompted us to pursue developing better DNA extraction protocols to improve DNA yield in an automated setup.

We found that the Life Technologies tissue lysis and extraction protocol (Table 3, line 4, in purple) was highly consistent in its DNA yield. A combination of the Life Technologies tissue lysis with the Qiagen BioSprint DNA extraction protocol (Table 3, line 3, in green) gave the highest average DNA yield (Wilcoxon rank sum test P-value=0.0031). The amount of magnetic beads added to tissue lysis had little effect on DNA yield. The amount of protease K (2µL vs 8µL) also showed little difference in DNA yield. Chemical lysis alone, without physical disruption, was not sufficient to produce consistency in DNA yield (Table 3, lines 5 and 6).

For a typical PCR-based assay, DNA quantity of 0.25–1.8 ng/µL in 200µL volume is sufficient. However, genomic approaches such as whole genome DNA library construction for next generation sequencing demand as little as 30ng of DNA. In our typical Qiagen BioSprint DNA extraction protocol, roughly 50% of DNA samples failed to yield 0.375ng/µL (Table 3, line 1, in blue), which leaves ~50µL of DNA for future study and allows for only a single trial of whole genome library construction. This constraint became a significant hindrance in our research involving whole genome sequencing from an individual mosquito.

This improved DNA extraction protocol will increase the chance of library construction from a single individual. Our observation is limited to trying out commercially available automated DNA extraction protocols and we do not have sufficient expertise on why certain protocols worked better or worse than others. As genomic approaches are more readily available to researchers, this improved DNA extraction protocol will facilitate such approaches that demand high-quantity DNA input from limited source material.

We have not rigorously tested all the possible combinations of extraction protocols. We only tested this on mosquito samples, and we only explored high-throughput automated DNA extraction protocols as we typically handle hundreds of mosquito samples at a...
Table 3. DNA yield for different combinations of lysis and DNA extraction protocols. Group 1 is the standard protocol used at the Vector Genetics Laboratory over ten years.

<table>
<thead>
<tr>
<th>Group</th>
<th>Lysis</th>
<th>Physical disruption</th>
<th>DNA extraction</th>
<th>Mean conc. (ng/µL)</th>
<th>SD</th>
<th>% (&gt;0.375ng/µL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Qiagen proteinase K + ATL</td>
<td>Yes</td>
<td>Qiagen Biosprint</td>
<td>0.63</td>
<td>0.63</td>
<td>50.0%</td>
</tr>
<tr>
<td>2</td>
<td>Qiagen proteinase K + ATL</td>
<td>Yes</td>
<td>LifeTech MagMAX</td>
<td>0.74</td>
<td>0.20</td>
<td>83.3%</td>
</tr>
<tr>
<td>3</td>
<td>LifeTech proteinase K + PK + lysis buffer</td>
<td>Yes</td>
<td>Qiagen Biosprint</td>
<td>1.33</td>
<td>0.80</td>
<td>92.9%</td>
</tr>
<tr>
<td>4</td>
<td>LifeTech proteinase K + PK + lysis buffer</td>
<td>Yes</td>
<td>LifeTech MagMAX</td>
<td>0.84</td>
<td>0.07</td>
<td>100.0%</td>
</tr>
<tr>
<td>5</td>
<td>LifeTech proteinase K + PK + lysis buffer</td>
<td>No</td>
<td>Qiagen Biosprint</td>
<td>0.88</td>
<td>0.55</td>
<td>66.7%</td>
</tr>
<tr>
<td>6</td>
<td>LifeTech proteinase K + PK + lysis buffer</td>
<td>No</td>
<td>LifeTech MagMAX</td>
<td>0.56</td>
<td>0.33</td>
<td>66.7%</td>
</tr>
</tbody>
</table>
time for population genetics studies. Therefore, our findings should be noted as suggestion for improving people’s own DNA extraction protocols and not as an advertisement of a commercially available product.

Data availability


Author contributions

CCN and YL conceived the study. CCN, YY conducted DNA extractions. CCN conducted library preparations. TCC and YL conducted genome sequence data analysis. CCN and YL conducted data analysis for DNA quantity from different protocols. CCN, YY, TCC and YL wrote manuscript.

Competing interests

No competing interests were disclosed.

Grant information

Signature Research in Genomics (SRG) Program by the UC Davis Office of Research and School of Medicine supported this research.

I confirm that the funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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References


Open Peer Review

Current Referee Status: ✔ ✔ ✔

Version 1

Referee Report 10 February 2016
doi:10.5256/f1000research.7988.r11970

Karla Saavedra-Rodriguez
Department of Microbiology, Immunology and Pathology, Colorado State University, Fort Collins, CO, USA

The manuscript provides very useful data for researchers aiming to analyze whole mosquito genomes; especially when sequencing is still expensive and having a reliable DNA extraction protocol is essential. The abstract, justification, protocols and results are very well described.

Some comments:
• Did the preservation of individual mosquitoes – 80% alcohol or -20°C – affect the DNA yield? I am not sure if this was an additional variable. Please clarify.

• Mention the number of specimens you used for each combination of lysis and extraction methods.

• Table 1: Do ‘X, 2L, 2R, 3L, ..chrM’ columns correspond to mean coverage’s? Please add a brief description of these terms in the table legend.

• Discussion: the statement ‘the amount of magnetic beads…had little effect on DNA yield’. This variable was not mentioned in the methods section. Please add, mention specific bead amounts tested, sample size and statistical test.

• Table 2: Please mention in this table (or table 1) what ‘Mt, 2R….X’ stands for. Are ‘Mt and chM’ mitochondrial DNA?

• Table 2: State the difference between A. gambiae 1 and A. gambiae 2 groups.

• Table 2: There are significant differences between the genotype-calls in the WGA and original DNA. Would you suggest having a third method to validate your genotypes? Maybe regular DNA sequencing. I think this could add to your discussion.

• Table 3: Add a column with your sample sizes.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.
Thank you for your positive feedback. I will try to get the formal revision done in the near future.

For your first question, we typically use 80% for storing samples in Africa. Due to hot environmental conditions there we were expecting accelerated evaporation of ethanol which could make ethanol content lower than what we intended (70%). So we put extra to compensate the loss. We have tried this protocol on samples stored dry in silica, dry frozen at -20°C or 100% ethanol as well. If samples are stored in high ethanol content, they get harder than 70-80% ethanol. But this doesn’t affect DNA extraction because we do hydrate in water for an hour anyway to soften up the tissue except frozen ones.

We have not detected any decrease in yield for relatively fresh material (< 2 months). Quality goes down for samples stored without alcohol (either in silica or frozen) resulting in lower yield in target species DNA and lots of microbial content, requiring more sequencing reads to get the same coverage.

**Competing Interests:** No competing interests were disclosed.
We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

**Competing Interests:** No competing interests were disclosed.

**Referee Report 17 December 2015**

doi:10.5256/f1000research.7988.r11616

Beniamino Caputo¹, Verena Pichler²

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The article is an interesting update on extraction protocols and highlights the importance of testing different methods when extracting DNA for a wide range of downstream applications. The article’s title is sound and describes well the observations made, the same is true for the abstract which provides an adequate summary of the article. Also the aim of the article and the study design are well described and allow to reproduce the different methods tested. Results and discussions appear to be sound and coherent with the offered data.

There are only few minor corrections to make:

**In Methods:**
- please correct TissuLyser in TissueLyser
- specify the number of specimens analyzed

**In Results and Discussion:**
- in the second paragraph it might be noteworthy that genotype call errors do not appear only when considering the mitochondrial DNA, but, at least in *A.gambiae* listed in Table 2, many errors, such as opposed genotype calls can be observed also for autosomal sequences

**Table 1:**
- it could be useful to add some information in the caption such as the meaning of ChrM (I suppose it stands for mitochondrial DNA?) and units of measurements for the given data
- please correct *A.gambia* in *A.gambiae* (1st column)

We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

**Competing Interests:** No competing interests were disclosed.