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# Analysis of Alternative Storage Conditions for DNA Recovery from Field Samples

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## Analysis of Alternative Storage Conditions for DNA Recovery from Field Samples

### Abstract

As ecologists increasingly employ molecular methods, they find that tried and true preservation solutions (e.g. ethanol or formalin) may not be optimal when samples are targeted for genetic analyses. Before traveling to remote sample sites, researchers need to consider which preservation methods are likely to yield the largest quantity and highest quality of DNA based on their travel times and field conditions. They also need to consider whether they will have access to preservatives at remote sites and whether those preservatives can be safely transported. To determine which preservation methods would most reliably preserve tissue for genetic analysis under a range of field conditions, we examined total DNA recovery from female fruit flies (Drosophila melanogaster) individually held in various solutions (70% ethanol; 2% SDS, 100 mM EDTA; 1% SDS, 50 mM EDTA; 0.66% SDS, 33 mM EDTA; Zymo© lysis buffer; Zymo Xpedition© lysis buffer) at three different temperatures (22oC, 4oC and -20oC) for varying lengths of time (1 day, 4 weeks, and 8 weeks). We predicted that insects held in Zymo Xpedition<sup>©</sup> buffer would yield the overall highest DNA recovery since this buffer was designed for field collected animal tissue. We also predicted that variation in DNA recovery from insects held in different solutions would increase with preservation time and holding temperature. Although we observed significant differences in total DNA recovery from some of our samples, no trends were identified. Preliminary band quality analyses of PCR products utilizing stored DNA as template for amplification of the mCOI gene generally indicated decline in product quality as storage time increased. Future work will focus on better quantifying stored DNA quality and examining the relationship between total DNA recovered and overall DNA quality.

### About the Author

Authors Alison Schutt, Emily Stricklin, Britta Ten Haken, and Joseph Tolsma are currently students at Northwestern College.

An environmental scientist, Dr. Furlong holds a doctorate in ecology, evolution and marine biology. Her research and publications have been in the fields of stream ecology, entomology and biogeography, and she has experience as a biological consultant. Furlong teaches Introduction to Environmental Science, General Biology, Invertebrate Zoology and Aquatic Ecology. She has also been a stream ecology instructor for the Creation Care Study Program in Belize.

Dr. Tolsma's research efforts in cancer genetics and cell biology have been widely published in scientific journals and have received a number of awards. A Northwestern College alumna, she holds a doctorate in microbiology/immunology/virology from Northwestern University. During a sabbatical, she worked on a cell and molecular genetics textbook for non-science majors, as well as an adult Sunday school curriculum on genetic technologies. She has been a part of several symposia on bioethical issues surrounding genetic technologies, such as stem cells, cloning and genetic testing. Her current laboratory research extends her interest in genetics to populations in a study of mayfly genetic variation in Northwest lowa watersheds and her interest in cancer cell biology with a project that measures the anti-proliferative effects of plant extracts and chemicals in those extracts on human tumor cells in vitro. She is the 2015 recipient of Northwestern's annual Teaching Excellence Award.

### Authors

Alison Schutt, Emily Stricklin, Britta Ten Haken, Joseph Tolsma, Laurie Furlong, and Sara S. Tolsma

# Analysis of Alternative Storage Conditions for DNA Recovery from Field Samples

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### Abstract

As ecologists increasingly employ molecular methods, they find that tried and true preservation solutions (e.g. ethanol or formalin) may not be optimal when samples are targeted for genetic analyses. Before traveling to remote sample sites researchers need to consider which preservation methods are likely to yield the larges quantity and highest quality of DNA based on their travel times and field conditions. The also need to consider whether they will have access to preservatives at remote sites an whether those preservatives can be safely transported. To determine which preservation methods would most reliably preserve tissue for genetic analysis under a range of field conditions, we examined total DNA recovery from female fruit flies (Drosophil melanogaster) individually held in various solutions (70% ethanol: 2% SDS, 100 mM EDTA 1% SDS, 50 mM EDTA; 0.66% SDS, 33 mM EDTA; Zymo© lysis buffer; Zymo Xpedition@ lysis buffer) at three different temperatures (22°C, 4°C and -20°C) for varying lengths of time (1 day, 4 weeks, and 8 weeks). We predicted that insects held in Zymo Xpedition buffer would yield the overall highest DNA recovery since this buffer was designed for field collected animal tissue. We also predicted that variation in DNA recovery from insects hele in different solutions would increase with preservation time and holding temperature Although we observed significant differences in total DNA recovery from some of our samples, no trends were identified. Preliminary band quality analyses of PCR product utilizing stored DNA as template for amplification of the mCOI gene generally indicate decline in product quality as storage time increased. Future work will focus on bette guantifying stored DNA guality and examining the relationship between total DNA recovered and overall DNA quality.

#### Introduction

With molecular biological techniques rapidly becoming a standard component of ecological studies, scientists are rethinking strategies for specimen preservation in the field Fraditionally, it was most important to use a preservative solution that maintained structura characteristics but, if molecular biological analysis is the goal, structure is less importa than choosing a preservative solution that maintains the integrity of specimen DNA. Ethang solutions (typically 70%) and formaldehyde preserve specimen structural integrity and also may protect the integrity of DNA but are difficult to obtain at remote sites and can b problematic to transport due to their flammability and toxicity. Pokluda et al. (2014 proposed alternative preservative solutions containing various concentrations of sodiun dodecyl sulfate (SDS) and ethylenediaminetetraacetic (EDTA) acid. Their paper provider ntriguing evidence that these solutions may be as effective at preserving insect DNA as 70% ethanol, however their experimental design did not allow quantitative evaluation of DNA yield. To obtain clearer evidence that alternative preservative solutions were as o more effective than ethanol at maintaining DNA integrity in insect specimens, we designed a quantitative experiment. We reared genetically identical (wild type), female Drosophil melanogaster as a uniform organism and placed those organisms in six different solution (70% ethanol; 2% SDS, 100 mM EDTA; 1% SDS, 50 mM EDTA; 0.66% SDS, 33 mM EDTA Zymo© lysis buffer; Zymo Xpedition© lysis buffer), at three different temperatures (22°C 4°C and -20°C), for three different times (1 day, 4 weeks, and 8 weeks), After isolating DN we measured the amount of DNA recovered using a Nanodrop 2000 © Spectrophotomete We also measured the quality of the DNA by determining if it was amplifiable by PCR an ssessing band quality of the amplified PCR product.

#### Materials and Methods

#### Specimen Selection and Storage

Wild-type Drosophila melanogaster were offspring of those purchased from Carolina Biologicals (Burlington, NC Wite-type Urosopniai meianogaster were oftspring of those purchased from Caroina biologicas (duringion, NC). Files were ansethetized with FNApe (accimica biologicas), Burnington, NC). Finaré litis were selected, and place in microtoge tubes containing 70% ethanol, 2% SDS 100 mM EDTA, 1% SDS 50 mM EDTA, 68% SDS 33 mM EDTA, zymo Xpedrilom3 (JusiStatizian Buffer (Zymo Research, Invine, CA), or Lysis Buffer (ZR Tissue and Insect DNA Prep, Zymo Research, Invine, CA). Specimens, in solution, were placed at -20°C, 4°C, or 22°C for 24 hours, 4 weeks, or 8 weeks.

#### DNA Isolation

UNA Isociation After storage for the allotted time, DNA was isolated from the *Drosophila melanogaster* specimens using ZR Tissu and Insect DNA MiniPrey (Zymo Research, Invine, CA) for all specimens except those in Xpedition Lysis/Stabilization Buffer, DNA from specimens in Xpedition Buffer was isolated using Xpedition Tissue and Insec MiniPrey (Zymo Research, Invine, CA). Purified DNA was quantitated by measuring the £20/250 warelength risk with a NanoDrop 2000 UV-Vis Spectrophotometer (Thermo Scientific, Grand Island, NY). Significant differences total DNA amounts from samples held under varied storage conditions were determined by one-way ANOVAs ar significant subsets with the Tukey HSD Post Hoc Test.

#### Amplification by PCR

710-bp fragment of the mCOI gene was amplified using the primers LCO1490 (5 A /10-bp fragment of the mCUI gene was amplied using the primes LCU1440 (s GGTCAACABATCATAGASTATCGS) and HCO2189 (STAACCTCAGGGTGACCAAAAAATACT3) (Folmer, al, 1994), PCR conditions used were 60 ng of DNA, 2.5 uM MgC, 0.5 mM dNTP mix (Promega, Madison, WI), 1 Go-Tap Ubfer, 1 JL Go-Tap optimesa (Promega, Madison, WI) in a total valume of 50 µL Reactions wer amplified in 35 cycles at the following temperatures: 1 mixtue at 95°C, 1 mixtue at 92°C, 15 mixtue at 72°C wit a final actension of 10 minutes at 72°C followed by holding the samples at 42°C. FRO motocuts were visualized by oading 20 µl of the PCR reaction on 1% agarose gels run in TAE buffer at 100 mV. Bands were imaged using Bio-Rad Chemidoc MP Imager (Bio-Rad, Hercules, CA).

#### Estimation of DNA Quality via Band Analysis

Camilation to Drive scalarly via band relations of an got immediate DNA was used in each PCR reaction. DNA quality was estimated by analyzing the quantity PCR product producet. Five different individuals ranked band intensity relative to the molecular weight standard from 0 (no band) to 3 (more inferes than molecular weight standards). Differences in band quality at differe orage times (1 day vs. 4 weeks) were statistically ass

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Results

quality with greater storage time for most treatments (Figure 3). Regardless o storage conditions our DNA template produced a PCR product with an intensity that suggests it is could be sequenced more than 50% of the time. Template isolated from specimens stored in various solutions, regardless of time or temperature, produced quality PCR products between 63 and 75% of the time. When considering on emperature samples stored at -20°C resulted in template DNA that produce sequence-guality bands 80% of the time. That percentage dropped to 66% at 40 and 53% at 20°C. Samples stored for only 1 day resulted in PCR products that wer sequence-quality 79% of the time while extending storage to 4 weeks reduced that to

Total DNA from Samples Stored at -20°C

1%505 0.66% SDS

Storage Solutions

Total DNA from Samples Stored at 4°C

H-

0.66% SD:

Storage Solutions

Total DNA from Samples Stored at 22°C

0.66% SDS

1%SDS

Storage Solutions

Figure 1, Mean total DNA isolated from Drosophila melanogaster (n=5) stored a

three different temperatures, for three different lengths of time, in six different solutions (ETOH = 70% Ethanol; 2% SDS = 2% SDS, 100 mM EDTA; 1% SDS =

1% SDS, 50 mM EDTA; 0.66% SDS = 0.66% SDS, 33 mM EDTA; Zvmo = Zvmo@

lysis buffer, Expedition = Zymo Xpedition© lysis buffer). Within each solution type significantly different means are indicated by different letters (ANOVA p<0.05,

Tukey HSD p<0.05). Standard deviations are represented by error bars.

**2**24 k

P 1500

2 1500 2 1500

E 1000

200

a 150

ETOH 2% SDS

Mean

FTOP



Figure 2. Mean of total DNA after 8 weeks of storage at 3 temperatures in different storage solutions (ETOH = 70% Ethanol; 2% SDS = 2% SDS, 100 mM EDTA; 1% SDS = 1% SDS, 50 mM EDTA; 0.66% SDS = 0.66% SDS, 33 mM EDTA; Zvmo Zymo© lysis buffer, Expedition = Zymo Xpedition© lysis buffer). Total DNA differed significantly in 2%SDS 50mM EDTA (ANOVA p<0.05). The significantly samples are indicated by different letters (Tukey HSD p<0.05). Standard deviations are represented by error bars.



Figure 3. Mean band quality of PCR products stored in various solutions generally declined with increasing storage time (1 day vs. 4 weeks), however differences were rarely significant. Significantly different means are indicated with asterisks (Ttest p< 0.05). Standard deviations are indicated by error bars. Storage solutions: ETOH = 70% Ethanol; 2% = 2% SDS, 100 mM EDTA; 1% = 1% SDS, 50 mM EDTA; 0.66% = 0.66% SDS, 33 mM EDTA; Zymo = Zymo© lysis buffer. Expedition = Zymo Xpedition© lysis buffer.

Funding for this project was provided by Northwestern College Scholarship Grants (Tolsma) and a Northwestern College Endowed Research Fellowship (Furlong) We are grateful to Steve Bogaard and the Biology and Chemistry departments at Dordt College for allowing us to use their NanoDrop 2000 Spectropho The 2015 BIO 202 class provided skilled hands during a particularly busy DNA isolation day.

Figure 4. Representative PCR products amplified from Drosophila melanogaster DNA ated after specimens were stored under various conditions. Gel 1: Samples stored in Zymo lysis buffer at -20°C for 24 hours. Lane 1: no DNA control: Lanes 2-6: replicate samples. Gel 2: Samples stored at 4°C for 24 hours. Lane 1: no DNA control; Lane 2: 70% ethanol; Lane 3: 2%SDS, 100 mM EDTA; Lane 4: 1% SDS, 50 mM EDTA; Lane 5: 0.66% SDS, 33 mM EDTA; Lane 6: Zymo Expedition Buffer; Lane 7: Zymo lysis buffer. Gel 3: Samples stored in Zvmo Xpedition Buffer at -20°C for 24 hours. Lane 1: no DNA control Lanes 2-6; replicate samples, Lanes 7, 8, and 7 represent MW standards.



#### Discussion

Our purpose in tackling this project was very practical. We were having problems finding ethanol protable preservation options and that might yield better results in our genetic analyses. After readin Pokula et al. (2014), we were eager to try some of their suggested alternatives. In addition, we wer curious about buffers offered by Zymo®. However, if these alternatives are "good," we wanted t quantify "how good" before going out into the field again. Using replicates from a very uniform population (lab-reared, female, wild-type Drosophila melanogaster), we hoped to eliminate some of the variabilit inherent in "wild caught" insects. By comparing and quantifying total DNA and quality of DNA yield fro specimens held in various storage solutions and under a variety of storage conditions, we hoped to fin the best options for our own field collections.

We predicted that, regardless of the preservation solution, the quantity and quality of DNA recovered from our samples would decline with increasing storage time and increasing temperature. In addition, we assumed that the Zymo® Xpedition lysis buffer would give the best results overall since it was advertised as specially formulated for field preservation under a wide range of conditions.

The DNA recovered from our experiment was remarkably variable in quantity (Figures 1 and 2) and although some trends are evident, very few significant differences in treatment were observed. We saw evidence that ethanol preservation gives mediocre DNA recovery under a variety of conditions, but so do the other preservation fluids. In particular, we noted that the Zymo® Xpedition lysis buffer did not provid superior results. We also noticed that, while increasing storage time may result in decreased yield, thi was not always the case (Figure 1). Likewise, lower storage temperature tended to give higher, by arely significantly higher, yields (Figure2). We believe that variation in sample processing (tissu bashing) may mask the treatment effects.

Since the goal of our own research is the recovery of high quality DNA for amplification and sequencing we found the results of the band analyses particularly interesting (Figure 3) and plan to expand this pa of our study to include eight week samples. Furthermore, our DNA quality assessment is somewhat subjective and we plan to quantify it more precisely. We will measure the concentration of PCB product plified from the same amount of template DNA (60 ng) with the NanoDrop 2000 Spectrophoto and quantify the bands amplified using densitometry measurements of bands with the Biorad ChemiDo MP Imager. This will allow us to more fully explore the relationship between quantity and quality of DN solated from specimens held in under various conditions.

Additionally, we wonder if our highly variable DNA quantities were related to the extraction procedure rather than the preservation of our specimens. We experimented with various extraction procedure including bashing specimens with Zymo© bashing beads by hand, using a Disrupter Genie©, vortexin by hand, and shaking bashing bead tubes in a rack. Visual examination of the specimens after different disruption methods suggested that this step was highly variable. We are working on increasing th consistency of this step in the DNA isolation process before we repeat this experiment on more variable populations of insects

As our field season approaches, we know that we need to keep our specimens cold and process them i a timely manner for best results. However, we also now know that a few days at room temperature w not doom our efforts and that we can confidently use inexpensive and portable alternatives to ethanol.



