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© DNA Isolation from Reptile Blood using Gentra Puregene (Qiagen) DNA Isolation Kit

COMMENTS 0

DOI

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### **ABSTRACT**

WORKS FOR ME

This protocol is used to isolate DNA from reptile blood (either whole or red blood cell pellets). Reptile blood has nucleated red blood cells and will produce considerably more DNA from the same volume of mammalian blood. This protocol is adapted from Gentra Puregene DNA Isolation A Kit.

The time estimates assumes you are processing 24 samples and you are well practiced.

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PROTOCOL CITATION

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**KEYWORDS** 

DNA isolation, reptile blood, reptile whole blood

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### **Equipment**

- Pipettes (p1000, p200, p20)
- Shaking tube incubator
- Benchtop Centrifuge

### Consumables

- One 15 ml tube (DNA/RNA and DNAse free) to make 70% ETOH
- 1.5-mL microcentrifuge tubes (VWR Catalog Number 76332-068). Two tubes for each sample
- Filtered pipette tips (p1000, p200, p20)
- Serological pipette (10 or 20 ml) to make 75% ETOH
- latex or nitrile gloves

### Reagents

- Qiagen Puregene Core Kit A (Catalog Number 158467)
- ·Cell Lysis Buffer
- ·Protein Precipitation Solution
- ·DNA Hydration Solution
  - Proteinase K (20 mg/mL) (IBI Science, Product Number IB05406)
  - Isopropanol (ThermoFisher Scientific, Catalog Number AC327272500)
  - 70% ethanol (200 proof for dilutions: Deacon Labs Product Number 3916EA; Molecular Grade Water)
  - Molecular Grade Water (Quality Biological, Catalog Number 351-029-131)
  - 1.5 mL microcentriguge tubes (VWR Catalog Number 76332-068)

# Notice for high quality (long undamaged DNA)



\*Through out this protocol we **vortex** the samples for mixing. This works well and produces alot of good quality DNA that works for most everything. If you want to minimize damage to the DNA such as for long-read sequencing on the Nanopore or PacBIO, or assays to quantify DNA damage) **do not vortex**. Rather mix gently by flicking and inverting the tube. The quality will be higher but generally less yield.

## **Setting Up**

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3 Make fresh 70% ETOH. For example, to make 15 ml in a 15 ml tube use a 10 ml sterile pipette to take

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10m

△ 10.5 mL of IMI 100 % volume (200 proof) ETOH, and another 10 ml sterile pipette to add △ 4.5 mL of molecular grade water.

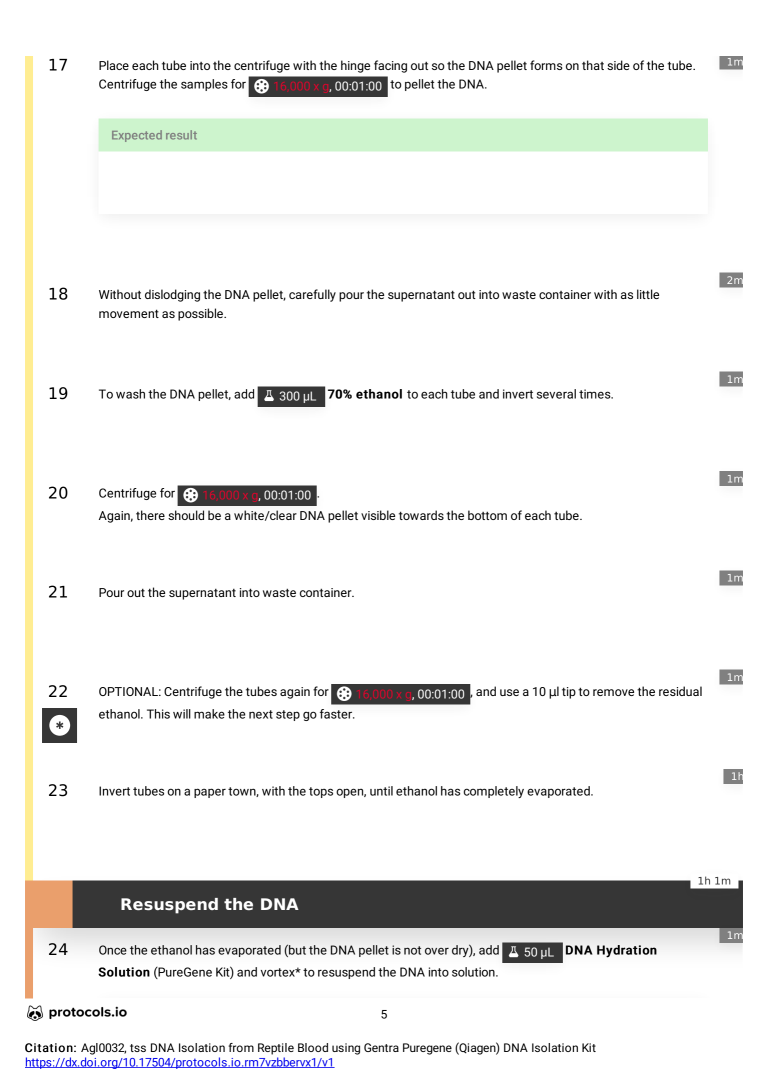
		1h 45m
	Red Blood Cell Lysis	
4	Set out the the blood samples to be used on ice to thaw, but stay cold.  Set out the correct number of 1.5 mL tubes into a clean tube rack.	5m
5	Add 300 µL of <b>Cell Lysis Solution</b> (from PureGene Kit) to each tube.	1m
	Note	
6	Add I 10 µL proteinase K (20 mg/ml) to each tube.	1m
7	Close tubes and label each tube according to Sample IDs.	3m
8	Add 15 µL of <b>whole blood</b> to the correct tube and <i>vortex well*</i> .	30m
	Note	
	Note	

9 Place each tube in the shaking incubator and incubate at Vortex\* the samples periodically.

1h

1h 33m

**Isolate DNA** Add 🗸 100 µL of **Protein Precipitation Solution** (from PureGene Kit) to each tube and vortex\* 10 vigorously. 5m 11 Place in ice for approximately 5 minutes. During this incubation do steps 12 and 13. 12 Obtain fresh 1.5 mL tubes and label top and side with DNA ID, Date, and "stock DNA" (these will be the tubes that the isolated DNA will be stored in). 2m 13 Use a pipette to add  $\perp$  300  $\mu$ L of **isopropanol** into each of the newly labeled tubes. 14 Centrifuge the samples for 00 x g, 00:01:00 After removing the tubes from the centrifuge, there should be a yellow/brown pellet at the bottom of the tube. If there is no pellet, repeat this step for each tube missing the pellet. 1m 15 Without dislodging the protein pellet, pour out the supernatant containing the DNA into the newly labeled tubes containing the isopropanol. Throw away the original tube with the protein pellet. 16 Mix the supernatant with the isopropanol by inverting 50 times. If there is a lot of DNA you can see the strands condensing at this step (looks like thin white threads).





Incubate the tubes for one hour at room temperature, or 4°C overnight to resuspend the DNA.

### 1h

## **Check Quality and Quantity**

Run 5 µL of resuspended DNA on 1% agarose gel to visual the quality and estimate quality. DNA can be quantified with the Nanodrop. For sensitive procedures (DNA sequencing library preparation we recommend using the Agilent TapeStation or BioAnalyzer).