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© Characterization of the Archaeome, Bacteriome and Eukaryome in Nasopharyngeal Swabs

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ABSTRACT

This protocol describes the Characterization of the Archaeome, Bacteriome and Eukaryome in Nasopharyngeal Swabs by sequencing with nanopore technology.

For a long time, archaea were under-represented in the literature, and less is known about their pathogenicity in human diseases. Using conventional methods, the cultivability particularly of archaea is challenging and they are still classified as the 'dark matter' of the microbiome. The evolution of advanced sequencing techniques in the twenty-first century, a strong focus on archaea research is interestingly observed. However, the influence on disease course or even pathogenesis in terms of respiratory disorders remain unexplored. Thus, more attention has to be paid on the characterization of the archaeome with the goal of translation into clinical contexts. Considering this important issues lacking good methodological reports in the literature, we evaluated previously developed primer sets and sequencing platforms. With these useful hints, we share potential alternative procedures with the aim how to increase the quality of research on archaeome and eukaryotes. The use of nasopharyngeal swab specimens derived from a cohort suffering from respiratory diseases enable to study translational aspects on disease course and eventually pathogenesis. The optimization of 'pre-sequencing' steps, starting from the DNA isolation, amplification, right choice of sequencing platforms e.g., MinION Oxford Nanopore rule some important traces to a high-qualitative in-depth sequencing success. However, those descriptive data significantly contribute to optimize existing archaic models with the aim to exploit translational approaches ex vivo.

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MATERIALS TEXT

DNA isolation:

QIAmp DNA-Mini Kit by Qiagen (Qiagen 51304)

PCR:

Q5 HotStart High Fidelity 2x MM (NEB M0492)

Library Preparation + Sequencing:

NEBNext FFPE DNA Repair Mix (NEB M6630)

NEBNext Ultra II End Repair/dA-Tailing Module (NEB E7546)

NEBNext Quick Ligation Module (NEB E6056)

Agencourt AMPure XP (Beckman Coulter A63880)

Long Amp Tag Polymerase MM (NEB M0323)

PCR Barcoding Expansion 1-96 (ONT EXP-PBC096)

Ligation Sequencing Kit (ONT SQK-LSK110)

DNA Isolation

2h

1 DNA was isolated from nasal swabs with amies medium, using approximately $\Box 1000~\mu L$. For the Isolation, the QIAmp DNA-Mini Kit by Qiagen was used, following the QiAamp tissue protocol from the 'QIAamp DNA Mini and Blood Mini Handbook 05/2016'

1.1



10m

Centrifugation of sample at **37500 rpm**, **00:10:00** until pellet formation.

1.2

5m

Resuspension of pellet in ■180 µL ATL

1.3



10m



3

Adding 20 µL Proteinase K, vortexing,

Incubate at § 56 °C until complete lysis. Occasionally vortexing.

1.4

2m

Brief Centrifugation of the sample.

1.5

15m

1.6

6m

Adding $\square 200 \ \mu L$ Ethanol (96-100%), and mix by pulse-vortexing for $\bigcirc 00:00:15$. Afterwards, short centrifugation of the sample.

1.7

5m

Transfer mixture (including precipitate) to the QIAamp Mini spin column. CAVE: without wetting the rim. Centrifugation: §8000 rpm, 00:01:00 Replace the QIAamp Mini spin column, use a clean 2 ml collection tube, Discard tube with the filtrate.

1.8

5m

Adding $\Box 500 \mu L$ Buffer AW1 CAVE: without wetting the rim.

Centrifuge: **8000 rpm, 00:01:00**.

Replace the QIAamp Mini spin column, use a clean 2 ml collection tube, Discard tube with the filtrate.

1.9

8m

Add **□500 µL Buffer AW2** to the QIAamp Mini spin column without wetting

the rim. Closing of the column, Centrifugation:

314000 rpm, 00:03:00, Centrifugation at full speed

1.10



Replace QIAamp Mini spin column with a new 2 ml collection tube. Discard tube with the filtrate. Centrifugation:

314000 rpm, 00:01:00, Centrifugation at full speed

1.11



7m

2m

Placing QIAamp Mini spin column in a new 1.5 ml microcentrifuge tube. Discard tube with the filtrate.

Add $\blacksquare 100~\mu L$ AE . Incubation at room temperature ($\bigcirc 00:05:00$), centrifugation: \$8000~rpm, 00:01:00

- 1.12 Repeat Step 1.11: Add the flowthrough of the previous step to the Mini spin column and incubation at room temperature for © 00:05:00, centrifugation \$8000 rpm, 00:01:00 at .
- 1.13 Concentration measurement with nanophotometer or qubit.

5m

PCR

2h 30m

2 PCR

Archaea: Nested PCR Eukaryotes: single PCR

2.1 Primer selection for archaea and eukaryotes

Α	В	С	D
Nr.	Name	Primer Name	Sequence (5' ♦ 3')
1	344F	S-D-Arch-0344- fw	5'-acggggygcagcaggcgcga-3'
2	1041R	S-D-Arch-1041- rev	5'-ggccatgcaccwcctctc-3'
3	519F	Arch-519F-Tag	5'-tttctgttggtgctgatattgccagcmgccgcggtaa-3'
4	786R	Arch-786R-Tag	5'-acttgcctgtcgctctatcctcggactacvsgggtatctaat-3'
5	563F	Euk-563F-Tag	5'-tttctgttggtgctgatattgcgccagcavcygcggtaay-3'
6	1132R	Euk-1132R-Tag	5'-acttgcctgtcgctctatcttcccgtcaatthcttyaart-3'

10m

2.2

1st PCR Mix:

■8 µL nuclease free water

■12.5 µL Q5 Polymerase

■2 µL Primer Mix

■2.5 µL DNA-Template

2.3 **\(\sigma\)**

PCR-Run. 1

Primer pair Arch-344-F-1041R / Eck. 563F-1132Rtag

Heated Lid: 110 C

Denaturation § 95 °C © 00:03:00

Cycles (30):

Denaturation § 95 °C © 00:00:30

Elongation § 72 °C © 00:00:30

End Cycle

Final Elongation § 65 °C © 00:05:00

2.4 10m

2nd PCR (archaea only) Mix:

■9.5 µL nuclease free water

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□12.5 μL Q5 Polymerase□2 μL Primer-Mix□1 μL DNA-Template

50m

2.5

PCR-Run. 2 (nested)

Primer pair Arch-519-F-786Rtag

Heated Lid: 110 C

Denaturation § 95 °C © 00:03:00

Cycles (28):

Denaturation § 95 °C ⊙ 00:00:30

Annealing § 55 °C © 00:00:30

Elongation ₹ 72 °C ♦ 00:00:30

End Cycle

Final Elongation § 65 °C © 00:05:00

2.6 Gel Electrophoresis

1h 30m

Check, if the wanted sequences were amplified. Ether through classic gel electrophoresis or through microcapillary gel electrophoresis.

Library preparation + sequencing:

10m

3



Library Preparation + Sequencing:

- 1st Purification
- PCR preparation
- 2nd Purification
- Concentration measurement

3.1

10m

1st Purification:

Add $\blacksquare 36~\mu L$ Beats AMPure XP and apply an external magnetic field for $\bigcirc 00:05:00$. Afterwards discard fluid supernatant.



7

3.2 Add \blacksquare 150 μ L Ethanol 70% and discard fluid supernatant.

8m

3m

3.3 Add another $\Box 150~\mu L$ Ethanol 70% . Afterwards discard the fluid supernatant and dry tube with open lid

5m

3.4 Resuspend pellet in \Box 15 µL nuclease free water

JII

3.5

10m

PCR-preparation: Mix

■12.5 µL Long Amp Tag Polymerase MM

■2 µL sample

■9.5 µL nuclease free water

■1 µL Barcode

30m

3.6

Heated Lid: 110 C

Denaturation ₹ 95 °C (© 00:03:00

Cycles (18):

Denaturation ₹ 95 °C (© 00:00:15

Annealing § 62 °C © 00:00:15

Elongation № 65 °C (© 00:00:45

End Cycle

Final Elongation § 65 °C © 00:05:00

3.7

2nd Purification

- 3.8 Add \square 36 μ L Beats and apply an external magnetic field for \bigcirc 00:05:00 . Afterwards discard fluid supernatant
- 3.9 Add **150 μL Ethanol 70%** and discard fluid supernatant.
- 3.10 Add another **150 μL Ethanol 70%** . Afterwards discard the fluid supernatant and dry tube with open lid
- 3.11 Resuspend pellet in **15 μL nuclease free water**
- 3.12 Concentration measurement with nanophotometer 5m
- 3.13 Library preparation:

We used a modified version of the PCR barcoding (96) genomic DNA (SQK-LSK109) protocol by Nanopore.

Quantify the barcoded library using a nanophotometer and pool all barcoded libraries in the desired ratios in a 1.5 ml DNA LoBind Eppendorf tube.

3.14 \(\sqrt{10m} \)

Prepare $\blacksquare 1 \mu g$ pooled barcoded libraries in $\blacksquare 47 \mu L$ nuclease free water .

- 3.15 DNA repair and end-prep
- 3.16 Thaw DNA CS (DCS) at RT, spin down, mix by pipetting, and place on ice. 5m

3.17 Prepare the NEBNext FFPE DNA Repair Mix and NEBNext Ultra II End repair MA-tailing Module reagents in accordance with manufacturer's instructions, and place on ice.

3.18

10m

In a 0.2 ml thin-walled PCR tube, mix the following:

■1 µL DNA CS

■47 μL DNA

■3.5 µL NEBNext FFPE DNA Repair Buffer

■2 µL NEBNext FFPE DNA Repair Mix

■3.5 µL Ultra II End-prep reaction buffer

■3 µL Ultra II End-prep enzyme mix

Mix gently by flicking the tube, and spin down.

3.19

10m

5m

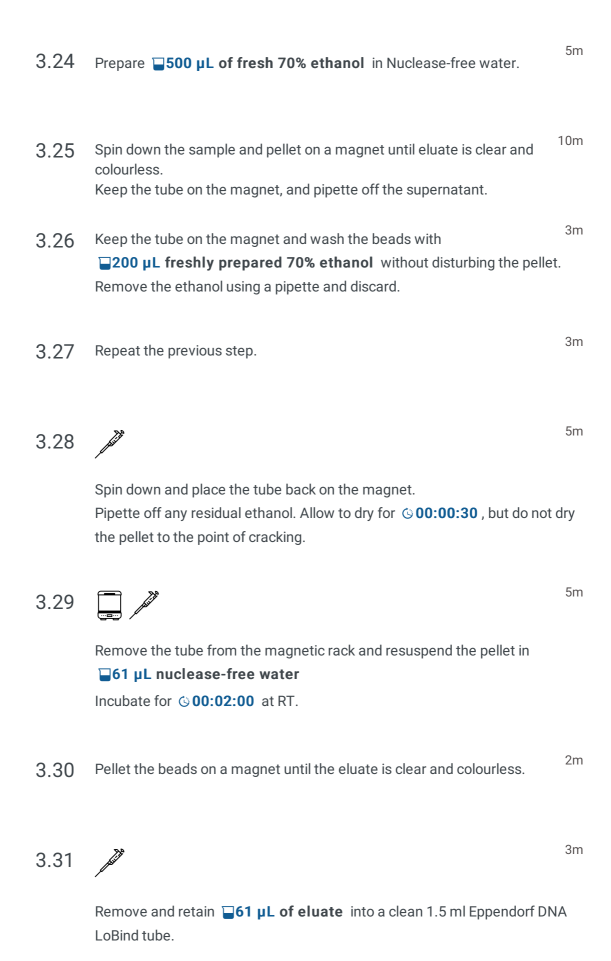
Using a thermal cycler, incubate at & 20 °C for \circlearrowleft 00:05:00 and & 65 °C for \circlearrowleft 00:05:00

3.20

AMPure XP bead clean-up

- 3.21 Resuspend the AMPure XP beads by vortexing.

 Transfer the DNA sample to a clean 1.5 ml Eppendorf DNA LoBind tube.
- 3.22 Add **□60 μL of resuspended AMPure XP beads** to the end-prep reaction and mix by flicking the tube.
- 3.23 Incubate on a Hula mixer (rotator mixer) for **© 00:05:00** at room temperature.



- 3.32 Take forward the repaired and end-prepped DNA into the adapter ligation step. However, at this point it is also possible to store the sample at § 4 °C overnight.
- 3.33 Adapter ligation and clean-up (PCR barcoding (96) genomic DNA (SQK-LSK109) protocol by Nanopore)

Although the recommended 3rd party ligase is supplied with its own buffer, the ligation efficiency of Adapter Mix (AMX) is higher when using Ligation Buffer supplied within the Ligation Sequencing Kit.

- 3.34 Spin down the Adapter Mix (AMX) and Quick T4 Ligase, and place on ice. 1m
- 3.35 Thaw Ligation Buffer (LNB) at RT, spin down and mix by pipetting.

 Due to viscosity, vortexing this buffer is ineffective. Place on ice immediately after thawing and mixing.
- 3.36 Thaw the Elution Buffer (EB) at RT, mix by vortexing, spin down and place on ice.
- 3.37 To retain DNA fragments of < 3 KB, thaw one tube of Short Fragment Buffer (SFB) at RT, mix by vortexing, spin down and place on ice.
- 3.38 In a 1.5 ml Eppendorf DNA LoBind tube, mix in the following order:
 - ■60 µL DNA sample from the previous step
 - ■25 µL Ligation Buffer (LNB)
 - ■10 µL NEBNext Quick T4 DNA Ligase
 - ■5 µL Adapter Mix (AMX)

Mix gently by flicking the tube, and spin down.

3.39 Incubate the reaction for **© 00:10:00** at RT. If you have omitted the AMPure purification step after DNA repair and end-prep, do not incubate the reaction for longer than **© 00:10:00**.

5m

Resuspend the AMPure XP beads by vortexing. Add

 \blacksquare 40 μ L of resuspended AMPure XP beads to the reaction and mix by flicking the tube.

3.41 Incubate on a Hula mixer (rotator mixer) for © 00:05:00 at RT.

5m

- 3.42 Spin down the sample and pellet on a magnet. Keep the tube on the magnet, and pipette off the supernatant.
- 3.43 Wash the beads by adding **250 μL Short Fragment Buffer (SFB)**. Flick the beads to resuspend, spin down, then return the tube to the magnetic rack and allow the beads to pellet. Remove the supernatant using a pipette and discard.
- 3.44 Repeat the previous step.

5m

3.45

3m

Spin down and place the tube back on the magnet. Pipette off any residual supernatant. Allow to dry for © 00:00:30 but do not dry the pellet to the point of cracking.

3.46 🗍 🥻

10m

Remove the tube from the magnetic rack and resuspend the pellet in $\blacksquare 15~\mu L$ Elution Buffer (EB) . Spin down and incubate for 00:10:00 at RT.

3.48

2m

Remove and retain $\ \Box 15 \ \mu L$ of eluate containing the DNA library into a clean 1.5 ml Eppendorf DNA LoBind tube.

- 3.49 Quantify **1 μL of eluted sample** using a Qubit fluorometer. The prepared library is used for loading into the flow cell. Store the library on ice until ready to load.
- 3.50 The prepared library is used for loading into the flow cell. Store the library on ice until ready to load.
- 3.51 Priming and loading the SpotON flow cell
- 3.52 Thaw the Sequencing Buffer (SQB), Loading Beads (LB), Flush Tether (FLT)^{1m} and one tube of Flush Buffer (FB) at RT.

3.53 Im

Mix the Sequencing Buffer (SQB), Flush Tether (FLT) and Flush Buffer (FB) tubes by vortexing and spin down at RT.

3.54 **^**

Open the MinION Mk1B lid and slide the flow cell under the clip. Slide the priming port cover clockwise to open the priming port.

Take care when drawing back buffer from the flow cell. Do not remove more than 20-30 μ l, and make sure that the array of pores are covered by buffer at all times. Introducing air bubbles into the array can irreversibly damage pores.

After opening the priming port, check for a small air bubble under the cover. ^{1m}

3.55 Draw back a small volume to remove any bubbles (a few µl):

Set a P1000 pipette to 200 μl Insert the tip into the priming port

Turn the wheel until the dial shows 220-230 μ l, or until you can see a small volume of buffer entering the pipette tip

3.56

1m

To prepare the flow cell priming mix, add

■30 µL of thawed and mixed Flush Tether (FLT) directly to the tube of thawed and mixed Flush Buffer (FB), and mix by vortexing at RT.

- 3.57 Load **B00** μ**L** of the priming mix mix into the flow cell via the priming port, avoiding the introduction of air bubbles. Wait for **© 00:05:00**. During this time, prepare the library for loading by following the steps below.
- 3.58 Thoroughly mix the contents of the Loading Beads (LB) by pipetting because it contains a suspension of beads which settle very quickly. It is vital that they are mixed immediately before use!

3.59

2m

In a new tube, prepare the library for loading as follows:

■37.5 µL Sequencing Buffer (SQB)

■25.5 µL Loading Beads (LB), mixed immediately before use

■12 µL DNA library

3.60 Complete the flow cell priming through Gently lifting the SpotON sample port cover to make the SpotON sample port accessible. Load

■200 µL of the priming mix into the flow cell via the priming port (not the SpotON sample port), avoiding the introduction of air bubbles.

3.61

1m

Mix the prepared library gently by pipetting up and down just prior to loading.

3.62 /⁸

Add $\blacksquare 75~\mu L$ of sample to the flow cell via the SpotON sample port in a dropwise fashion. Ensure each drop flows into the port before adding the next.

- 3.63 Gently replace the SpotON sample port cover, making sure the bung enters the SpotON port, close the priming port and replace the lid.
- 3.64 If you using a MinION Mk1C turn basecalling while sequencing on.

3.65 Ending the experiment

After your sequencing experiment is complete, if you would like to reuse the flow cell, please follow the Wash Kit instructions and store the washed flow cell at $2-8^{\circ}$ C, OR

Follow the returns procedure by washing out the flow cell ready to send back to Oxford Nanopore.

Bioinformatics: 1d

4

4.1 If you were unable to basecall in real time, perform the basecalling now using the Guppy basecaller (newest version).

https://community.nanoporetech.com/docs/prepare/library_prep_protocols/Guppy-protocol/v/gpb_2003_v1_revag_14dec2018/guppy-software-overview

4.2 Now, using the resulting .fastq files, run the WIMP workflow from the Epi2Me software.

(https://nanoporetech.com/resource-centre/epi2me-wimp-workflow-quantitative-real-time-species-identification-metagenomic)

4.3 If the graphical output from the WIMP workflow is not sufficient for your analysis, you can download the results in a .csv dataset. Due to the size of this

dataset, further analyses may be performed by creating an SQL database.

The data contains the

- filename of the .fastq file
- Read ID --> is the unique primary key, wich enables to identify the read and therefore the sequence
- Run ID
- exit_status (of the read from the WIMP workflow)
- barcode
- taxID (every phylogenetic rank of each species has its own ID, with these IDs the lineage is composed
- name (of the organism)
- score
- lineage

4.4 Python scripts

While working on this project, a few Python scripts may be useful, depending on analysis you want to perform.

This script we used to split large files into smaller ones:

```
#IMPORTANT: this script must be started from the same
file directory as your input file!
filecounter=0
filelinecounter=0
inputfilename="file i want to split.txt" #set the
correct name of the file, you want to #split.
filename=inputfilename.split(".")[0]
file lines= open(inputfilename, 'r').readlines()
print(len(file lines))
while filelinecounter
outputfilename=filename+"_"+str(filecounter).zfill(3)+".
txt" #set #the correct ending for your file here
print(outputfilename)
while filelinecounter
outfile.write(file lines[filelinecounter])
  filelinecounter=filelinecounter+1
 else: filecounter=filecounter+1
```

This script was used, to append the lenght of each analysed read (or with

```
#IMPORTANT: this script must be started from the same
file directory as your input file!
# This script, the .fastq files from the run you want to
analyse and the WIMP.csv file must be in the same
directory!
inputfilename="WIMP inputfile.csv" #change the inputfile
here
import os
from multiprocessing import Pool
import concurrent.futures #imports the multithreading
library
import shutil
from pathlib import Path
filecounter=0
filelinecounter=0
i=1
# Define a function for the thread
def search fasta(WIMP inputline):
WIMP inputline=WIMP inputline.rstrip()
 fastqfilename=WIMP inputline.split("-",2)[0]+".fastq"
 #print(str(fastqfilename))
 readID=WIMP inputline.split(",",3)[1]
 #print(str(readID))
 fqfile=open(fastqfilename, 'r').readlines()
 #print("fqfile is open")
 #print(str(fqfile[0]))
 fqcounter=0
 found= False
 while found == False:
  fgreadID= fgfile[fgcounter*4].split()[0][1:37]
 #print(str(fqreadID))
 if (readID == fgreadID):
   readlenght=len(fqfile[fqcounter*4+1]) # if you want
to get the sequence instead of the lenght, remove the
len() function.
   #print(str(readlenght))
   found=True
  else: fqcounter=fqcounter+1
 completeline=WIMP inputline+","+str(readlenght)+"\n"
```

```
#print("Thread")
 return completeline
if name == " main ":
 dirname = os.path.join("C:/WIMPlenght tmp")
os.mkdir(dirname)
 filename=inputfilename.split(".")[0]
 print(filename)
 file lines= open(inputfilename, 'r').readlines()
print(len(file lines))
while filelinecounter
throughputfilename=filename+" "+str(filecounter).zfill(6
)+".csv"
 print(throughputfilename)
 while filelinecounter
outfile=open(dirname+"/"+throughputfilename, 'a')
   print(filelinecounter)
   outfile.write(file lines[filelinecounter])
   filelinecounter=filelinecounter+1
  #print(filelinecounter)
 else:
   filecounter=filecounter+1
   print("Filenumber: ", filecounter)
 print("tmpfiles complete")
 outputfilename=inputfilename.split(".")
[0]+" Output WIMP&Seqlenght.csv"
print(outputfilename)
while i < filecounter:
 tmpfilename=filename+"_"+str(i).zfill(6)+".csv"
 WIMP lines =
open(os.path.join(dirname+"/"+tmpfilename),
'r').readlines() #opens the tmp WIMP outputfile and
creates a list with each line as one item in the list
 p=Pool()
 with open(outputfilename, 'a') as outfile:
   result=p.map(search fasta, WIMP lines)
   p.close()
   p.join()
  #print(result)
   for f in result:
   #print(f)
    outfile.write(f)
```

```
print(i)
else:
  print("task complete")
  shutil.rmtree(dirname)
  print("tmpfiles deleted")
```

