

# Task 1: Phylogenetic Analysis of FAP Sequences

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## 1 Retrieving FAP Sequences via BLAST Search

### 1.1 Overview

To initiate the sequence alignment and subsequent phylogenetic analysis, 25 additional FAP sequences were retrieved from the NCBI database and saved as `FAP_BLAST.fas`. An R script was developed that retrieves the first 25 BLAST-aligned sequences by providing a query accession number, with results automatically saved to `data_raw/FAP_BLAST.fas`.

### 1.2 R Implementation

The following R script implements the BLAST query and sequence retrieval workflow:

```

# Load required packages
if (!requireNamespace("rentrez", quietly = TRUE)) {
  install.packages("rentrez")
}
if (!requireNamespace("xml2", quietly = TRUE)) {
  install.packages("xml2")
}
library(rentrez)
library(xml2)

# Set email for NCBI
Sys.setenv(ENTREZ_EMAIL = "mobofo3315@feralrex.com")

# Function to submit BLAST query and get RID and estimated wait time
submit_blast <- function(query_seq) {
  blast_url <- paste0(
    "https://blast.ncbi.nlm.nih.gov/Blast.cgi?CMD=Put&PROGRAM=blastp&DATABASE=nr&QUERY=",
    URLencode(query_seq)
  )
  res <- httr::GET(blast_url)
  content <- httr::content(res, as = "text", encoding = "UTF-8")

  rid <- sub(".*RID = ([A-Z0-9]+).*", "\\\1", content)
  rtoe <- as.numeric(sub(".*RTOE = ([0-9]+).*", "\\\1", content))

  if (nchar(rid) == 0) stop("Failed to retrieve RID from BLAST submission")

  return(list(rid = rid, wait = rtoe))
}

# Function to poll BLAST results until ready and parse XML
wait_for_blast_results <- function(rid, max_attempts = 20, wait_sec = 15) {
  for (i in seq_len(max_attempts)) {
    url <- paste0(
      "https://blast.ncbi.nlm.nih.gov/Blast.cgi?CMD=Get&RID=",
      rid, "&FORMAT_TYPE=XML"
    )
    res <- httr::GET(url)
    content <- httr::content(res, as = "text", encoding = "UTF-8")

    if (grepl("<BlastOutput>", content)) {
      cat(sprintf("BLAST results ready on attempt %d.\n", i))
      return(read_xml(content))
    } else {
      cat(sprintf(
        "Attempt %d: Results not ready, retrying in %d seconds...\n",
        i, wait_sec
      ))
    }
  }
}

```

```

        ))
    Sys.sleep(wait_sec)
}
}
stop("Failed to retrieve valid BLAST XML after retries.")
}

# Function to extract unique accession IDs from BLAST XML
extract_accessions <- function(xml_doc, max_hits = 25) {
  hits <- xml_find_all(xml_doc, "//Hit")
  accessions <- unique(xml_text(xml_find_all(hits, "Hit_accession")))
  accessions <- sub("\\..*$", "", accessions) # remove version suffix
  head(accessions, max_hits)
}

# Query protein accession
query_protein <- "XP_001703004"

# Submit BLAST and get RID + wait time
blast_info <- submit_blast(query_protein)
cat("RID:", blast_info$rid, "\nEstimated wait time (seconds):", blast_info$wait, "\n")

Sys.sleep(blast_info$wait + 5) # initial wait

# Poll until BLAST results are ready and parse XML
blast_xml <- wait_for_blast_results(blast_info$rid)

# Extract accession IDs from hits
accessions <- extract_accessions(blast_xml)
cat(sprintf("Found %d accessions.\n", length(accessions)))

# Fetch sequences in FASTA format using Entrez
fasta_seqs <- rentrez::entrez_fetch(
  db = "protein",
  id = paste(accessions, collapse = ","),
  rettype = "fasta",
  retmode = "text"
)

# Save sequences to a FASTA file in data_raw folder
dir.create("data_raw", showWarnings = FALSE)
output_file <- "data_raw/FAP_BLAST.fas"
writeLines(fasta_seqs, con = output_file)
cat(sprintf("Saved %d sequences to %s\n", length(accessions), output_file))

```

### 1.3 Alternative Python Implementation

If retrieval is unsuccessful with the R script, you can increase the `max_attempts` and `wait_sec` parameters and retry. Alternatively, the following Python implementation is generally more robust:

```
# Requirements for the script
# pip install biopython

from Bio.Blast import NCBIWWW, NCBIXML
from Bio import Entrez, SeqIO
import os

Entrez.email = "mobofo3315@feralrex.com"

# Create output directory if needed
os.makedirs("data_raw", exist_ok=True)

# 1. Run BLASTp for XP_001703004
result_handle = NCBIWWW.qblast("blastp", "nr", "XP_001703004", hitlist_size=25)
blast_records = NCBIXML.read(result_handle)

# 2. Collect accession IDs from hits
accessions = []
for alignment in blast_records.alignments:
    # Extract accession (remove version)
    acc = alignment.accession.split('.')[0]
    if acc not in accessions:
        accessions.append(acc)
    if len(accessions) >= 25:
        break

print(f"Found {len(accessions)} accessions.")

# 3. Fetch sequences in FASTA format
handle = Entrez.efetch(
    db="protein",
    id=", ".join(accessions),
    rettype="fasta",
    retmode="text"
)
records = list(SeqIO.parse(handle, "fasta"))

# 4. Save to file
output_file = r"C:\Users\...\data_raw\FAP_BLAST.fas"
SeqIO.write(records, output_file, "fasta")
print(f"Saved {len(records)} sequences to {output_file}.")
```

## 2 Data Preparation

### 2.1 Sequence Consolidation and Duplicate Removal

After acquiring the FAP sequences, they were combined with the existing `FAP_genbank.fas` dataset. All duplicate sequences were removed, and the refined dataset was saved as `FAP_all.fas`. The following code was executed to determine the number of duplicate sequences:

```
duplicates <- duplicated(FAP_all)
sum(duplicates)

# Output: [1] 4
```

**Result:** Four duplicate sequences were identified and removed from the combined dataset.

## 3 Phylogenetic Analysis

### 3.1 Model Selection and Testing

Following model testing, the results were evaluated to select the best-fitting substitution model:

```
head(mt, 5)

#           Model df    logLik      AIC
# 1          WAG 63 -18678.95 37483.90
# 2        WAG+I 64 -18581.27 37290.54
# 3     WAG+G(4) 64 -18333.38 36794.76
# 4 WAG+G(4)+I 65 -18320.26 36770.52
# 5          JTT 63 -18899.60 37925.19
#
#           AICw      AICc      AICcw
# 1 1.231348e-155 37499.23 2.074524e-155
# 2 1.197702e-113 37306.39 1.558333e-113
# 3 5.452270e-06 36810.61 7.093960e-06
# 4 9.999945e-01 36786.89 9.999929e-01
# 5 1.845146e-251 37940.52 3.108624e-251
#
#           BIC
# 1 37759.85
# 2 37570.87
# 3 37075.09
# 4 37055.23
# 5 38201.14
```

#### 3.1.1 Best Fit Model

The model selection analysis revealed **WAG+G(4)+I** as the optimal substitution model with the highest AIC weight (0.9999945). The fitted model parameters are presented below:

```

fit

# model: WAG+G(4)+I
# loglikelihood: -18302.52
# unconstrained loglikelihood: -3743.818
# Proportion of invariant sites: 0.02643574
# Model of rate heterogeneity: Discrete gamma model
# Number of rate categories: 4
# Shape parameter: 2.081987
#      Rate Proportion
# 1 0.0000000 0.02643574
# 2 0.3110929 0.24339107
# 3 0.6814733 0.24339107
# 4 1.1010182 0.24339107
# 5 2.0150298 0.24339107
# Rate matrix: WAG

```

## 3.2 Maximum Likelihood Tree Construction

A maximum likelihood phylogenetic tree was constructed and midpoint-rooted for better visualization. Bootstrap values greater than 70% were displayed to indicate branch support:

```

tree_rooted <- midpoint(fit$tree)

plotBS(tree_rooted,
       trees = fit$bs,
       p = 70,
       type = "phylogram",
       cex = 0.6,
       main = "Maximum Likelihood Tree of FAP Sequences")

add.scale.bar()

```

# Maximum Likelihood Tree of FAP Sequences

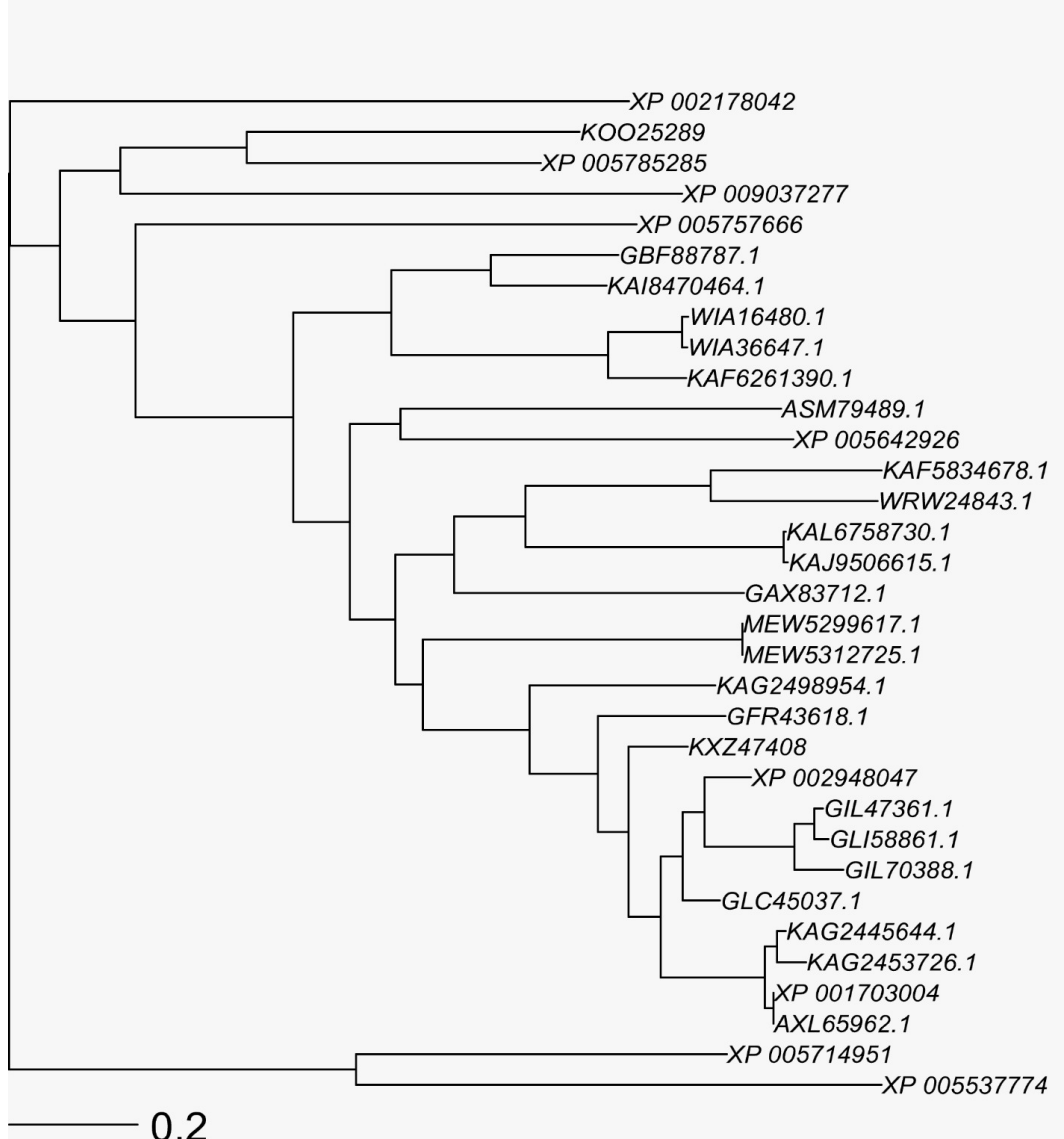


Figure 1: Maximum Likelihood phylogenetic tree of FAP sequences estimated under the WAG+G(4)+I model. Bootstrap values (>70%) are displayed at internal nodes.

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