







```
sr/bin/env python3
from utils import time_me, print_args, print_retval
from utils import get_gtf_value
from utils.rma import RMMTranslationTable
import logging
logger = logging.getlogger() # root logger
logging.besicConfig(level=logging.DMPO,format='%(message)s')
otine ne
@print_args
def get_all_transcripts(filename="Nono_supiens.GRCh38.87.gtf", chromosome='7', gene='895G80000001635');
     transcripts = ()
     # First pass: Fetch all transcripts for the given gene and chromosome
     lagger.debug('First pass on file %s' % filename)
lagger.debug('Chr %s | Gene %s' % (chromosome.gen
with open(filename, modew'rt") as gtf:
              se re = re.compile(r'gene id\s+"?()"?',format(gene))
           for line in gtf:
               blocks = line.split("\t") # Only that chromosome and
               if (
                    continue # skip to the next line
               # Otherwise, it is a transcript for the given gene and chrom
attributes = blocks[8]
transcript_id = get_gtf_value("transcript_id",attributes)
               assert (transcript_id ) # is not None assert (transcript_id not in transcripts), ("How come I see transcript %s already? \n\nLine:\n\n%s" % (transcript_id,line))
               start = int(blocks[3])
end = int(blocks[4])
               strand = 1 if blocks[6] == '+' else -1
               # Adding it to the table
transcripts[transcript_id] = {
                     'start'istart,
'end':end,
                     'strand':strand,
'exons':①, # exons will be added in the second pass. Empty so far.
'start_codon': None,
'stop_codon': None
                logger.debug('Added record: () --- ()'.format(transcript_id,transcripts[transcript_id]))
     logger.debug('Transcripts after first pass')
logger.debug(transcripts)
     # Second pass, fetching the exons for those transcripts
# Must rescan, can't reuse the gtf iterator: it's at the end already.
    logger.debug('Second pass')
with open(filename, mode="rt") as gtf:
    for line in gtf:
        blocks = line.split("\t")
                    len(blocks) < 9 or
                    continue # Skip that line
               feature = blocks[2]
attributes = blocks[8]
               transcript_id = get_gtf_value('transcript_id',attributes)
               if transcript_id not in transcripts: # checking the k
continue # Skip cuz not a transcript for that giv
               if not gene_re.search(attributes):
    print("Weird! I should have a gene_id (gene) in {attr}".format(gene-gene_attr-attributes))
```



```
*scratch* — (179 x 99)
#!/usr/bin/env python3
# -*- coding: utf-8 -*-
 from utils import time_me, print_args, print_retval
 from utils import get_gtf_value
 from utils.rna import RNATranslationTable
 import re
 #from os.path import splitext
 import logging
 logger = logging.getLogger() # root logger
 logging.basicConfig(level=logging.INFO,format='%(message)s')
@time_me
 @print_args
def get_all_transcripts(filename="Homo_sapiens.GRCh38.87.gtf", chromosome='7', gene='ENSG00000001626'):
    transcripts = {}
    # First pass: Fetch all transcripts for the given gene and chromosome
     logger.debug('First pass on file %s' % filename)
    logger.debug('Chr %s | Gene %s' % (chromosome,gene))
    with open(filename, mode="rt") as gtf:
    #gene_id = 'gene_id "%s"' % gene
        gene_re = re.compile(r'gene_id\s+"?{}"?'.format(gene))
         for line in gtf:
            blocks = line.split("\t")
            # Only that chromosome and
             if (
                                                # no comments, please
# only that chromosome. Careful: not comparing integers!
                 len(blocks) < 9 or
                 blocks[0] != chromosome or
                 blocks[2] != 'transcript' or # the line should be a transcript
                 not gene_re.search(blocks[8]) # Is that the right gene?
            ):
                 continue # skip to the next line
            # Otherwise, it is a transcript for the given gene and chromosome
             attributes = blocks[8]
             transcript_id = get_gtf_value('transcript_id',attributes)
             assert( transcript_id ) # is not None
             assert (transcript_id not in transcripts), ("How come I see transcript %s already? \n\nLine:\n\n%s" % (transcript_id,line))
             start = int(blocks[3])
             end = int(blocks[4])
             strand = 1 if blocks[6] == '+' else -1
            # Adding it to the table
             transcripts[transcript_id] = {
                 'start':start,
                 'end':end,
                 'strand':strand,
                 'exons':{}, # exons will be added in the second pass. Empty so far.
                  'start_codon': None,
                 'stop_codon': None
             logger.debug('Added record: {} => {}'.format(transcript_id,transcripts[transcript_id]))
     logger.debug('Transcripts after first pass')
     logger.debug(transcripts)
    # Second pass, fetching the exons for those transcripts
    # Must rescan, can't reuse the gtf iterator: it's at the end already.
     logger.debug('Second pass')
    with open(filename, mode="rt") as gtf:
         for line in gtf:
            blocks = line.split("\t")
             if (
                 len(blocks) < 9 or
                                             # no comments, please
                 blocks[0] != chromosome or # only that chromosome
                 not (blocks[2] == "exon" or blocks[2] == "start_codon" or blocks[2] == "stop_codon")
             ):
                 continue # Skip that line
             feature = blocks[2]
             attributes = blocks[8]
             transcript_id = get_gtf_value('transcript_id',attributes)
            if transcript_id not in transcripts: # checking the keys
                 continue # Skip cuz not a transcript for that given gene
             if not gene_re.search(attributes):
                 print("Weird! I should have a gene_id {gene} in {attr}".format(gene=gene,attr=attributes))
             if feature == "exon":
                 logger.debug('Found an exon')
                 exon_id = get_gtf_value('exon_id',attributes)
                 exons = transcripts[transcript_id].get('exons',None)
                 assert( exons is not None )
                 if exon_id in exons:
                     print("Weird! Have I seen that exon %s before?" % exon_id)
U:--- main.py
                       Top L26 Git-vt17 (Python MMM) 02:37 1.21
```