FastQCParser User Guide:	
is recommended to open this via browser: <a href="https://ms2206.github.io/FastQCParser/">https://ms2206.github.io/FastQCParser/</a>	
ntroduction	
Python program to parse FastQCtext files, and generate reports and plots.	
1. Clone directory: git clone https://github.com/ms2206/FastQCParser.git	

- 2. Make a new python environment based from requirments.yaml conda env create -f requirements.yaml --<NAME>
- 3. Load environment env. conda activate <NAME>
- 4. Change directory into FastQCParser
- 5. Example Usage: python3 src/main.py data/raw/fastqc\_data2.txt fastqc\_2 -a

## Set up

Example Usage: With python3, run executable found at src/main.py, pass in data/raw/fastqc\_data2.txt - as input file, and fastqc\_2 as output directory. Use optional argument -a.

python3 src/main.py data/raw/fastqc\_data2.txt fastqc\_2 -a

#### **Entry Point:**

python3 src/main.py data/raw/fastqc\_data2.txt fastqc\_2 -a

#### Args Help

#### python src/main.py -h

usage: main.py [-h] [-b] [-t] [-s] [-c] [-g] [-n] [-l] [-d] [-o] [-p] [-k] [-a] input\_file output\_dir

Parse and plot FASTQC data.

positional arguments:

input\_file Path to the FASTQC file. output dir Directory to save plots.

optional arguments:

-h, --help show this help message and exit

-b, --per\_base\_seq\_qual

Extract and plot per base sequence quality.

-t, --per\_tile\_seq\_qual

Extract and plot per tile sequence quality.

-s, --per\_seq\_qual\_scores

Extract and plot per sequence quality scores

-c, --per\_base\_seq\_content

Extract and plot per base sequence content

-g, --per\_seq\_GC\_cont

Extract and plot per sequence GC content

-n, --per\_base\_N\_cont

Extract and plot per base N content

-I, --seq\_len\_dist Extract sequence length distribution

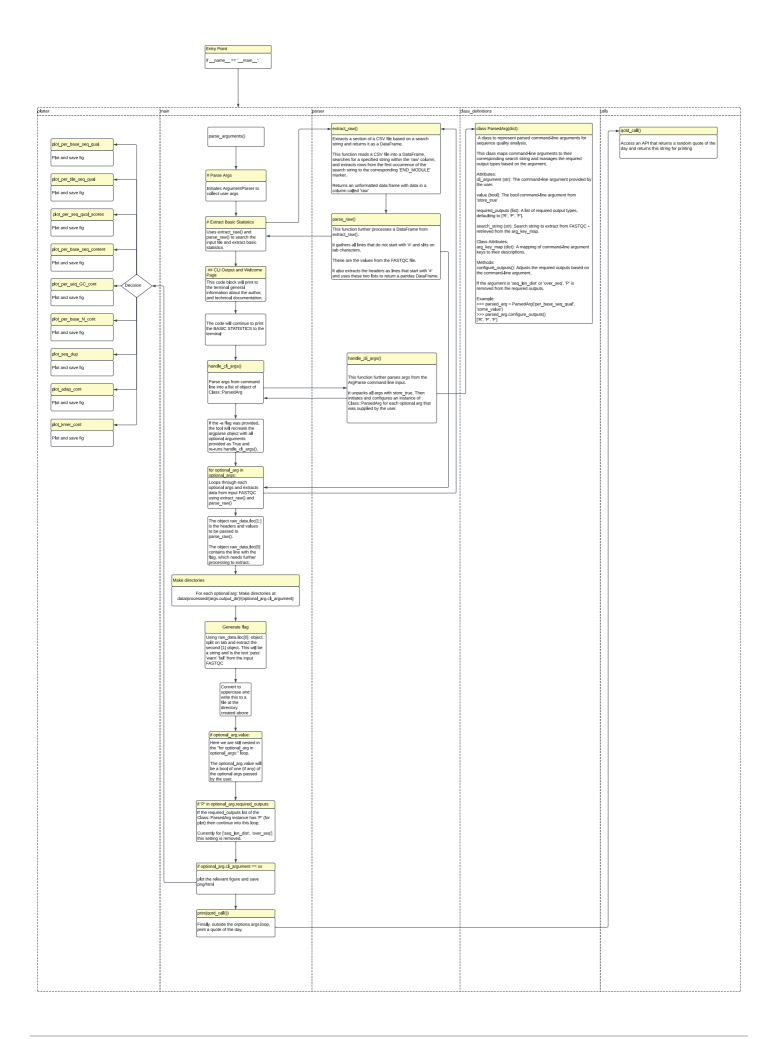
-d, --seq\_dup Extract and plot sequence duplication levels

-o, --over\_seq Extract overrepresented sequences
-p, --adap\_cont Extract and plot adapter content
-k, --kmer\_cont Extract and plot K-mer Content

-a, --all Extract and plot all metrics

#### **Optional Args**

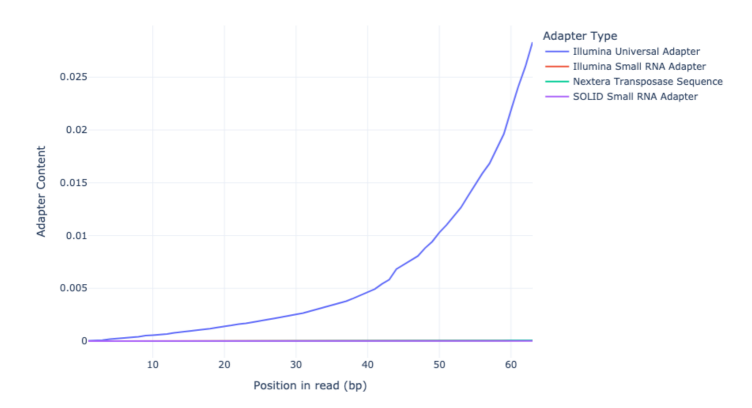
Help and misc information provided by ArgeParse for optional arguments.



#### Adapter Content

Plot's adapter content by position.

## Adapter content by position



#### Kmer Content

Plot's kmer content by position.

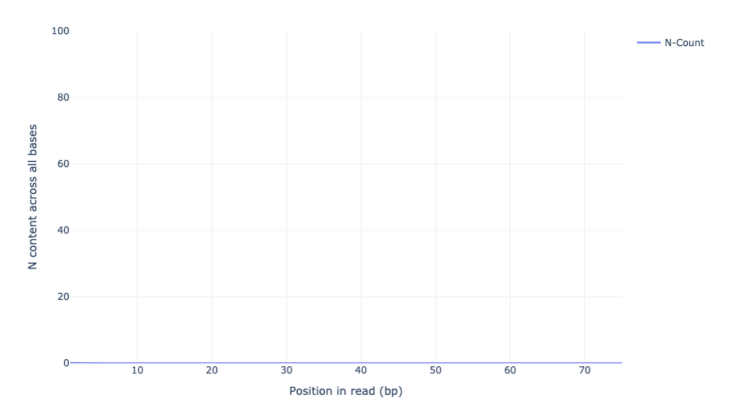
#### Sequence Counts



### Overrepresented sequences

Plot's Per base N content.

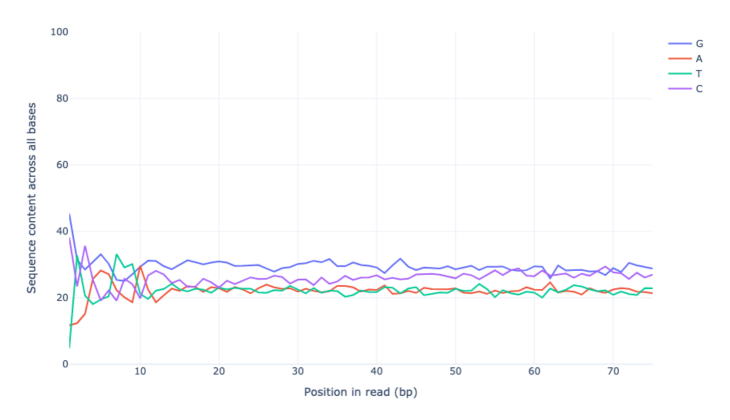
# Adapter content by position



Per base sequence content

Plot's Per base sequence content.

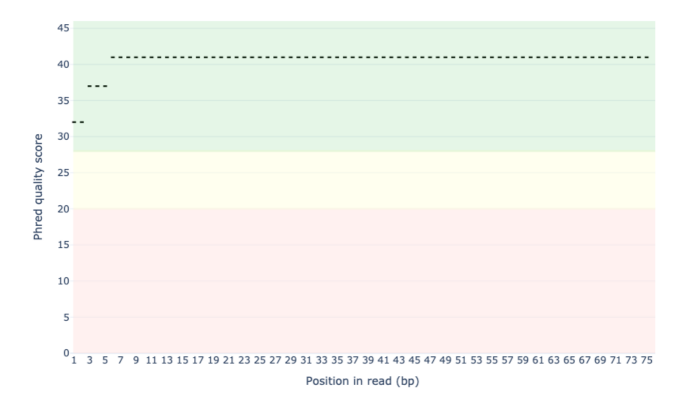
## Per base sequence content



Per sequence quality scores

Plot's Per sequence quality scores.

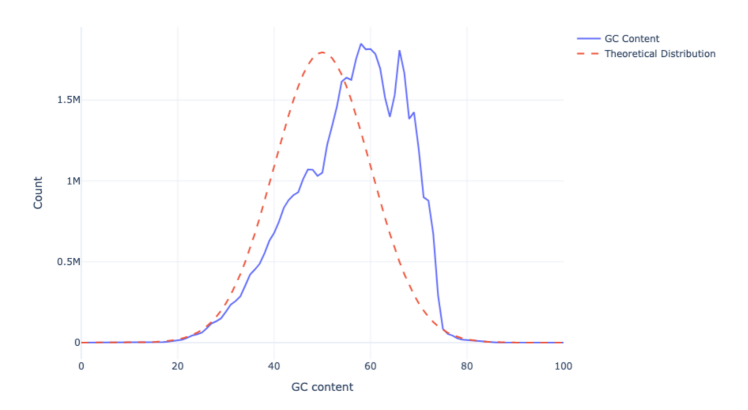
## Per base sequence quality



Per sequence GC content

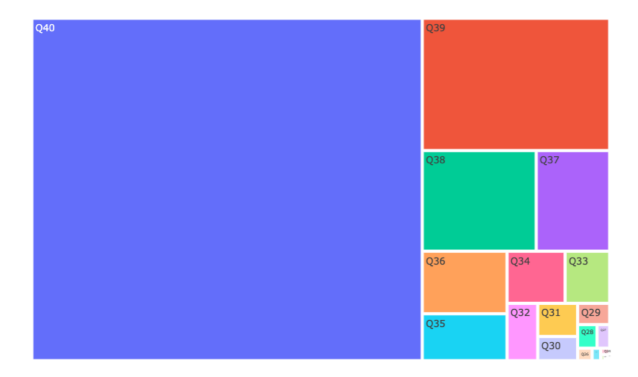
Plot's Per sequence GC content.

# Per sequence GC content



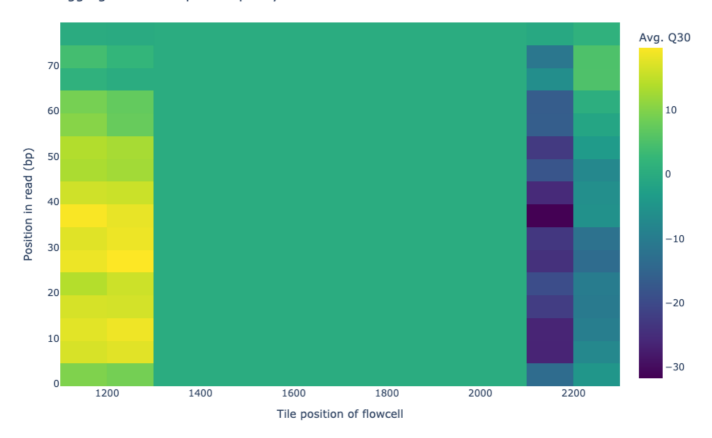
Per sequence quality scores Plot's Per sequence quality scores.

#### Count of Phred score



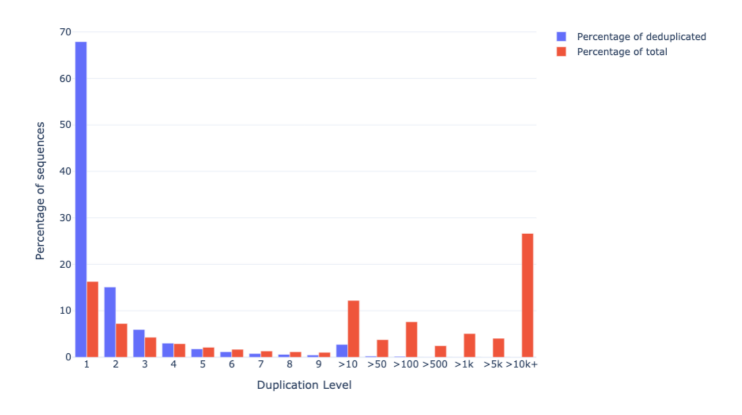
Per tile sequence quality
Plot's Per tile sequence quality.

# Per aggregated tile sequence quality



Sequence Duplication Levels Plot Sequence Duplication Levels.

## Sequence Duplication Levels



# GitHub https://github.com/ms2206/FastQCParser.git

Documentation built with MkDocs.