Next generation sequencing (NGS) database for tandem repeats with multiple pattern 2°-shaft multicore string matching

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A B S T R A C T

Next generation sequencing (NGS) technologies have been rapidly applied in biomedical and biological research in recent years. To provide the comprehensive NGS resource for the research, in this paper, we have considered 10 loci/repeats TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA. Then we developed the NGS Tandem Repeat Database (TandemRepeatDB) for all the chromosomes of Homo sapiens, Callithrix jacchus, Chlorocebus sabaeus, Gorilla gorilla, Macaca fascicularis, Macaca mulatta, Nomascus leucogenys, Pan troglodytes, Papio anubis and Pongo abelii genome data sets for all those loci. We find the successive occurrence frequency for all the above 10 SSR (simple sequence repeats) in the above genome data sets on a chromosome-by-chromosome basis with multiple pattern 2° shaft multicore string matching.

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1. Introduction

Since the completion of the first human genome sequence, demand for cheaper and faster sequencing methods has increased greatly. This demand has driven the development of second-generation sequencing methods, or next-generation sequencing (NGS). In this paper we developed NGS Tandem Repeat Database (TandemRepeatDB) that stores the successive occurrence frequency of SSRs from the considered genomes.

Simple sequence repeats (SSRs) are tandemly repeated DNA sequences found in varying abundance in most genomes [1,2]. These repeats have been extensively used for genetic mapping and population studies [3]. SSRs also provide molecular tools to understand spatial relationships between chromosome segments, which in turn, aid in analyzing temporal relationships between species and genera [4]. In humans about 3% of the genome is occupied by SSRs [5].

The study of repeat frequency and its distribution pattern in the genome is expected to help in understanding their significance. There is accumulating evidence to suggest that SSRs function to regulate gene expression [6,7].

The availability of complete genome sequences for many organisms has made it possible to carry out genome-wide analyses. In the present study we have screened all the chromosomes of Homo sapiens, Callithrix jacchus, Chlorocebus sabaeus, Gorilla gorilla, Macaca fascicularis, Macaca mulatta, Nomascus leucogenys, Pan troglodytes, Papio anubis and Pongo abelii [8] and studied the distribution and successive occurrence frequency of TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA loci [9,10].

Earlier, few studies [11–14] have attempted to analyze the distribution of tandem repeats in human like genomes but they are confined to a single or a small set of genomes. This tandem repeat mining helps in understanding and addressing biological questions. It is used in various diverse applications like DNA fingerprinting, maternity identification, paternity identification, theft identification, suspect findings, and disease identification [15,16,9,10].

2. Methodology

In the present paper, we constructed the TandemRepeatDB with multiple pattern 2° shaft multicore string matching algorithm. String matching [17–20] is a process of identifying the pattern (P) in a given text (T). In the present paper chromosomes of genomes are considered as text (T) and the loci are considered as patterns. The multiple pattern 2° shaft multi core string matching algorithm searches the multiple patterns concurrently in a single part (2° = 1 shaft) with multi core processors.
In the TandemRepeatDB construction, the text file and the patterns are read and the patterns are searched in text file (T) with multiple pattern 2° shaft multi core string matching algorithm. If perfect tandem repeat occurs then the successive logic is applied. The successive logic means continuous perfect occurrence of similar tandem repeats. If the successive tandem repeat size > 1 then the successive occurrence of tandem repeat information is stored in the database. The database is constructed in MySQL using JAVA. The TandemRepeatDB construction process comprises four stages and its complete architecture is shown in Fig. 1.

STAGE I: Reading
(a) The Text file is read.
(b) The Pattern set is read.

STAGE II: Searching
(a) All the patterns from the given set are read and categorized basing on their right most characters.
(b) One of the patterns from one category is selected.
(c) The shift_left_to_right (Pm-1) function is applied for shift position.
(d) The multiple pattern 2° shaft multicore string matching algorithm is selected for searching.

STAGE III: Search results
(a) If a perfect repeat occurs then successive occurrences are searched.

STAGE IV: Storing
(a) If the perfect successive repeat size > 1 then it is stored in the TandemRepeatDB with the following information.
   - sample_name
   - sample_chromosome_name
   - position
   - noofoccurences
   - codi/repeat name

The multiple pattern 2° shaft multicore string matching algorithm consists input and output, initialization, main function, search function and shift_left_to_right function.

In the input and output the genome sets and patterns are taken as input and the sample_id, sample_name, sample_chromosome_name, position, noofoccurrences, and codi are returned as output. In the initialization, multi_pattern (all pattern in the set), n (text length), m (pattern length) and all other variables are initialized. In the main function the genome set is read on chromosome by chromosome basis, the individual chromosome is given to shift_left_to_right function. Once the shift value is received the search function is called for all the patterns. The shift_left_to_right function, applies the shift operation with the pattern’s rightmost character and the shift position is returned to main function. The search process compares character by character from both the directions until a complete match or mismatch occurs.

Fig. 1. Architecture of TandemRepeatDB.
If match occurs the successive occurrence of the pattern is searched. If the successive occurrence size is greater than 1 then the data is stored into TandemRepeatDB. If the mismatch or complete match occurs the same procedure is repeated until the end of the T.

2.1. Structure of the database

In this paper a table is created with the given sample name. The table contains sample_id, sample_name, sample_chromosome_name, position indicating the occurrence position of the codi, noocurrences and codi.

| Table 1 | Table Structure. |
|---------|------------------|
| Type    | Collation        |
| sample_id | text              |
| sample_name | text              |
| sample_chromosome_name | text          |
| position | int(10)           |
| noocurrences | int(10)         |
| codi    | text              |

2.1.1. Structure of the database

In this paper a table is created with the given sample name. The table contains sample_id, sample_name, sample_chromosome_name, position indicating the occurrence position of the codi, noocurrences and codi.

| Table 2 | Genome sequences used in the study. |
|---------|-----------------------------------|
| Genome sequence name | Name & number of chromosomes | Total number of tandem repeats extracted (>1) |
| Homo sapiens | 1 to 22, MT, X, Y and Un (26) | 11,99,985 |
| Callithrix jacchus | 1 to 22, X, Y and Un (25) | 11,40,529 |
| Chlorocebus sabaeus | 1 to 29, MT, X and Un (33) | 11,13,445 |
| Gorilla gorilla | 1, 2A, 3 to 22, MT, X and Un (26) | 11,83,843 |
| Macaca fascicularis | 1 to 20, MT, X and Un (23) | Once (12,31,029) |
| Macaca mulatta | 1 to 20, MT, X and Un (23) | 12,74,556 |
| Nomascus | 1a to 2, 7b to 8, 22a to 25, X and Un (27) | 11,71,594 |
| Pan troglodytes | 1, 2A, 3 to 22, MT, X and Un (27) | 12,76,766 |
| Papio anubis | 1 to 20, MT, X and Un (23) | 13,51,393 |
| Pongo abelii | 1, 2A, 3 to 22, MT, X and Un (26) | 13,81,887 |

3. Tandem repeat size analysis

In this section, the perfect successive tandem repeats are extracted and analyzed by executing SQL queries on the TandemRepeatDB for all the chromosomes of the considered genomes corresponding to TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, TCTG, TCTG and TCTG repeats have been presented.

In this study 259 chromosomes of H.sapiens, C.jacchus, C.sabaeus, G.gorilla, M.fascicularis, M.mulatta, N.leucogenys, P.troglodytes, P.anubis and P.abelii genomes have been used as shown in Table 2.

| Table 3 | Tandem repeat successive occurrences for all chromosomes of H.sapiens. |
|---------|-----------------------------------------------|
| codi/Repeat name | MAX number of codi in the successive occurrences | Number of times the MAX number appeared |
| TAGA | 21 | Once |
| AGAA | 42 | Twice |
| GATA | 22 | Once |
| TATC | 25 | Once |
| TCAT | 12 | Twice |
| GAAT | 12 | Once |
| AGAT | 21 | Once |
| CTTG | 78 | Once |
| TATC | 25 | Once |
| TCTG | 12 | Four Times |

indicating the number of occurrences of the repeat and codi indicating the name of the repeat. The table structure is shown in Table 1.

Availability of NGS techniques leads to the accessibility of genome sequences. Studying the perfect successive occurrences of the tandem repeats using Bioinformatics approach would be very interesting and informative.

In the remaining part of the study, perfect tandem repeats of all chromosomes in H. sapiens, C. jacchus, C. sabaeus, G. gorilla, M. fascicularis, M. mulatta, N. leucogenys, P. troglodytes, P. anubis and P. abelii genomes are analyzed and a brief note on the successive occurrence frequency of TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats have been presented.

In this study 259 chromosomes of H.sapiens, C.jacchus, C.sabaeus, G.gorilla, M.fascicularis, M.mulatta, N.leucogenys, P.troglodytes, P.anubis and P.abelii genomes have been used as shown in Table 2.

3.1. H. sapiens

H. sapiens are the binomial nomenclature for the human species. Homo is the human genus, which also includes Neanderthals and many other extinct species of hominid.
In the paper, multiple pattern 2° shaft multicore string matching algorithm is used to retrieve the perfect successive tandem repeats from H. sapiens genomes which consists 1 to 22, MT, X, Y and Un chromosomes as shown in Table 2. A total of 11,99,985 perfect successive repeats are extracted from the above chromosomes, which are stored in the homo_sapiens table.

Table 3 gives the summary of extracted MAX number of successive occurrences from the homo_sapiens table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats. The TAGA results are extracted from the table by executing Query1 and Query2.

\[
\text{Query}_1 = \pi_{\text{max number of occurrences}}(\sigma_{\text{taga}} \text{ homo_sapiens}) \\
\text{Query}_2 = \pi_{\text{max number of occurrences}}(\sigma_{\text{taga}} \Lambda \text{ no occurrences} = \text{Query}_1 \text{ homo_sapiens})
\]

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats. The extracted MAX number of successive occurrences from the homo_sapiens table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 2.

From the Fig. 2, the following observations can be made:

- CTTT tandem repeat has maximum of 78 successive base pairs,
- AGAA tandem repeat has maximum of 42 successive base pairs twice,
- TCTG tandem repeat has maximum of 12 successive base pairs four times,
- The remaining Tandem repeats have successive base pairs from a minimum of 12 to a maximum of 25,
- All the above observations have a significant role in the bio-informatics studies.

3.2. C. jacchus

The common marmoset is a New World monkey. It originally lived in the Northeastern coast of Brazil.

In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from C. jacchus genomes which consists 1 to 22, X, Y and Un chromosomes as shown in Table 2. A total of 11,40,529 perfect successive repeats are extracted from the above chromosomes, which are stored in the callithrix_jacchus table.

Table 4 gives the summary of extracted MAX number of successive occurrences from the callithrix_jacchus table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats.

The TAGA results are extracted from the table by executing Query1 and Query2.

\[
\text{Query}_1 = \pi_{\text{max number of occurrences}}(\sigma_{\text{taga}} \text{ callithrix_jacchus}) \\
\text{Query}_2 = \pi_{\text{max number of occurrences}}(\sigma_{\text{taga}} \Lambda \text{ no occurrences} = \text{Query}_1 \text{ callithrix_jacchus})
\]

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats.

The extracted MAX number of successive occurrences from the callithrix_jacchus table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 3.

From the Fig. 3, the following observations can be made:

- AGAA tandem repeat has maximum of 57 successive base pairs,
- CTTT tandem repeat has maximum of 51 successive base pairs,
- TATC tandem repeat has maximum of 18 successive base pairs thrice,
- The remaining Tandem repeats have successive base pairs from a minimum of 13 to a maximum of 21,
- All the above observations have a significant role in the bio-informatics studies.

3.3. C. sabaeus

The green monkey, also known as the sabaeus monkey or the Callithrix monkey, is an Old World monkey with golden-green fur and pale hands and feet. The tip of the tail is golden yellow as are the backs of the thighs and cheek whiskers.

In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from C. sabaeus genomes.
which consists 1 to 29, MT, X, Y and Un chromosomes as shown in Table 2. A total of 11,13,445 perfect successive repeats are extracted from the above chromosomes, which are stored in the chlorocebus_sabaeus table.

Table 5 gives the summary of extracted MAX number of successive occurrences from the chlorocebus_sabaeus table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats. The TAGA results are extracted from the table by executing Query1 and Query2.

\[
\text{Query}_1 = \pi_{\text{max \ noofoccurrences}} (\sigma_{\text{codi = \text{TAGA} (chlorocebus_sabaeus) }} )
\]

\[
\text{Query}_2 = \pi_{\text{max \ noofoccurrences}} (\sigma_{\text{codi = \text{TAGA} \land \text{noofoccurrences = Query1 (chlorocebus_sabaeus) }}})
\]

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA.

The extracted MAX number of successive occurrences from the gorilla_gorilla table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 5.

From the Fig. 5, the following observations can be made:

- CTTT tandem repeat has maximum of 66 successive base pairs,
- AGAA tandem repeat has maximum of 41 successive base pairs,
- GAAT tandem repeat has maximum of 12 successive base pairs ten times,
- The remaining tandem repeats have successive base pairs from a minimum of 14 to a maximum of 26,
- All the above observations have a significant role in the bio-informatics studies.

3.5. M. fascicularis

The crab-eating macaque, also known as the long-tailed macaque, is a cercopithecine primate native to Southeast Asia. It is referred to as the cynomolgus monkey in laboratories.

In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from M. fascicularis genomes which consists 1 to 20, MT, X and Un chromosomes as shown in Table 2. A total of 12,31,029 perfect successive repeats are extracted from the above chromosomes, which are stored in the macaca_fascicular table.

Table 6 gives the summary of the extracted MAX number of successive occurrences from the gorilla_gorilla table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats. The TAGA results are extracted from the table by executing Query1 and Query2.

\[
\text{Query}_1 = \pi_{\text{max \ noofoccurrences}} (\sigma_{\text{codi = \text{TAGA} (gorilla_gorilla) }} )
\]

\[
\text{Query}_2 = \pi_{\text{max \ noofoccurrences}} (\sigma_{\text{codi = \text{TAGA} \land \text{noofoccurrences = Query1 (gorilla_gorilla) }}})
\]

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA.

The extracted MAX number of successive occurrences from the gorilla_gorilla table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 5.

From the Fig. 5, the following observations can be made:

- CTTT tandem repeat has maximum of 66 successive base pairs,
- AGAA tandem repeat has maximum of 41 successive base pairs,
- GAAT tandem repeat has maximum of 12 successive base pairs ten times,
- The remaining tandem repeats have successive base pairs from a minimum of 14 to a maximum of 26,
- All the above observations have a significant role in the bio-informatics studies.

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In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from M. fascicularis genomes which consists 1 to 20, MT, X and Un chromosomes as shown in Table 2. A total of 12,31,029 perfect successive repeats are extracted from the above chromosomes, which are stored in the macaca_fascicular table.

Table 5

| codi/Repeat name | MAX number of codi in the successive occurrences | Number of times the MAX number appeared |
|------------------|-----------------------------------------------|----------------------------------------|
| TAGA             | 19                                            | Twice                                  |
| AGAA             | 54                                            | Once                                   |
| GATA             | 20                                            | Once                                   |
| TCTA             | 20                                            | Twice                                  |
| GAAT             | 14                                            | Once                                   |
| TATC             | 14                                            | Once                                   |
| CTTT             | 42                                            | Once                                   |
| TCTG             | 14                                            | Once                                   |

Fig. 3. Max number of successive occurrences of all repeats for all chromosomes of C. jacchus.
Table 7 gives the summary of the extracted MAX number of successive occurrences from the *macaca_fascicularis* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats. The TAGA results are extracted from the table by executing Query1 and Query2.

Query1 = π_{max(no of occurrences)}(σ_{codi−TAGA} (macaca_fascicularis))

Query2 = π_{max(no of occurrences)}(σ_{codi−TAGA and no of occurrences−query1} (macaca_fascicularis))

Query1, and Query2, are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA.

The extracted MAX number of successive occurrences from the *macaca_fascicularis* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 6.

From the Fig. 6, the following observations can be made:
- CTTT tandem repeat has maximum of 221 successive base pairs,
- AGAA tandem repeat has maximum of 218 successive base pairs,
- GAAT tandem repeat has maximum of 14 successive base pairs thrice,
- The remaining tandem repeats have successive base pairs from a minimum of 16 to a maximum of 33,
- All the above observations have a significant role in the bio-informatics studies.

3.6. *M. mulatta*

The rhesus macaque (*M. mulatta*), is one of the best-known species of Old World monkeys. It is listed as the least concern in the IUCN Red List of Threatened Species in view of its wide distribution, presumed large population, and its tolerance of a broad range of habitats.

In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from *M. mulatta* genomes which consists 1 to 20, MT, X and Un chromosomes as shown in Table 2. A total of 12,74,556 perfect successive repeats are extracted from the above chromosomes, which are stored in the *macaca_mulatta* table.

Table 8 gives the summary of the extracted MAX number of successive occurrences from the *macaca_mulatta* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats.

The TAGA results are extracted from the table by executing Query1 and Query2.

Query1 = π_{max(no of occurrences)}(σ_{codi−TAGA} (macaca_mulatta))

Query2 = π_{max(no of occurrences)}(σ_{codi−TAGA and no of occurrences−query1} (macaca_mulatta))

Query1, and Query2, are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA.

The extracted MAX number of successive occurrences from the *macaca_mulatta* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 7.

From the Fig. 7, the following observations can be made:
- AGAA tandem repeat has maximum of 84 successive base pairs,
- CTTT tandem repeat has maximum of 79 successive base pairs,
- TCTA tandem repeat has maximum of 21 successive base pairs twice,
- The remaining tandem repeats have successive base pairs from a minimum of 12 to a maximum of 31,
- All the above observations have a significant role in the bio-informatics studies.

3.7. *N. leucogenys*

The northern white-cheeked gibbon is a species of gibbon native to South East Asia. It is closely related to the southern white-cheeked gibbon, with which it was previously considered conspecific.

In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from *N. leucogenys* genomes which consists 1 to 6, 7b, 8 to 21, 22a, 23 to 25, X and Un chromosomes as shown in Table 2. A total of 11,71,594 perfect successive repeats are extracted from the above chromosomes, which are stored in the *nomascus_leucogenys* table.

Table 9 gives the summary of the extracted MAX number of successive occurrences from the *nomascus_leucogenys* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats.
The TAGA results are extracted from the table by executing Query1 and Query2.

Query1 = \( \pi_{\text{max}(\text{occurrences})} \left( \sigma_{\text{taga}} \right) \) (nomascus_leucogenys )

Query2 = \( \pi_{\text{max}(\text{occurrences})} \left( \sigma_{\text{taga}} \times \text{\text{occurrences}} \right) \) (nomascus_leucogenys )

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTIT, TCTG, and TCTA.

The extracted MAX number of successive occurrences from the nomascus_leucogenys table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTIT, TCTG, and TCTA repeats is graphically shown in Fig. 8.

From the Fig. 8, the following observations can be made:

- AGAA tandem repeat has maximum of 52 successive base pairs,
- CTTT tandem repeat has maximum of 33 successive base pairs,
- TCTG tandem repeat has maximum of 17 successive base pairs thrice,
- The remaining tandem repeats have successive base pairs from a minimum of 11 to a maximum of 23,
- All the above observations have a significant role in the bio-informatics studies.

### 3.8. P. troglodytes

The common chimpanzee (P. troglodytes), also known as the robust chimpanzee, is a species of great apes. Colloquially, the common chimpanzee is often called the chimpanzee.

In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from P. troglodytes genomes which consists 1, 2A, 2B, 3 to 22, MT, X, Y and Un chromosomes as shown in Table 2. A total of 12,76,766 perfect successive repeats are extracted from the above chromosomes, which are stored in the pan_troglodytes table.

Table 10 gives the summary of the extracted MAX number of successive occurrences from the pan_troglodytes table for TAGA, TCAT, GAAT,
AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats. The TAGA results are extracted from the table by executing Query1 and Query2:

$$\text{Query}_1 = \pi_{\max(nofoccurences)\left(\sigma_{\text{codi}}\left(\text{TAGA}\right)\right)}$$

$$\text{Query}_2 = \pi_{\max(nofoccurences)\left(\sigma_{\text{codi}}(\text{TAGA} \land \text{nofoccurences}) - \text{Query}_1\right)}$$

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA.

The extracted MAX number of successive occurrences from the pan_troglodytes table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 9.

From the Fig. 9, the following observations can be made:

- AGAA tandem repeat has maximum of 43 successive base pairs,
- CTTT tandem repeat has maximum of 30 successive base pairs,
- TCAT tandem repeat has maximum of 10 successive base pairs five times.
The remaining tandem repeats have successive base pairs from a minimum of 11 to a maximum of 19.

All the above observations have a significant role in the bio-informatics studies.

3.9. *P. anubis*

The olive baboon, also called the Anubis baboon, is a member of the family Cercopithecidae. The species is the most widely ranging of all baboons.

In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from *P. anubis* genomes which consists 1 to 20, MT, X and Un chromosomes as shown in Table 2. A total of 13,51,393 perfect successive repeats are extracted from the above chromosomes, which are stored in the *papio_anubis* table.

Table 11 gives the summary of the extracted MAX number of successive occurrences from the *papio_anubis* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats. The TAGA results are extracted from the table by executing Query1 and Query2.

\[
\text{Query}_1 = \pi_{\text{max number of occurrences}}(\sigma_{\text{TAGA}}(\text{papio_anubis}))
\]

\[
\text{Query}_2 = \pi_{\text{max number of occurrences}}(\sigma_{\text{TAGA}} \Lambda \text{noofoccurrences} \text{Query1}(\text{papio_anubis}))
\]

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA.

The extracted MAX number of successive occurrences from the *papio_anubis* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 10.

From the Fig. 10, the following observations can be made.

- AGAA tandem repeat has maximum of 54 successive base pairs,
- CTTT tandem repeat has maximum of 47 successive base pairs twice,
- TCTG tandem repeat has maximum of 21 successive base pairs twice,
- The remaining tandem repeats have successive base pairs from a minimum of 14 to a maximum of 32,
- All the above observations have a significant role in the bio-informatics studies.

3.10. *P. abelii*

The Sumatran orangutan is one of the two species of orangutans. Found only in the island of Sumatra, in Indonesia, it is rarer than the Bornean orangutan. In the paper, the proposed string matching algorithm is used to retrieve the perfect successive tandem repeats from *P. abelii* genomes which consists 1, 2A, 2B, 3 to 22, MT, X and Un chromosomes as shown in Table 2. A total of 13,81,887 perfect successive repeats are extracted from the above chromosomes, which are stored in the *pongo_abelii* table.

Table 12 gives the summary of extracted MAX number of successive occurrences from the *pongo_abelii* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA repeats. The TAGA results are extracted from the table by executing Query1 and Query2.

\[
\text{Query}_1 = \pi_{\text{max number of occurrences}}(\sigma_{\text{TAGA}}(\text{pongo_abelii}))
\]

\[
\text{Query}_2 = \pi_{\text{max number of occurrences}}(\sigma_{\text{TAGA}} \Lambda \text{noofoccurrences} \text{Query1}(\text{pongo_abelii}))
\]

Query1 and Query2 are executed for remaining repeats TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG and TCTA.

The extracted MAX number of successive occurrences from the Table 10 and Table 11 for all chromosomes of *P. troglodytes* and *P. anubis* are shown in the diagrams.

![Fig. 9. Max number of successive occurrences of all repeats for all chromosomes of *P. troglodytes*.](image-url)
The extracted MAX number of successive occurrences from the *pongo_abelli* table for TAGA, TCAT, GAAT, AGAT, AGAA, GATA, TATC, CTTT, TCTG, and TCTA repeats is graphically shown in Fig. 11.

From the Fig. 11, the following observations can be made.

- CTTT tandem repeat has maximum of 63 successive base pairs,
- AGAA tandem repeat has maximum of 37 successive base pairs,
- TCTA tandem repeat has maximum of 18 successive base pairs twice,
- The remaining tandem repeats have successive base pairs from a minimum of 11 to a maximum of 20,
- All the above observations have a significant role in the bio-informatics studies.

### Table 12

| codi/Repeat name | MAX number of codi in the successive occurrences | Number of times the MAX number appeared |
|------------------|-------------------------------------------------|----------------------------------------|
| TAGA             | 20                                              | Once                                   |
| AGAA             | 37                                              | Once                                   |
| GATA             | 19                                              | Once                                   |
| TCTA             | 18                                              | Twice                                  |
| TCAT             | 12                                              | Once                                   |
| GAAT             | 13                                              | Once                                   |
| AGAT             | 20                                              | Once                                   |
| CTTT             | 63                                              | Once                                   |
| TATC             | 19                                              | Twice                                  |
| TCTG             | 11                                              | Twice                                  |

### 4. Conclusions

In this paper we developed the TandemRepeatDB that provides a single portal access to perfect successive repeats in genomes of *H. sapiens*, *C. jacchus*, *C. sabaeus*, *G. gorilla*, *M. fascicularis*, *M. mulatta*, *N. leucogenys*, *P. troglodytes*, *P. anubis* and *P. abelli*. The database is known to be the first of its kind to host all types of perfect successive tandem repeats for the considered genomes. From the analysis of all the records existing in the TandemRepeatDB, it is observed that CTTT tandem repeat and AGAA tandem repeat occupy the major role. This TandemRepeatDB will be a very valuable resource for researchers studying repeats in the above mentioned genomes.

### Conflict of interests

Authors did not have any conflict of interests.

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