RESEARCH ARTICLE



Development and Characterization SSR Makers Based on Next-generation Sequencing Technology in Iranian Turkmen Camel

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ARTICLE INFO

Article history: Received 17 May 2025

Accepted 06 July 2025 Available 24 July 2025

Keywords:

Primer SSR marker Turkmen camel Whole genome sequencing

Supplementary information:

Supplementary information for this article is available at http://sc.journals.umz.ac.ir/

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p-ISSN 2423-4257 e-ISSN 2588-2589

ABSTRACT

The Iranian Turkmen one-humped camel (Camelus dromedarius) is an economically and culturally significant species valued for its milk and meat production. It is well-adapted to the arid and semi-arid climates of Middle East. Despite its importance, the genetic diversity and historical demography of this species remain poorly characterized compare to two-humped camel species, which exhibit limited genetic differentiation, the genetic diversity and historical demography of one-humped camels remain poorly understood. This study developed novel simple sequence repeat (SSR) markers for the Iranian Turkmen camel genome to facilitate genetic studies and marker-assisted breeding. We aimed to identify polymorphic loci to support analyses of genetic diversity, population structure, and evolutionary biogeography. Using Illumina HiSeq 2000 sequencing, we generated 589,326,158 clean 150 bp paired-end reads at 50x coverage. De novo assembly produced 235,978 contigs (N50= 8,526 bp) from which 151,556 SSR motifs were identified, primarily 2-mer (38.80%), 4mer (21.85%), and 1-mer (16.20%) motifs. Primer pairs were designed for 144,184 loci (95%) (amplicon sizes of 100-180 bp). The Turkmen camel exhibited high SSR diversity (6,201 unique motifs), significantly exceeding than of Arabian and Bactrian camels ($\chi^2 = 7.14$, p= 0.007, df= 1) with an estimated 78 to 510 additional unique motifs. These markers enable genetic diversity analysis, historical demography studies, and identification of breed-specific motifs (e.g., 1,179 novel motifs in Turkmen camels). Additionally, 19,425 codon-repeat loci, predominantly leucine-rich repeats (42%), were identified, potentially linked to stress response genes, offering insights into functional genomics. These SSR markers support conservation genomics and breeding programs for Camelus species, enhancing their long-term sustainability. Limitations include potential bias from using short contigs rather than scaffolds, and the need for functional validation of 19,425 codon-repeat loci, particularly leucine-rich repeats (42%).

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Please cite this paper as: Rezyannejad, E., Nobari, K., & Bakhtiari Zadeh, M. R. (2025). Development and characterization SSR makers based on next-generation sequencing technology in Iranian Turkmen camel. Journal of Genetic Resources, 11(2), 191-202. doi: 10.22080/jgr.2025.28696.1424

Introduction

The Camelidae family, originating in North America during the Tortonian stage of the Miocene, later diverged into two tribs: Lamini

and Camelini. Camelini, or the old-world camels, belong to the order Artiodactyla and migrated to the Eastern Hemisphere via the Bering land bridge (Wu et al., 2014; Wu et al., 2015). The genus Camelus comprises three extant species:

the one-humped dromedary (*Camelus dromedarius*), also known as Arabian camels; the two-humped domestic Bactrian (*Camelus bactrianus*); and the wild Bactrian camel (*Camelus ferus*) (Rao *et al.*, 1970). The extinction of wild dromedaries underscores the importance of conserving domestic popolations (Roth and Merz, 2013).

Dromedaries, constituting ~90% of the global camel population of 27.7 million, are primarily found in North Africa and parts of Asia (Kadim *et al.*, 2008). They valued for their milk, meat, fiber, transportation, and even dung as fuel (Burger, 2016). Their physiological traits enable them to survive in ability to survive in arid and semi-arid environments with low rainfall, extreme heat, and sparse vegetation, underscoring their role in food security. However, limited research and genetic improvement efforts, particularly in developing countries, have hindered their sustainable management (Fitak *et al.*, 2016).

In Iran, the Turkmen camel, *Camelus dromedaries* breed, holds significant cultural and economic value in Golestan Province. Prized for its milk and meat, this breed exhibits unique adaptations to local conditions, distinguishing it from other dromedary and Bactrian populations. Despite its importance, its genetic diversity and conservation status remain poorly characterized, exacerbated by threats like climate change, habitat loss, and genetic dilution through crossbreeding.

Microsatellites, or simple sequence repeats (SSRs), are powerful tools for studying genetic variation, population structure, and evolutionary relationships due to their high polymorphism, codominance, and multiallelic nature (Mahmoud et al., 2012; Piro et al., 2020; Yousefzadeh et al., 2024). Compared to single nucleotide polymorphisms (SNPs), SSRs offer higher discriminatory power per locus, requiring fewer markers for population-level analyses and are cost-effective for resource-limited settings (Nobari et al., 2010; Guichoux et al., 2011; Fatemi et al., 2025). However, fragmented draft genomes for C. dromedarius, C. bactrianus, and C. ferus limit comprehensive genetic studies, necessitating de novo SSR marker development using next-generation sequencing (NGS) (Fitak et al., 2016; Ming et al., 2020). NGS enables efficient identification of genome-wide SSRs,

even without fully annotated reference genomes, facilitating the discovery of novel, breed-specific markers.

This study aims to address these gaps by developing novel SSR markers for the Iranian Turkmen camel using NGS. By comparing these markers with those from *Camelus dromedarius*, *Camelus bactrianus*, and *Camelus ferus*, we characterize genetic diversity, population structure, and breed-specific adaptations. These markers provide a foundation for targeted breeding and conservation strategies to ensure the long-term sustainability of this culturally and economically significant breed.

Material and Methods

Sampling, DNA extraction, and sequencing

Blood was collected from the jugular vein of a single female Iranian Turkmen camel (C. dromedaries), reared for hand milking, on a private farm in Golestan province, Iran. A 5ml syringe was used, and the sample was stored in a 5ml EDTA tube to prevent coagulation. Genomic DNA was extracted from whole blood using the salting-out method. DNA quality and integrity were assessed via 1% agarose gel electrophoresis strained with ethidium bromide, visualized under UV light. DNA concentration was quantified using a Oubit Fluorometer, yielding 304 ng/ul, confirming high molecular weight DNA suitable for sequencing. The DNA sample was sent for high-throughput whole genome sequencing using the Illumina HiSeqTM2000 platform generating 150 bp paired-end reads at 50x coverage. *De novo* assembly was performed to generate contigs. Although based on a single individual, this approach is sufficient for de novo SSR discovery. However, it limits the assessment of allele frequencies and population-level polymorphism, which requires a larger sample sizes and validation across multiple individuals.

Quality filtering and de novo assembly

Raw sequence reads were quality-checked using FastQC (https://www.bioinformatics.babraham.a c.uk/projects/fastqc/), confirming no adapter contamination. Reads shorter than 40 bp were removed, and a quality threshold of Q20 was applied, retaining sequences with at least 50% of bases meeting this score, using CLC Genomics Workbench (CLC Bio, Aarhus, Denmark). De

novo assembly was conducted with CLC Genomics Workbench using the following parameters: mismatch cost= 3, insertion/deletion cost= 3, length fraction= 0.5, and similarity fraction= 0.8. The assembled contings were compared to the dromedary reference genome (accession number: GCA_000803125.1) using MUMmer3 (nucmer).

SSR identification and primer design

Microsatellites (SSRs) were defined as repeat motifs of 1-10 nucleotides to include longer motifs (hepta- to deca-nucleotide repeats), which are less prone to replication slippage and stutter artifacts, enhancing specificity for population genetics and linkage mapping. SSRs were identified using SSR Locator software with the following criteria: mononucleotide motifs with repeats \geq 20, dinucleotide motifs with repeats \geq 10, trinucleotide motifs with repeats \geq 7, tetranucleotide motifs with repeats ≥ 5 , pentanucleotide and hexanucleotide motifs with repeats ≥ 4 , and hepta-, octa-, nona-, and decanucleotide motifs with repeats ≥ 3 . Primer pairs were designed in Primer3 for amplicons of 100-180 bp, with primer lengths of 15-25 bp (optimal: 20 bp), temperature range of 55-65°C (default 60°C), and the GC content was targeted to be between 40-60% (optimal: 50%) to ensure robust amplification and specificity (no experimental

validation was performed in this study). SSR identification and primer design were also performed for Arabian dromedary (CamDro3), domestic Bactrian (Ca_bactrianus_MBC_1.0), and wild Bactrian (BCGSAC_Cfer_1.0) genomes, downloaded in FASTA format from the NCBI genome database. Consistent criteria were applied using SSR Locator to enable comparative analyses.

Results

DNA extraction and sequence assembly

Genomic DNA extracted from a female Iranian Turkmen camel (Camelus dromedaries) yielded a concentration of 304 ng/ul, quantified using a Oubit Fluorometer. Whole-genome sequencing on the Illumina HiSeq™ 2000 platform produced 589,326,158 clean 150 bp paired-end reads at 50x coverage, ensuring robust SSR detection. De novo assembly generated 235,978 contigs with an N50 of 8,526 bp, ranging from 200 bp to 683,514 bp (SD = 25,268). Contig size distribution (Fig. 1) showed a peak at ~200 bp, reflecting fragmentation typical of short-read sequencing. Approximately 92% of contigs aligned to the genome Arabian dromedary reference (GCA 000803125.1) using MUMmer3. confirming high sequence homology.

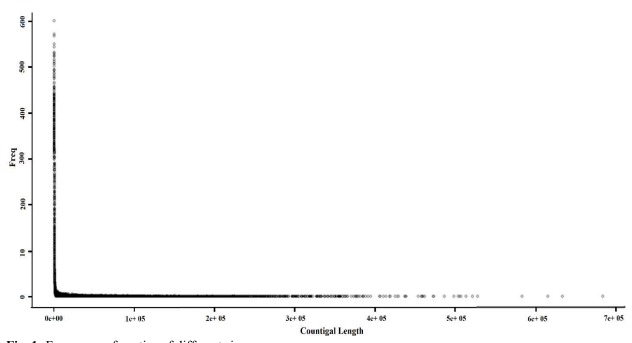


Fig. 1: Frequency of contigs of different sizes

SSR identification and comparative analysis

From the Turkmen camel genome, 152,381 microsatellites (SSRs) were identified, with 151,556 retained after filtering for ≥90%

sequence identity and excluding low-complexity regions. Dinucleotide repeats were most abundant (38.80%), followed by tetranucleotide (21.85%), and mononucleotide (16.20%) repeats (Table 1).

Table 1. Number of different motifs for Turkmen dromedaries, Arabian dromedaries, domestic Bactrian camels, and wild Bactrian camels.

| Ecotypes | Frequency | 1-mer | 2-mer | 3-mer | 4-mer | 5-mer | 6-mer | 7-mer | 8-mer | 9-mer | 10-mer | Total |
|--|-----------|-------|-------|-------|-------|-------|-------|-------|-------|-------|--------|--------|
| Camelus dromedarius (Turkmen camel) | N | 24550 | 58808 | 8414 | 33112 | 14182 | 4121 | 4583 | 2373 | 923 | 502 | 151556 |
| | % | 16.2 | 38.8 | 5.5 | 21.8 | 9.3 | 2.7 | 3.0 | 1.6 | 0.6 | 0.3 | |
| Camelus dromedarius (Arabian camel) | N | 29369 | 62208 | 8131 | 36188 | 13963 | 4042 | 4296 | 2496 | 918 | 497 | 162108 |
| | % | 18.1 | 38.4 | 5.0 | 22.3 | 8.6 | 2.5 | 2.7 | 1.5 | 0.6 | 0.3 | |
| Camelus bacterianus | N | 28929 | 59913 | 8084 | 37820 | 14379 | 4311 | 4394 | 2660 | 907 | 484 | 161881 |
| (Bactrian camel) | % | 17.9 | 37.0 | 5.0 | 23.4 | 8.9 | 2.7 | 2.7 | 1.6 | 0.6 | 0.3 | |
| Camelus ferus (Wild Bactrian camel) | N | 30182 | 60399 | 8142 | 36674 | 14208 | 4270 | 4382 | 2508 | 882 | 450 | 162097 |
| | % | 18.6 | 37.3 | 5.0 | 22.6 | 8.8 | 2.6 | 2.7 | 1.5 | 0.5 | 0.3 | |

Primer pairs were designed for 144,184 loci (95%) for genotyping. Repeat thresholds were: mononucleotide \geq 20, dinucleotide \geq 10, trinucleotide \geq 7, and tetranucleotide \geq 5, pentanucleotide and hexanucleotide \geq 4, and hepta- to deca-nucleotide \geq 3. Fig. 2 illustrates the

number of repeats for each identified motif across the genome of the Turkmen camel and other camels for comparison purposes. The 10 motifs with the highest number of repeats for each camel are listed in Table 2.

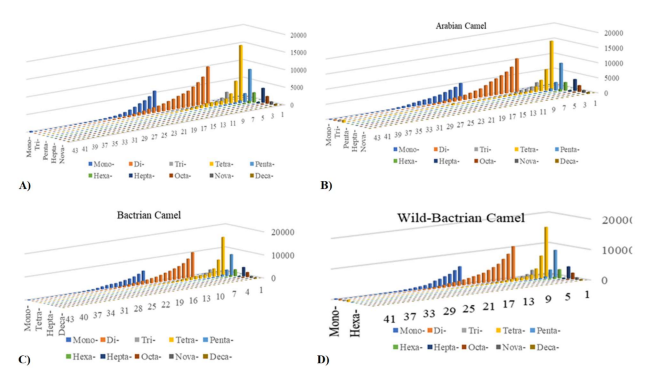


Fig. 2. Number of repeats for motifs ranging from Mono-nucleotide to Deca-nucleotide in the genomes: A) Turkmen camels; B) Arabian camels; C) *Camelus bacterianus*; D) Wild Bactrian camels.

Table 2. Frequency (percentage) of the 10 most repeated motifs in Turkmen camels, Arabian camels, Bactrian camels, and Wild Bactrian camels.

| Strand1/Strand2 | Turkmen camels | Arabian camels | Bactrian camels | Wild Bactrian camels |
|-----------------|----------------|----------------|-----------------|----------------------|
| A/T | 14.49 | 16.8 | 17.44 | 18.20 |
| C/G | 1.64 | - | - | - |
| AC/GT | 16.40 | 16.6 | 16.71 | 16.56 |
| CA/TG | 13.16 | 13.1 | 13.24 | 13.52 |
| TA/TA | 3.22 | 3.0 | 2.06 | 2.14 |
| AT/AT | 2.96 | 2.6 | 1.80 | 1.89 |
| GA/TC | 1.60 | 1.7 | 1.73 | 1.72 |
| AG/CT | - | 1.4 | 1.44 | 1.44 |
| AAAC/GTTT | 2.12 | 2.1 | 2.12 | 2.12 |
| AAAAC/GTTTT | 1.72 | 1.7 | 1.71 | 1.72 |
| TAAA/TTTA | 1.39 | 1.4 | - | - |
| CAAA/TTTG | - | - | 1.44 | 1.37 |

The Turkmen camel exhibited 6,201 unique motifs, significantly higher than Arabian dromedary (5,895), domestic Bactrian (5,979), and wild Bactrian (5,847) camels ($\chi^2 = 7.14$, p= 0.007, df= 1) (Table 3). Genomic site frequency for unique motifs (Fig. 3) showed larger motifs (hepta- to deca-nucleotide) typically occurred at single sites, while mono-, di-, and trinucleotide motifs appeared at multiple sites. Shared motifs across species are summarized in Table 4.

The Turkmen camel had 1,179 novel motifs unique to this breed, absent from NCBI databases, compared to 932 for Arabian dromedary, 971 for domestic Bactrian camel, and 945 for wild

Bactrian camel. Dromedaries (Turkmen and Arabian) shared 2,870 novel motifs unique to their clade, while Bactrian camels shared 2,633. These novel motifs, defined as previously unreported in the NCBI databases, enhance breedspecific genetic characterization and conservation efforts. Of the 151,556 SSRs, primers were designed for 89-98% of motifs across categories (e.g., 89% for dinucleotide, 97% for trinucleotide) (Table 5), with counts of 24550, 58808, 8414, 33112, 14182, 4121, 4583, 2373, 923, and 502 for di, tri, tetra, penta, hexa, hepta, octa, nona, and deca nucleotid repeats, respectively.

Table 3. Number of unique motifs for each group and their nucleotide lengths across the genome of different camels.

| Ecotypes | Motif (Xmer) | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
|---------------------|------------------------|--------|---------|--------|---------|--------|--------|--------|-------|-------|-------|
| Torkaman camel | Unique Motifs | 4 | 12 | 60 | 225 | 573 | 871 | 1864 | 1440 | 716 | 436 |
| | $NoMG^1$ | 579835 | 863316 | 88329 | 234579 | 70704 | 19356 | 14336 | 7640 | 2907 | 1544 |
| | $MotNucCou^2$ | 579835 | 1726632 | 264987 | 938316 | 353520 | 116136 | 100352 | 61120 | 26163 | 15440 |
| Arabian camel | Unique Motifs | 4 | 12 | 60 | 215 | 566 | 821 | 1756 | 1385 | 663 | 413 |
| | $NoMG^1$ | 714047 | 913615 | 110567 | 343895 | 79592 | 21128 | 13834 | 10583 | 3058 | 1658 |
| | $MotNucCou^2$ | 714047 | 1827230 | 331701 | 1375580 | 397960 | 126768 | 96838 | 84664 | 27522 | 16580 |
| Bactrian camel | Unique Motifs | 4 | 12 | 59 | 223 | 572 | 855 | 1854 | 1361 | 630 | 409 |
| | $NoMG^1$ | 705731 | 884515 | 86045 | 282223 | 72380 | 20460 | 14048 | 10766 | 3297 | 1564 |
| | $MotNucCou^2$ | 705731 | 1769030 | 258135 | 1128892 | 361900 | 122760 | 98336 | 86128 | 29673 | 15640 |
| Wild Bactrian camel | Unique Motifs | 4 | 12 | 60 | 217 | 544 | 839 | 1802 | 1342 | 634 | 393 |
| | $NoMG^1$ | 741827 | 933429 | 129738 | 338148 | 87538 | 23934 | 14277 | 11101 | 3293 | 1562 |
| | MotNucCou ² | 741827 | 1866858 | 389214 | 1352592 | 437690 | 143604 | 99939 | 88808 | 29637 | 15620 |

NoMG1: The number of motifs in the genome; MotNucCou2: The motif's nucleotide counts on the genome

Table 4. Number of shared unique motifs among different camels.

| Camels | Turkmen camel | Wild Bactrian camel | Arabian camel | Bactrian camel |
|---------------------|---------------|---------------------|---------------|----------------|
| Torkaman camel | | | | |
| Wild Bactrian camel | 3327 | | | |
| Arabian camel | 3777 | 3384 | | |
| Bactrian camel | 3508 | 3745 | 3376 | |

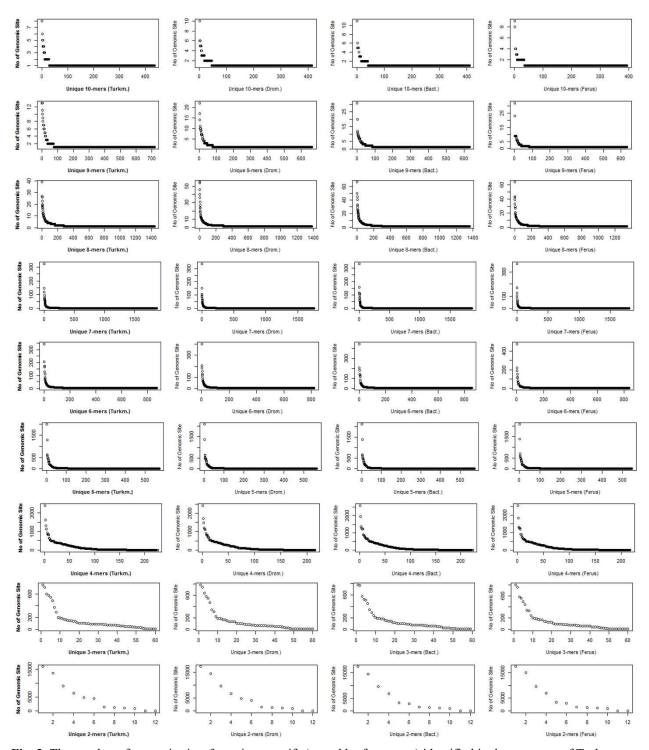


Fig. 3. The number of genomic sites for unique motifs (sorted by frequency) identified in the genomes of Turkmen, Arabian, Bactrian, and wild Bactrian camels.

Additionally, 19,425 codon-repeat loci were identified, predominantly leucine-rich (42%), with 68% in coding regions, suggesting possible roles in stress response and immune function genes (e.g., HSP70 and IL10), pending functional

validation (Table 6). Repeat count distributions are shown in Fig. 4, and motif frequencies in the Fig. 5. A supplementary table details unique motifs, their genomic positions, and nucleotide counts across species (Supplement 1).

Table 5. Number and percentage of primers developed for each SSR motif

| | 1-mer | 2-mer | 3-mer | 4-mer | 5-mer | 6-mer | 7-mer | 8-mer | 9-mer | 10-mer |
|---------|----------------|-------|-------|-------|-------|-------|-------|-------|-------|--------|
| Number | 21928 | 57205 | 8001 | 31186 | 13617 | 4056 | 4450 | 2332 | 907 | 490 |
| Percent | 0.89 | 0.97 | 0.95 | 0.94 | 0.95 | 0.97 | 0.96 | 0.94 | 0.94 | 0.98 |
| Total | 144184 (0.95%) | | | | | | | | | |

Table 6. Amino acid codon SSR loci, their frequencies, and repeat counts.

| Amino acid | Number loci | Total repeats | Amino acid | Number loci | Total repeats |
|------------|-------------|---------------|------------|-------------|---------------|
| Leu | 2526 | 20490 | Pro | 885 | 4470 |
| Asn | 1812 | 16021 | Arg | 793 | 4324 |
| Ile | 1335 | 12901 | Gly | 858 | 4253 |
| Term | 1070 | 10589 | His | 651 | 4013 |
| Tyr | 959 | 9687 | Gln | 585 | 3753 |
| Lys | 1309 | 6531 | Ala | 610 | 3598 |
| Phe | 1362 | 6424 | Met | 302 | 2513 |
| Ser | 905 | 5681 | Asp | 306 | 2276 |
| Cys | 786 | 4926 | Glu | 368 | 2039 |
| Val | 870 | 4852 | Trn | 349 | 1922 |
| Thr | 784 | 4499 | total | 19425 | 135762 |

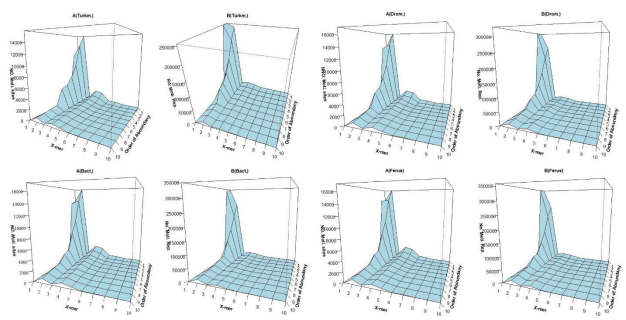


Fig. 4. Motifs identified in the genomes of Turkmen, Arabian, Bactrian, and wild Bactrian camels: A) The number of genomic sites; B) Repeat counts; Arabian showed with Drom.

Discussion

Contigs length is a critical factor in *de novo* whole-genome sequencing, influencing assembly quality for downstream analyses like SSR detection. Longer contigs reduce gaps, improving continuity and accuracy (Chen *et al.* 2015). The N50 metric, representing the contig length at which 50% of the genome is covered by contigs

of that length or longer, assembly contiguity (Desai *et al.*, 2013). Here, de novo assembly yielded 235,978 contigs with an N50 of 8,526 bp, sufficient to support robust SSR detection, despite a peak in short ~ 200 bp (Fig. 1). This fragmentation, typical of Illumina HiSeq 2000 short-read sequencing, may underrepresent SSR-rich regions, introducing bias in motif identification (Rhie *et al.*, 2021). Further studies

using long-read sequencing (e.g., PacBIO, Oxford Nanopore) could enhance contiguity and resolve repetitive regions (Srivastava et al., 2019). Sequencing generated 589,326,158 clean reads at 50x coverage, calculated using SAMtools depth (v1.9, minimum mapping quality of 20) ensuring reliable SSR detection, including lowfrequency motifs (Tóth et al., 2000; Jeanjean et al., 2025). Alignment of 92% of Turkmen camel contigs to the Arabian dromedary reference genome (GCA 000803125.1) using MUMmer3 confirmed high sequence homology and assembly quality. However, reliance on the Arabian dromedary reference may introduce mapping biases for Bacterian camels, where genomic divergence could lead to underrepresentation of lineage-specific SSRs.

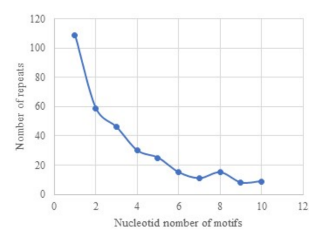


Fig. 5. The highest repeat counts for each SSR motif

The Turkemen camel genome contained 151,556 SSRs, with dinucleotide repeats (38.80%) most prevalent, followed by tetranucleotide (21.85) and mononucleotide (16.20%) repeats (Table 1). This distribution aligns with patterns in other camels (Hedayat-Evrigh *et al.* 2018; Barazandeh, *et al.* 2020; Shaltenbay *et al.*, 2025) and animals (Yan *et al.*, 2008; Bakhtiarizadeh *et al.*, 2012, Abe & Gemmell 2014, Sadder *et al.*, 2015), where shorter repeats predominate due to higher mutation rates and replication slippage (Tóth *et al.*, 2000). In contrast, some plants show higher trinucleotide repeat frequencies (Varshney *et al.*, 2002; Durand *et al.*, 2010; Joshi *et al.*, 2010; Zhou *et al.*, 2016; Wang *et al.* 2017).

Table 2 demonstrated that the frequent dinucleotide across all camels was AC/GT, with

an approximate frequency of 16% for all camels. This was followed by the mononocleotide A/T, which exhibited variable frequencies among different camels. While breed-specific motifs (*e,g,* C/G in Turkmen, CAAA/TTTG in Bacterian camels) suggest evolutionary divergence, valuable for breed identification and conservation efforts.

Compared to Barazandeh et al. (2020), who reported AC/TG (54%) and AT/TA (32.8%), with GC/CG having the lowest frequency (1.2%). Their most frequent 3-mer motif was GCC/GGC (19.2%), followed by AGC/GCT (10.3%). These differences likely stem from methodological distinctions: our study used DNA-seq, capturing both coding and non-coding regions, whereas Barazandeh et al. (2020) used RNA-seq, which focuses on transcribed regions. Coding regions, subject to purifying selection, may limit SSR expansion to maintain protein function, leading to lower 2-mer frequencies in RNA-seq data. In contrast, non-coding regions, under relaxed selection. allow greater SSR variability, explaining the higher 2-mer frequency in our DNA-sea This highlights data. complementary nature of DNA-seq and RNA-seq approaches in camel genomics, as they capture different compartments and selective pressures (Tóth et al., 2000).

The Turkmen camel exhibited 6,201 unique motifs, surpassing Arabian dromedary (5,895), domestic Bactrian (5,979) and wild Bactrian (5,847) camels (Table 3), indicating greater SSR diversity potentially linked to environmental adaptations (Pandey et al., 2018). Of these, 1,179 novel motifs were unique to the Turkmen camel, enhancing its genetic characterization. Dromedaries shared 2,870 novel motifs, and Bactrian camels 2,633, reflecting clade-specific variation. Although our study analyzed a single individual, Turkmen camels belong to a broader dromedary population shaped by geographic isolation, migration, and crossbreeding. Future multi-individual studies will be required to assess population substructure, genetic drift, and gene flow, which may significantly influence SSR diversity.

The results presented in Table 5 showed that out of a total of 151,556 SSRs identified in the Turkmen camel, primers could be designed for 144,184 (95%) of them. Therefore, the SSR

markers developed and their corresponding primers in this study provide a valuable resource for future research on genetic diversity, historical and geographical demography, evolution, and conservation genomics of various camel species. SSR markers, along with their corresponding primers, can also aid in enhancing genetic studies of the species with limited genetic characterization and registration.

Among 19,425 codon-repeat loci, 68% were in coding regions, with leucine-rich repeats (42%) (Table 6) suggesting roles in stress response and immune function (*e.g.*, HSP70, IL10) genes (Kumar *et al.*, 2016; Glazko *et al.*, 2023). These findings warrant functional validation.

Limitations include the reliance on a singleindividual sample, restricting allele frequency and polymorphism assessments. Short-read sequencing technologies (e.g., Illumina) are prone to systematic biases in repetitive and GC-rich regions, which may result in underrepresentation of certain SSR motifs. This limitation highlights the need for integrating long-read sequencing platforms in future camel genomics to overcome repetitive region biases. In the end, the SSR primers were designed in silico and were not experimentally validated; while computational primer design using Primer3 provides a strong foundation, laboratory-based testing is necessary to confirm amplification efficiency, specificity, and reproducibility.

Conclusion

This study identified 151,556 SSR motifs in the Turkmen camel genome, with dinucleotide tetranucleotide (21%),mononucleotide (16%) repeats most prevalent. The Turkmen camel exhibited the highest SSR diversity (6,201 unique motifs) significantly exceeding other camels ($\chi^2 = 7.14$, p= 0.007, df= 1), with 78 to 510 additional unique motifs. Primer pairs for 144,184 loci (95%) support population genetic studies, linkage mapping, and conservation. Additionally, 19,425 codon-repeat loci, predominantly leucine-rich (42%), with 68% in coding regions, suggest roles in stress response and immune function. Despite robust 50x coverage and N50 of 8,526 bp, limitations include single-individual sample. population-level analyses, and short contings potentially missing SSR-rich regions. Although

144,184 primer pairs were computationally designed using Primer3, reliance on the Arabian dromedary reference genome may also introduce alignment bias for Bactrian camels. Further studies should use longer-read sequencing (e.g., PacBio, Oxford Nanopore), larger sample sizes, and functional asseys to validate codon-repeat roles and marker polymorphism across Camelus species, enhancing breeding and conservation strategies for the Turkmen.

Acknowledgment

The authors gratefully acknowledge for the research funding provided for this project (No. 02/3365) by Institute of Science and High Technology and Environmental Sciences, Graduate University of Advanced Technology, Kerman, Iran.

Data availability statement

The sequencing data generated in this study are available from the corresponding author upon request.

Conflict of interest

There is no conflict of interest between authors.

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