

Journal of Advances in Biology & Biotechnology

Volume 27, Issue 8, Page 1316-1324, 2024; Article no.JABB.121173 ISSN: 2394-1081

Characterization of Diverse Carrot (*Daucus carota* **L.) Genotypes Using Long Noncoding RNA Based Microsatellite Markers**

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

Article Information

DOI[: https://doi.org/10.9734/jabb/2024/v27i81255](https://doi.org/10.9734/jabb/2024/v27i81255)

Open Peer Review History:

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: <https://www.sdiarticle5.com/review-history/121173>

> *Received: 04/06/2024 Accepted: 06/08/2024 Published: 10/08/2024*

Original Research Article

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Cite as: Hundekar, Aishwarya, Sarvamangala S C, Manjunath H, Mahantesha B. N. Naika, and Bapurayagouda P. 2024. "Characterization of Diverse Carrot (Daucus Carota L.) Genotypes Using Long Noncoding RNA Based Microsatellite Markers". Journal of Advances in Biology & Biotechnology 27 (8):1316-24. https://doi.org/10.9734/jabb/2024/v27i81255.

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ABSTRACT

In the pursuit of enhancing carrot breeding programs, our study focused on the experimental validation of selected long non-coding RNAs (lncRNAs) using end-point polymerase chain reaction (PCR). Twenty-seven carrot (*Daucus carota* L.) genotypes, including Asiatic and Western varieties, were grown in RCBD design and evaluated for root traits at UHS Bagalkot, Karnataka, India, during 2022-2023. These genotypes were used to validate five SSR markers developed for lncRNAs in carrots. From an initial set, five lncRNAs were chosen for further experimental validation based on their bit score and the presence of simple sequence repeat (SSR) motifs. Our novel approach involved the development of 27 imperfect SSR markers and five perfect SSR markers specifically targeting these lncRNAs. The integration of SSR markers with lncRNA studies is unprecedented in carrot breeding, opening new avenues for genetic improvement. Among the five lncRNA-derived SSR (lncSSR) markers developed, *Dc*LNC55 and *Dc*LNC62 exhibited polymorphisms across 27 diverse carrot genotypes. In conclusion, our study highlights the innovative use of lnc-SSR markers in carrot breeding, demonstrating their utility in genetic diversity studies and their potential role in uncovering the functional importance of lncRNAs in crop phenotypes. The findings present significant implications for future carrot breeding strategies, providing a foundation for integrating molecular markers and lncRNA research in carrot and other related horticultural crop improvement programs.

Keywords: Carrot; long non-coding RNA; polymerase chain reaction; simple sequence repeat; polymorphism.

1. INTRODUCTION

Carrot (*Daucus carota* L.) is a biennial herb with a chromosome number of $2n = 2x = 18$ with a genome size of 473Mb and belongs to the Apiaceae family [1]. Fewer studies are available on lncRNAs in carrots compared to model organisms like Arabidopsis despite the availability of high-quality genome sequences in carrots as large numbers of lncRNAs have been found in nature [2].

Long non-coding RNAs (lncRNAs), which exceed 200 nucleotides in length and do not code for proteins, play a crucial role in the regulation of gene expression at multiple levels [3]. To classify a transcript as "noncoding," it is assessed based on the absence of an extensive open reading frame (typically over 100 codons) and the lack of codon conservation [4]. In the nucleus, lncRNAs regulate gene expression through transcriptional and epigenetic mechanisms, including the modulation of chromatin compaction and the establishment of chromosomal connections [5]. They also influence gene expression by regulating the loading of histone or chromatin modifiers to chromatin [6]. Additionally, lncRNAs can directly affect transcription through the formation of R-loops, interference with RNA polymerase machinery, and transcription of the lncRNA locus. In the cytoplasm, a subset of lncRNAs is involved in controlling mRNA turnover and translation, highlighting their

multifaceted roles in gene expression regulation [7].

Plant lncRNAs, transcribed by RNA polymerases PolI, PolII, and PolV, originate from various genomic regions, including enhancers, promoters, gene introns, pseudogenes, and as antisense transcripts to other genes [8,9]. These transcripts can be classified into different types based on their genomic position relative to protein-coding genes, such as natural antisense
transcripts (lncNATs), intronic lncRNAs, transcripts (lncNATs), intronic lncRNAs, intergenic lncRNAs (lincRNAs), and sense lncRNAs. lncNATs align with or diverge from sense strand transcripts, potentially modulating gene regulation, while intronic lncRNAs are transcribed from within introns of protein-coding genes without exon overlap. Sense lncRNAs share promoters with protein-coding genes and transcribe from regions overlapping exons [5,10]. Recent studies have revealed the involvement of lncRNAs in various biological processes, including flowering, reproduction, photomorphogenesis, vernalization, organ development, cell cycle control, and responses to biotic and abiotic stresses. The experimental validation of lncRNAs is essential to elucidate their interactions with transcriptional and translational components, providing a strong rationale for exploring their regulatory roles in different metabolic pathways in plants [11].

Simple Sequence Repeat (SSR) markers, also known as microsatellites, are highly polymorphic and co-dominant markers widely used in genetic research. They consist of short, repetitive DNA sequences that are scattered throughout the genome, making them ideal for assessing genetic diversity and mapping genetic traits. Due to their high reproducibility and ease of analysis, SSR markers are invaluable tools in plant breeding and molecular biology [12]. In reality, there are few SSR markers unique to the carrot genome, and the earliest accounts of the developing SSRs date back only a few years [13,14]. Cavagnaro et al., 2011 used BACend sequences and an SSR-enriched genomic collection to create 300 SSR markers in carrots. SSRs were discovered by Iorizzo et al. in a assembly of the carrot transcriptome from four genotypes and provided a collection of 114 markers they identified by in silico investigations as polymorphic. Considering Since then, most research on the genomics of carrots have employed these SSR markers [15].

2. MATERIALS AND METHODS

2.1 Plant Material

The present study was conducted at the Department of Biotechnology and Crop Improvement, University of Horticultural Sciences, Bagalkot. Total of 27 genotypes carrot (*Daucus carota* L.) germplasm lines were grown, inclusive of Asiatic/Eastern and European/Western were used. The detailed names and colour description of the 27 carrot genotypes are listed in Table 1. The vegetative phase carrot root evaluation experiment was conducted at sector 1 farm (16°12'N, 75°12'45 E), University of Horticultural Sciences (UHS), Bagalkot, Karnataka, India during 2022-2023. It is part of a semi-arid tropical region with a mild climate, with typical temperatures between 23 °C and 35 °C and receives an average rainfall of 318 mm.

2.2 Primer Designing for lncRNAs

The lncRNAs were identified in the Alnc database and Primers were designed by searching microsatellite regions in the lncRNA transcripts. Sequences flanking the microsatellite regions in lncRNA were targeted for primer design using the Krait tool [16] by targeting the microsatellite flanking regions. The primer pairs were designed based on standard criteria, such as predicted melting temperature of 50-60^oC and Guanine-Cytosine (GC) content of 45-55 percent for designing the primer pairs to obtain PCR amplicon length of 100-200 bp (Table 2).

Table 1. List of carrot (*Daucus carota* **L.) genotypes used for evaluation of genetic variability**

SI. No.	Genotypes	Description	
1.	UHSBC-34-2	Orange	
2.	UHSBC-17	Orange	
3.	UHSBC-31-2	Orange	
4.	UHSBC-31-1	Orange	
5.	UHSBC-32-2	Orange	
6.	UHSBC-65	Released variety, Red colour (Pusa Rudhira)	
7.	UHSBC146	Orange (Improved-population-2)	
8.	UHSBC-23-1-1	Orange-red	
9.	UHSBC-66	Black, released (Pusa Asita)	
10.	UHSBC-31-4	Orange	
11.	UHSBC-34-2-1	Orange	
12.	UHSBC-100	Dark Orange (Western)	
13.	UHSBC-46-1-1	Orange	
14.	UHSBC-151	Reddish orange	
15.	UHSBC-23-1	Orange	
16.	UHSBC-150	Reddish orange	
17.	UHSBC-1	Pale orange	
18.	UHSBC-152	Orange	
19.	UHSBC-160	Orange	
20.	UHSBC-17-2	Orange	
21.	UHSBC-59-66	White	
22.	UHSBC-59	White	
23.	UHSBC-17-1	Orange	
24.	UHSBC-117	Orange	
25.	UHSBC-150-1	Orange	

2.3 Genomic DNA Extraction and Amplification of lncSSR Markers

The genomic DNA was extracted from carrot leaves using CTAB extraction method [17]. DNA amplification was carried out using lncRNA-SSR primers for 27 genotypes of carrot to check their utility and functional validation. Amplification of DNA for the respective target lncRNA was performed using polymerase chain reaction (PCR) in a 10 μl reaction mixture for lncRNA validation each containing 2 μl template DNA (50 ng/μl), 0.5 μl forward primers (5 μmol/l), 0.5 μl reverse primers (5 μmol/l), 2 μl ddH2O and 5 μl 2 × Taq PCR Master mix. The PCR program was as follows: 1 min at 95 °C, 20 s denaturing at 95 °C, 45 s annealing at 47-53 °C (Annealing temperature was optimised for each primer set using gradient PCR before actual amplification) and 30 s elongation at 72 °C followed by a final step at 72 °C for 2 min. The details of primer sequences used for amplification of lncRNA is presented in the Table 3. After the PCR, fractionation was done using agarose gel electrophoresis to assess the amplification of target genes followed by EtBr staining. 3.5 % agarose gel was used for separation of lnc-SSR markers in 1 \times TBE buffer at initial 85V for 15 minutes followed by 120 V for 1 h. Subsequently, the banding pattern was visualized using a gel documentation system and clear images of the bands were captured and scoring was performed. The clear and unambiguous bands of all the polymorphic lncSSR markers were scored for 27 carrot genotypes and calculated manually.

3. RESULTS AND DISCUSSION

3.1 Identification of SSR Regions in lncRNA and Marker Development

The identified sets of transcripts from the Alnc database were used to search for simple sequence repeats (SSR) in the Krait tool. 32 sequences were considered to contain microsatellites or SSR motifs. The identified SSRs have mono, di, tri, tetra, and hexa nucleotide repeat motifs. Among 32, perfect SSRs were five (Table 2a) and the remaining 27 were imperfect SSRs (Table 2b). From the perfect SSRs- two SSRs contain mono- a nucleotide repeat motif, two SSRs contain di- a nucleotide repeats motif and the remaining one has hexa- nucleotide repeat motif. Among the imperfect SSRs- five SSRs contain mononucleotide repeat motif, eleven SSRs contain dia nucleotide repeat motif, six SSRs contain trinucleotide repeat motif, four SSRs contain quadnucleotide repeat motif and the remaining one has hexa- nucleotide repeat motif. One perfect SSR (STRG.55.1) was chosen for validation as it consists of a hexa-nucleotide repeat motif. Four among the imperfect SSRs, STRG.25.1 (dinucleotide repeat motifs), STRG.62.1 and STRG.70.1 (tri-nucleotide repeat motifs), and STRG.91.1 (tetra nucleotide repeat motifs) were chosen for validation because of the presence of a greater number of repeats and higher chances of polymorphism.

3.2 Validation of lncRNA by PCR

A total of five SSR markers were used for the validation of 27 genotypes which yielded clear and scoreable bands. Among the five SSR markers considered for validation *Dc*Lnc91, *Dc*Lnc25 and *Dc*Lnc70 showed monomorphic bands of 140, 100, and 140bp respectively. While markers *Dc*LNC55 and *Dc*LNC62 showed polymorphism with band sizes ranging from 100- 125 bp and 200-480 bp respectively (Table 4) and gel profile (Fig. 1) indicating their suitability in the carrot genetic diversity assessment. For validation of lncRNA, 5 lnc-SSR markers were used. Among them, 3 lnc-SSR markers showed monomorphic bands, while 2 markers showed polymorphism among 27 diverse carrot genotypes. Polymorphic SSR markers can reveal genetic diversity or variations in lncRNA sequences among different genotypes and would be further useful for mapping and understanding its role in phenotypic expression of economic traits in carrots.

3.3 Genotypic Scores for lnc-SSR Markers

For lnc-SSR allele sizing, the genotypes were analysed by comparing the sizes of the amplified bands against a 100bp reference ladder. Each genotype's band size was measured and recorded manually by comparing the DNA fragment lengths to a molecular weight marker on an agarose gel ensuring accurate identification of allele variations. The use of the 100bp ladder provides a reliable standard for consistent and reproducible results across different samples. The observation of monomorphic bands in 3 and-SSR markers and polymorphic bands in two markers suggests potential genetic variation.

Table 2a. List of lncRNA perfect SSRs designed

SI. No.	$\frac{a}{2}$ Ω မ္ဟ	짇 LĹ.		↽ ن ق	tability	Φ e Re	Tm ₂	N ن ق	Ď
19	STRG.62.1	TCTTCCTTTGATTATCCCACAGG	57.88	43.48	4	TGTGCAAACCTCCAAAGTCC	58.6	50	3.85
20	STRG.66.1	CCCTCTTTCTCAGCGCC	57.3	64.71	6.53	CGACGATCTCGAAATCATGG	56.53	50	3.66
21	STRG.70.1	AAAAGGGGAAAGTGCGGC	58.54	55.56	6.53	CAATGACCATTTTTATCAAACCCC	57.02	37.5	4.95
22	STRG.70.1	AATTGTGGGGTTTGATAAAAATGG	56.61	33.33	3.16	AATTTTGCCCCCACCACC	58.1	55.56	4.61
23	STRG.75.1	TCACTACACTAGTCTCTCTCTCC	57.65	47.83	3.71	AGGTGTATGCAGAGAGGCC	58.78	57.89	5.19
24	STRG.78.1	CCAACAACCCAGCAGCCC	61.31	66.67	5.19	GGTGTGTCTCCCCAGTAATGG	60.07	57.14	3.16
25	STRG.84.1	AGCTCGCTCACTTCTGGC	59.74	61.11	4.85	GCGACAATGATTTCTCCGGC	59.97	55	6.13
26	STRG.91.1	GGGAGGGAGGGAAGATCC	57.68	66.67	3.36	ATTATCACAATGCTCTTTTCTTCC	55.74	33.33	3.46
27	STRG.91.1	GTGATAATGTAAGTAAGTAGCTGC	55.46	37.5	5.25	TTTGTTTCCCCTCCTTGC	54.99	50	4.01

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Fig. 1. Validation of lncSSR markers

SI.	Primer ID	Forward primer sequence	Reverse primer sequence
No.		$(5^{\circ}-3^{\circ})$	$(5^{\circ}-3^{\circ})$
	DcLNC55	TTTCTTGGGTAAAGCAAGG	GACTTTCTTAAAGGCGATACC
2.	DcLNC62	CTCCTTTATAATTTAACAGGTGGG	CCTCCAAAGTCCAAAGTGC
3.	DcLNC91	GTGATAATGTAAGTAAGTAGCTGC	TTTGTTTCCCCTCCTTGC
4.	DcLNC ₂₅	TGCTGTTTGCTTTCCCGG	GCGTGTTTATTCCAAAATGGCC
5.	DcLNC70	AAAAGGGGAAAGTGCGGC	CAATGACCATTTTTATCAAACCCC

Table 3. The details of lnc-SSR markers used for validation in endpoint PCR

Table 4. List of the lnc-SSRs amplification data in carrot (*Daucus carota* **L.) genotypes**

SI. No.	Primer ID	Annealing temperature (°C)	Observed amplicons size (bp)
. .	DcLNC55	47	100-125
າ	DcLNC62	48	200-480
3.	DcLNC91	48	140
	DcLNC ₂₅	53	100
-5.	DcLNC70	50	140

4. CONCLUSION

The identification of polymorphic bands in two out of five lncRNA SSR markers used for validation indicates genetic variation in the genotypes. This study underscores the pivotal role of long non-coding RNAs (lncRNAs) in the genetic and phenotypic diversity of carrot genotypes. By developing novel lncRNA-derived SSR markers, we have provided useful tools for assessing genetic variation in carrots. The polymorphism observed in these markers among diverse carrot genotypes highlights their potential application in carrot breeding programs. Our findings open new avenues for integrating molecular markers and lncRNA research in crop improvement strategies.

FUTURE SCOPE

Generating genome-wide lncRNAs and utilizing the polymorphic markers in future analyses will enhance insights into the genetic diversity of regulatory roles of these non-coding sequences. They help breeders to understand the functional roles and regulation of important genes influencing carrot morphology and other phenotypic characteristics. This knowledge can be leveraged to develop new carrot varieties with desirable traits, enhancing breeding efficiency and outcomes.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that NO generative AI technologies such as Large Language Models (ChatGPT, COPILOT, etc) and text-to-image

generators have been used during writing or editing of manuscripts.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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