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MOLECULAR SIGNATURE OF nrDNA-ITS MARKER IN ISODON RUGOSUS (LAMIACEAE)

Devendra Kumar Srivastava^{1*}, Pekhna Bansal¹, Pradeep Kumar Singh¹, Manjit Inder Singh Saggoo²

Eternal University, Department of Botany, Baru-Sahib, 173101 Himachal Pradesh, India

Abstract

Srivastava D.K., Bansal P., Singh P.K., Saggoo M.I.S., 2021: Molecular signature of *nrDNA-ITS* marker in *Isodon rugosus* (Lamiaceae). – Botanica, 27(1): 53–61.

Molecular features of *nrDNA-ITS* sequences in medicinal plant *Isodon rugosus* were assessed with universal *ITS-1* and *ITS-2* primers. *ITS* sequences of 46 *Isodon* species were defined for their respective *ITS-1*, 5.8S and *ITS-2* regions through *in-silico* datamining and analysis. Only in 32 species, sequence of all the three defined regions was complete, while in the rest of 14 species was partial. *Isodon rugosus* revealed five genic regions, namely, *I8S rRNA* gene (partial sequence, > 1....38), *ITS-1* (complete sequence, > 39....236), 5.8S rRNA gene (complete sequence, > 237....400), *ITS-2* (complete sequence, > 401....607) and large subunit *26S rRNA* gene (partial sequence, > 608....672). The sequence regions were recorded with variable sites (VS), singleton (ST) and species-specific parsimonious-informative (PI) sites. The information was also collected in a phylogenetic tree obtained using the maximum likelihood (ML) method.

Keywords: *Isodon rugosus*, Lamiaceae, *nrDNA-ITS* marker.

INTRODUCTION

The genus name Isodon (Schrad. ex Benth.) Spach comes from the Greek word 'isos' means equal and 'odous' (or odontos) means tooth (Bentham, 1832; Spach, 1840). It consists of more than 150 species (LIU et al., 2017), distributed chiefly in tropical to temperate East Asia and tropical Africa (MABBERLEY, 2008; Chen et al., 2017). In India, the genus is represented with nearly 16 species (including one variety), mainly from the Himalayas, Western Ghats, and Nilgiri regions (Sharma, 2018). Isodon rugosus (Wall. ex Benth.) Codd. is an essential indigenous member of Indian Lamiaceae. Species is usually considered quite variable floristically due to obscure molecular identity, taxonomic relationship at inter and intracontinental level, and thus, studied under different names: Isodon plectranthoides, Lumnitzera densiflora, Ocimum densiflorum, Plectranthus rugosus and Rabdosia rugose.

DNA based markers such as nrDNA-ITS, rbcL, matK, etc. are reliable, reproducible and unaffected by the physical form and environment. They provide a convenient tool for identifying medicinal plants, crude drugs and even commercial products (SRIVAS-TAVA & SAGGOO, 2014; ZHOU et al., 2014). Among the different DNA markers, nrDNA-ITS has been supported as a remarkable marker where plastid markers fail to discriminate the plants at the species level (Hollingsworth, 2011). Therefore, nrDNA-ITS was chosen for study in the plant *Isodon rugosus*. This research is also in addition to our previous work in the plant *I. rugosus*, where the molecular autograph of plastid marker/gene (matK) was studied (SRIVAS-TAVA et al., 2020). The present study was undertaken to provide alternative DNA marker information in the taxon *I. rugosus* that could be employed along with other marker/s for its molecular identification and barcoding.

²Punjabi University Patiala, Department of Botany, 147002 Punjab, India

^{*}Corresponding author. E-mail: devsrivastv@rediffmail.com, devsrivastv@gmail.com

MATERIALS AND METHODS

Plant leaf material was used from the same specimen of I. rugosus (IRBSA-01), which has been studied previously by Srivastava et al. (2020) in the field population of Baru-Sahib valley of Himachal Pradesh (India). Except for primer pairs 18S (forward) ITS-U1:5'-GGAAGKARAAGTCGTAACAAGG-3' and 26S (reverse) ITS-U4: 5'-RGTTTCTTTTCCTC-CGCTTA-3' (CHENG et al., 2015), genomic DNA isolation (CHASE & HILLS, 1991), software applications DNAMAN version 7.0 (Lynnon Biosoft Corporation, USA), MEGA version 7.0 (Kumar et al., 2016) and ITS sequence analysis methodology were similar to that followed in previous of *mat-k* sequence study (Srivastava et al., 2020). Phylogeny with maximum parsimony (MP) criterion was avoided due to inconsistency with low scores. Sequence boundaries of different regions (i.e. ITS-1, 5.8S and ITS-2) were

determined by comparing the aligned sequences with the sequences of other related species of *Isodon* registered in NCBI GenBank (Table 1). The complete sequence of the *ITS* region was deposited at NCBI GenBank (accession No. MH931390).

RESULTS AND DISCUSSION

The sequence amplified by the primers *18S* (forward) *ITS-U1* and *26S* (reverse) *ITS-U4* showed the overall length of amplified *nrDNA-ITS* regions with 569 nucleotides (excluding partial sequences of *18S* and *26S rRNA* gene). The *ITS* sequences of 46 different *Isodon* spp. from the BLASTn result were retrieved from the NCBI GenBank (Table 1) and analysed with a sequence of *I. rugosus* (MH931390).

In-silico datamining showed only 32 *Isodon* species with complete *nrDNA-ITS* sequences, out of which nine species (accessions No. KF032259.1,

Table 1. Isodon species from the BLASTn result and their respective GenBank accession numbers

No.	Species (abbreviations)	Gene Bank No.	No.	Species (abbreviations)	Gene Bank No.
1.	Isodon rugosus (I–1)*	MH931390	25.	I. loxothyrsus (I–25)	FJ593381.1
2.	I. adenanthus (I–2)	FJ593355.1	26.	I. lungshengensis (I–26)	FJ593382.1
3.	I. adenolomus (I–3)	FJ593356.1	27.	I. megathyrsus (I–27)#	MH117581.1
4.	I. amethystoides (I-4)#	KF032259.1	28.	I. melissoides (I–28)	FJ593385.1
5.	I. anisochilus (I–5)	KF855420.1	29.	I. nervosus (I–29)#	KF032257.1
6.	I. brachythyrsus (I–6)	KF855427.1	30.	I. oresbius (I–30)	KF855434.1
7.	I. bulleyanus (I–7)	FJ593358.1	31.	I. parvifolius (I–31)	KF855425.1
8.	I. coetsa (I–8)#	KM877351.1	32.	I. pharicus (I–32)	KF855430.1
9.	I. dawoensis (I–9)	KF855429.1	33.	I. phyllostachys (I–33)	FJ593388.1
10.	I. effuses (I–10)	KF855444.1	34.	I. pleiophyllus (I–34)	FJ593389.1
11.	I. enanderianus (I–11)#	JQ389514.1	35.	I. rosthornii (I–35)	FJ593390.1
12.	I. eriocalyx (I–12)	FJ593364.1	36.	I. rubescens (I–36)#	KF032249.1
13.	I. flabelliformis (I–13)	FJ593365.1	37.	I. rugosiformis (I–37)	FJ593391.1
14.	I. flavidus (I–14)	FJ593366.1	38.	I. scoparius (I–38)	FJ593392.1
15.	I. flexicaulis (I–15)	FJ593367.1	39.	I. sculponeatus (I–39)	FJ593393.1
16.	I. forrestii (I–16)	FJ593368.1	40.	I. serra (I–40)#	KF032262.1
17.	I. gesneroides (I–17)	FJ593369.1	41.	I. setschwanensis (I–41)	KF855412.1
18.	I. glutinosus (I–18)	FJ593370.1	42.	I. shikokianus (I–42)	KF855449.1
19.	I. grandifolius var. atuntzeensis (I–19)	FJ593371.1	43.	I. shikokianus var. occidentalis (I–43)	KF855450.1
20.	I. inflexus (I–20)#	KF032266.1	44.	I. taliensis(I–44)	FJ593394.1
21.	I. irroratus (I–21)	KF855439.1	45.	I. wikstroemioides (I–45)	FJ593396.1
22.	I. japonicas (I–22)#	KF032254.1	46.	I. xerophilus (I–46)	FJ593397.1
23.	I. leucophyllus (I–23)	KF855421.1	47.	Rabdosia setschwanen-sis var. yungshengensis (I–47)**	KF855436.1
24.	I. longitubus (I–24)	KF855445.1	48.	Mentha suaveolens***; M. canadensis***	AF369161; KC473228

^{*}Query taxon, **Rabdosia setschwanensis var. yungshengensis – Syn. – Isodon setschwanensis, ***out group species, # sequences in nine taxa already defined for their respective regions in NCBI databases.

KM877351.1, JQ389514.1, KF032266.1, KF032254.1, MH117581.1, KF032257.1, KF032249.1 and KF032262.1) were observed with defined regions (e.g. ITS-1, 5.8S, ITS-2), which were originally provided in the NCBI database (Table 1). Remaining all the species that were not defined by their respective regions hitherto is now defined and presented in Table 2. The information regarding accession numbers, length, sequence span and GC content [calculated as $G + C\% = \{(G + C) / (A + T + G)\}$ + C) $\times 100$ of the different ITS regions (i.e. ITS-1, 5.8S and ITS-2) of various Isodon species are provided in Table 2.

Amplified sequence in *nrDNA-ITS* sequence in *I. rugosus* was reported with five regions: *18s* ribosomal RNA gene (partial sequence, > 1....38), *ITS-1* (complete sequence, > 39....236), *5.8S rRNA* gene (complete sequence, > 237....400), *ITS-2* (complete sequence, > 401....607) and large sub unit (*26S rRNA* gene partial sequence, > 608....672).

ITS-1 region

The sequence of ITS-1 in I. rugosus was observed with a length of 198 nt bases. Excluding the sites with missing (or ambiguous) data and gaps produced significant alignment (SA) in the ITS-1 region at 198 nt sites (> 39.....236). From the aligned positions of the ITS-1 region of 47 taxa, in I. rugosus, 49 sites were variable (VS), 30 sites were singleton (ST) and 19 sites of which were observed as parsimonious-informative (PI). Distribution of informative sites with their respective nucleotide bases (e.g. A, T, G or C) was observed to be positioned at 60 (C), 67 (A), 77 (C), 82 (T), 84 (C), 94 (A), 100 (C), 106 (C), 114 (G), 116 (G), 147 (A), 166 (A), 167 (T), 180 (C), 183 (C), 205 (G), 213 (C), 221 (G) and 229 (C) (Table 3). The GC content in the ITS-1 region of I. rugosus was 66.16%. The spanning length of the ITS-1 region varied from 194 nt to 198 nt, with a more frequent (in 44 species) occurrence of 196 nt (Table 2). Length variation was observed in three taxa, namely, I. rugosus (> 39.....236; 198 nt); *I. brachythyrsus* (> 01.....194; 194 nt) and *I. coesta* (> 33.....227; 195 nt).

5.8S region

The spanning length of the 5.8S region of *nrDNA* in *I. rugosus* was 164 nt bases (> 237.....400). In *I. rugosus*, out of 164 aligned positions, four sites were

variable, three sites were singleton, and one (1) site was parsimonious informative. The parsimonious informative site in the aligned sequences of the 5.8S region was observed (Table 4) at position 237 (A). The GC content (G + C%) in the region was 54.88%. In 45 species, the length of the 5.8S region was observed with a similar value of 164 nt bases, while GC content was identical in 40 species (Table 2). One species, i.e. I. phyllostachys (I-33; > 197....359), was reported with single nucleotide deletion, making its spanning length of 163 nt bases (Table 2). The GC content differed in six species, namely, I. coesta (I-08, G + C%: 55.49), I. inflexus (I-20, G + C%: 54.27), I. lungshengensis (I–26, GC%: 55.49), I. parvifolius (I-31, G + C%: 54.27), I. phyllostachys (I-33, G + C%: 55.21) and I. shikokianus var. occidentalis (I-43, G + C%: 55.49).

ITS-2 region

Alignment in the ITS-2 region of I. rugosus was significant at 207 nt positions (> 401.....607). There were 38 nt sites, which were variable, 26 sites were singleton, and 12 sites were parsimonious informative. Parsimonious informative sites (Table 4) were observed at positions 418 (A), 419 (C), 424 (G), 425 (C), 429 (C), 433 (G), 450 (T), 502 (C), 551 (C), 558 (T), 565(T) and 568 (C). The GC richness was reported as 68.12% (G + C%). Among the 47 *Isodon* spp. (including *I. rugosus*), the spanning length of *ITS-2* regions was partial in 15 species (193 nt in 3 sp., 194 nt in 7 sp., 195 nt in 1 sp., 197 nt in 2 sp., 199 nt in 1 sp., 205 nt in 1 sp.) and complete in 32, excluding I. rugosus, species (207 nt in 4 spp., 208 nt in 20 spp., 209 nt in 4 spp., 210 nt in 1 sp., 211 nt in 2 spp. and 212 nt in 1 sp.) (Table 2).

It was found that only nine species of *Isodon*: *I. amethystoides* (I-4, KF032259.1), *I. coetsa* (I-8, KM877351.1), *I. enanderianus* (I-11, JQ389514.1), *I. inflexus* (I-20, KF032266.1), *I. japonicas* (I-22, KF032254.1), *I. megathyrsus* (I-27, MH117581.1), *I. nervosus* (I-29, KF032257.1), *I. rubescens* (I-36, KF032249.1) and *I. serra* (I-40, KF032262.1) were defined previously as complete for all the three regions of *nrDNA-ITS* (i.e. *ITS-1* + 5.8S + *ITS-2*) by their respective authors in NCBI database (Table 1). By taking reference to the similar definition or defined regions of these nine taxa, only 32 species (I-01, I-03, I-04, I-07, I-08, I-11, I-12, I-13,

Table 2. Defined length (L), spanning region (>) and nucleotide composition (G + C%) of the ITS-1, 5.8S, ITS-2 regions of Isodon rugosus and related species

	ITS-1		5.8S		ITS-2			
Accession No.	L > region	G + C%	L > region	G + C%	L > region	G + C%		
MH931390 (I-1)	198 > 39236	66.16	164 > 237400	54.88	207 > 401607	68.12		
FJ593355.1 (I-2)	196 > 01196	63.78	164 > 197360	54.88	205 > 361565	65.85		
FJ593356.1 (I-3)	196 > 01196	65.82	164 > 197360	54.88	208 > 361568	67.79		
KF032259.1 (I-4)	196 > 15210	65.82	164 > 211374	54.88	207* > 375567	67.15		
KF855420.1 (I-5)	196 > 01196	64.80	164 > 197360	54.88	197 > 361557	69.04		
KF855427.1 (I–6)	194 > 01194	65.46	164 > 195358	54.88	194*** > 359552	68.04		
FJ593358.1 (I–7)	196 > 01196	64.29	164 > 197360	54.88	211 > 361571	68.72		
KM877351.1 (I–8)	195 > 33227	64.62	164 > 228391	55.49	208 > 392599	66.35		
KF855429.1 (I–9)	196 > 01196	65.31	164 > 197360	54.88	194 > 361554	68.56		
KF855444.1 (I-10)	196* > 01196	64.79	164 > 197360	54.88	194* > 361554	67.53		
JQ389514.1 (I-11)	196 > 06201	65.82	164 > 202365	54.88	208 > 366573	66.35		
FJ593364.1 (I–12)	196** > 01196	64.29	164 > 197360	54.88	208* > 361568	67.31		
FJ593365.1 (I–13)	196 > 01196	64.80	164 > 197360	54.88	212 > 361572	66.03		
FJ593366.1 (I–14)	196 > 01196	65.82	164 > 197360	54.88	208 > 361568	66.83		
FJ593367.1 (I–15)	196 > 01196	65.82	164 > 197360	54.88	208 > 361568	67.79		
FJ593368.1 (I–16)	196* > 01196	65.82	164 > 197360	54.88	208 > 361568	68.27		
FJ593369.1 (I–17)	196 > 01196	65.31	164 > 197360	54.88	209 > 361569	67.94		
FJ593370.1 (I–18)	196* > 01196	65.31	164 > 197360	54.88	209 > 361569	67.94		
FJ593371.1 (I–19)	196 > 01196	64.80	164 > 197360	54.88	208 > 361568	67.79		
KF032266.1 (I-20)	196 > 15210	64.80	164 > 211374	54.27	211 > 375585	67.77		
(/	196*** > 01196	63.78	164 > 197360	54.88	194* > 361554	68.56		
KF032254.1 (I-22)	196 > 15210	65.31	164 > 211374	54.88	208 > 375582	67.31		
KF855421.1 (I-23)	196 > 01196	66.33	164 > 197360	54.88	195 > 361555	69.74		
KF855445.1 (I-24)	196 > 01196	65.31	164 > 197360	54.88	193 > 361553	67.36		
FJ593381.1 (I–25)	196* > 01196	65.82	164 > 197360	54.88	208 > 361568	67.79		
FJ593382.1 (I–26)	196 > 01196	65.82	164 > 197360	55.49	208 > 361568	67.79		
MH117581.1 (I–27)	196 > 33228	64.80	164 > 229392	54.88	208 > 393600	67.79		
FJ593385.1 (I–28)	196* > 01196	65.31	164 > 197360	54.88	208 > 361568	67.79		
KF032257.1 (I-29)	196 > 17212	66.33	164 > 213376	54.88	209 > 377585	67.94		
KF855434.1 (I–30)	196 > 01196	65.82	164 > 197360	54.88	199**** > 361559	67.34		
KF855425.1 (I–31)	196 > 01196	66.33	164 > 197360	54.27	197* > 361557	69.04		
` ′	196*** > 01196	63.78	164 > 197360	54.88	194* > 361554	68.56		
FJ593388.1 (I–33)	196 > 01196	64.29	163 ^d > 197359	55.21	208 > 360567	67.79		
, ,	196*** > 01196	62.76	164 > 197360	54.88	208*** > 361568	66.35		
FJ593390.1 (I–35)	196 > 01196	65.31	164 > 197360	54.88	208* > 361568	67.31		
KF032249.1 (I-36)	196 > 17212	64.80	164 > 213376	54.88	208 > 377584	67.79		
FJ593391.1 (I–37)	196 > 01196	65.82	164 > 197360	54.88	208 > 361568	66.83		
FJ593392.1 (I–38)	196 > 01196	65.31	164 > 197360	54.88	208 > 361568	67.79		
FJ593393.1 (I-39)	196 > 01196	64.80	164 > 197360	54.88	207 > 361567	66.67		
KF032262.1 (I-40)	196 > 17212	65.81	164 > 213376	54.88	210 > 377586	68.10		
KF855412.1 (I–41)	196* > 01196	65.31	164 > 197360	54.88	194 > 361554	69.07		
KF855449.1 (I-42)	196 > 01196	64.80	164 > 197360	54.88	193 > 361553	68.39		
KF855450.1 (I-43)	196 > 01196	64.80	164 > 197360	55.49	193* > 361553	66.84		
FJ593394.1 (I–44)	196 > 01196	65.31	164 > 197360	54.88	209* > 361569	66.99		
	196 > 01196	64.80	164 > 197360	34.88	208 > 361568	67.79		
FJ593396.1 (I–45) FJ593397.1 (I–46)	196 > 01196 196 > 01196	64.80	164 > 197360 164 > 197360	54.88 54.88	208 > 361568 207 > 361567	67.79 67.15		

L-length in nucleotide bases, N-any nucleotide base, *-N(1), **-N(2); ***-N(3), ; ****N(4), d-deletion 1.

Table 3. Distribution of informative (PI) sites and nucleotide base positions in the aligned ITS-1 region of Isodon rugosus

	ITS-1 region (> 39236)																		
Species*	60	67	77	82	84	94	100	106	114	116	147	166	167	180	183	205	213	221	229
I-01	C	A	C	T	C	A	C	C	G	G	A	A	T	C	C	G	C	G	C
I-02	_	_	_	_	_	_	_	_	A	T	C	_	_	_	_	_	_	A	_
I-03	_	_	_	_	_	_	_	_	_	T	C	_	_	_	_	_	_	_	_
I-04	_	_	_	_	_	_	_	_	_	_	C	_	_	_	_	_	_	_	Α
I-05	_	_	_	_	_	_	Т	_	_	_	C	_	_	_	_	Т	_	_	_
I-06	_	_	_	_	_	_	**	_	_	Т	C	_	_	_		_	_	_	_
I-07	Т	_	_	_	_	_	_	_	_	_	С	_	_	_	_	Т	_	_	Т
I-08	Т	_	_	_	_	_	_	_	_	_	С	_	_	_	_	_	_	_	_
I-09	_	_	_	_	_	_	_	_	_	T	Т	_	_	_	_	_	_	_	_
I-10	_	_	_	Α	_	_	_	_	_	_	С	_	_	_	Α	_	_	_	A
I-11	_	_	_	_	_	G	_	_	_	A	С	_	_	_	_	_	_	A	_
I-12	_	_	_	_	_	G	_	_	_	T	_	_	_	**	_	_	_	_	_
I-13	_	_	_	_	_	_	_	_	_	Т	С	_	_	_	_	_	_	_	_
I-14	_	_	_	_	_	_	_	_	_	T	C	_	_	_	_	_	_	_	<u> </u>
I-15	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_
I-16	_	_	_	_	_	_	_	_	_	**	С	_	_	_	_	_	_	_	_
I-17	_	_	_	_	_	_	_	_	_	_	C	_	_	Т	_	_	_	_	A
I-18	_	_	_	_	_	_	_	_	_	**	С	_	_	_	_	_	_	_	_
I–19	_	_	_	_	_	_	_	_	Т	Т	_	_	_	_	_	_	_	_	_
I-20	_	_	_	_	Т	_	_	_	_	_	С	Т	_	_	Α	_	_	_	A
I-21	_	_	Α	_	_	_	_	_	_	Т	С	_	_	_	_	-	-	-	**
I-22	_	_	_	_	_	_	_	_	_	_	С	_	_	_	_	_	_	_	A
I-23	_	_	_	_	_	_	_	_	_	_	С	_	_	_	_	_	_	_	-
I-24	_	_	_	_	Т	_	_	_	_	_	С	Т	_	_	_	Α	_	_	_
I-25	_	Т	_	_	_	_	_	_	_	_	С	_	_	_	_	_	_	**	-
I-26	_	_	_	_	_	_	_	_	_	Т	С	_	_	_	_	_	_	_	_
I-27	_	_	_	_	_	_	_	Α	_	_	С	_	_	_	_	_	_	_	_
I-28	_	Т	_	Α	_	_	_	_	_	**	С	_	_	_	_	_	_	_	_
I-29	_	_	_	_	_	_	_	_	_	_	С	_	_	_	_	_	_	_	_
I-30	_	_	_	-	-	-	-	_	_	T	С	_	_	_	_	_	_	_	_
I-31	_	_	_	-	-	-	-	_	_	_	С	_	-	-	_	_	_	_	_
I-32	_	_	Т	_	_	_	_	_	Т	T	**	_	_	_	_	-	-	_	_
I-33	_	_	_	_	_	_	_	_	_	_	Т	_	_	_	_	_	_	_	_
I-34	_	_	_	_	_	_	Y	_	_	_	С	_	-	-	_	-	Т	-	-
I-35	_	_	_	_	_	_	_	_	_	_	С	_	-	_	_	_	Т	_	A
I-36	_	_	_	_	_	_	_	_	_	A	С	_	_	_	_	_	Т	_	A
I-37	_	Т	_	_	_	_	_	_	_	_	С	_	_	_	_	_	_	_	_
I-38	_	_	_	-	-	-	_	A	_	_	С	_	_	Т	_	-	-	-	_
I-39	_	_	_	_	_	_	_	_	_	Т	С	_	-	_	_	_	_	-	-
I-40	_	_	_	_	-	-	_	_	_	_	С	_	_	_	_	-	-	-	_
I-41	_	_	A	_	_	_	_	_	_	_	С	_	_	_	_	-	-	-	_
I-42	_	_	_	_	_	_	_	_	_	_	_	Т	A	Т	_	_	_	_	-
I-43	_	_	_	_	Т	_	Т	_	_	_	С	Т	A	_	_	A	-	-	-
I-44	_	_	_	_	_	_	Т	_	_	_	С	_	_	_	_	_	_	Α	-
I-45	_	_	_	_	_	_	_	_	Т	Т	С	_	_	_	_	_	_	_	_
I-46	_	_	_	-	-	-	-	_	_	_	С	_	_	-	_	_	_	_	Ī -
I-47	_	_	_	_	_	_	_	_	_	Т	С	_	_	_	_	_	_	_	_
				T. 1.	1. 1 6	2 1.									_				

^{*}Abbreviation of species see in Table 1, '-' – bars represents the identical nucleotides as that of *I. rugosus* in their respectively aligned sequence sites, **N – any nucleotide base.

Table 4. Distribution of informative (PI) sites and nucleotide base positions in the aligned 5.8S nrDNA and ITS-2 region of Isodon rugosus

G : 44	5.8S region (> 237400)		ITS-2 region (> 401607)											
Species*	237	418												
I-01	A	A	С	G	С	С	G	Т	С	С	Т	Т	С	
I-02	_	_	_	_	Т	_	-	_	_	gp	С	-	Т	
I-03	_	С	A	Т	_	_	-	_	_	-	С	_	Т	
I-04	_	С	A	Т	_	_	-	_	_	-	С	_	Т	
I-05	_	С	A	С	_	_	-	_	_	-	С	_	_	
I-06	_	С	A	**	_	_	-	_	_	-	С	_	_	
I-07	_	С	A	С	_	_	_	_	_	_	С	_	_	
I-08	G	-	_	_	_	_	-	_	_	_	-	_	T	
I-09	_	С	A	T	_	_	_	_	_	_	С	_	_	
I-10	_	С	A	T	_	**	_	_	_	_	С	_	T	
I-11	_	C	A	T	_	_	_	_	T	T	C	_	T	
I-12	_	С	A	T	_	_	_	_	_	_	С	_	T	
I-13	_	С	_	С	_	_	_	_	_	T	C	_	T	
I-14	_	С	A	Т	T	_	_	_	_	_	С	С	T	
I-15	_	С	A	Т	_	_	_	_	_	_	С	_	T	
I-16	_	С	A	Т	_	_	_	_	_	_	С	С	Т	
I-17	_	С		T	_	_	_	_	_		С	_	T	
I-18	_	С	A	Т			_	С	_		C		T	
I–19	_	С	A	T		_	_	_			С	_	T	
I-20	_	_	_	Т	_	A	_	_	_	_	С	_	T	
I-21	_	С	A	Т	_	_	_	_	_	_	С	_	T	
I-22	_	С	A	Т	_	_	_	_	_	_	C	_	T	
I-23	-	С	_	T	_	_	_	_	_	_	С	С	_	
I-24	-	_	A	_	T	_	_	_	T	_	C	_	Т	
I-25	-	С	A	T	_	_	_	_	_	_	С	_	T	
I-26	G	С	A	T	_	_	_	_	_	_	С	С	T	
I-27	-	С	A	Т	_	_	A	_	_	_	С	С	T	
I-28	_	С	A	Т	_	_	_	_	_	_	C	_	T	
I-29	_	С	A	Т		_	-	_		_	C	-	Т	
I-30	_	С	A	С	_	_	-	_	_	-	C	_	T	
I-31	_	С	A	С	**	_	-	_	_	_	С	C	T	
I-32	_	С	A	Т	_	_	-	_	_	-	C	**	T	
I-33	_	C	A	Т		_	-	_	_	-	C	С	T	
I-34	_	C	A		_	_	-	-	_	**	C	_	T	
I-35	_	C	A	T	_	_	-	С	_	T	C	_	T	
I-36	_	C	A	T		_	-	_	_	_	C	_	T	
I–37	_	С	A	Т	_	A	_	_	_	_	C	-	T	
I-38	_	С	A	Т	_	_	A	_	_		C	С	T	
I-39	_	С	A	A	_	_	_	_	_	T	C	_	T	
I-40	_	-	A	T	_	_	-	_	_		C	_	Т	
I-41	_	С	A	С	_ 	_	_	_		T	C	_	_	
I-42	_	_		_	T	_	-	_	T	_	C	_	T	
I-43	_	_		_	Т	_	_	_	Т	-	C	_	T	
I-44	_	-	gp	T		_	-	_		**	С	-	T	
I-45	_	С	A	Т	_	_	_	_	_	_	-	С	T	
I-46	_	-	_	_	Т	_	-	_	Т	_	C	-	T	
I–47	_	С	A	T	_	_	_	_	_	_	С	**	**	

^{*}Abbreviation of species see in Table 1, '-' – bars represent the identical nucleotides as that of *Isodon rugosus* in their respectively aligned sequence sites, ** N – any nucleotide base, gp – gap.

I-14, I-15, I-16, I-17, I-18, I-19, I-20, I-22, I-25, I-26, I-27, I-28, I-29, I-33, I-34, I-35, I-36, I-37, I-38, I-39, I-40, I-44, I-45 and I-46) were found complete in all the three regions of *nrDNA-ITS* (*ITS-1*, 5.8S and *ITS-2*). From the above 32 species, for *ITS-2* region, 19 species: *I. adenolomus* (I-3), *I. coetsa* (I-8), *I. enanderianus* (I-11), *I. eriocalyx* (I-12), *I. flavidus* (I-14), *I. flexicaulis* (I-15), *I. for-*

restii (I–16), *I. grandifolius* var. atuntzeensis (I–19), *I. japonicas* (I-22), *I. loxothyrsus* (I-25), *I. lungshengensis* (I–26), *I. megathyrsus* (I–27), *I. melissoides* (I–28), *I. phyllostachys* (I–33), *I. pleiophyllus* (I–34), *I. rosthornii* (I–35), *I. rubescens* (I–36), *I. rugosiformis* (I–37), *I. taliensis* (I–44) possessed similar spanning length of 208 nt (Table 2). Alignment in these taxa reflected a maximum length span of 212 nt

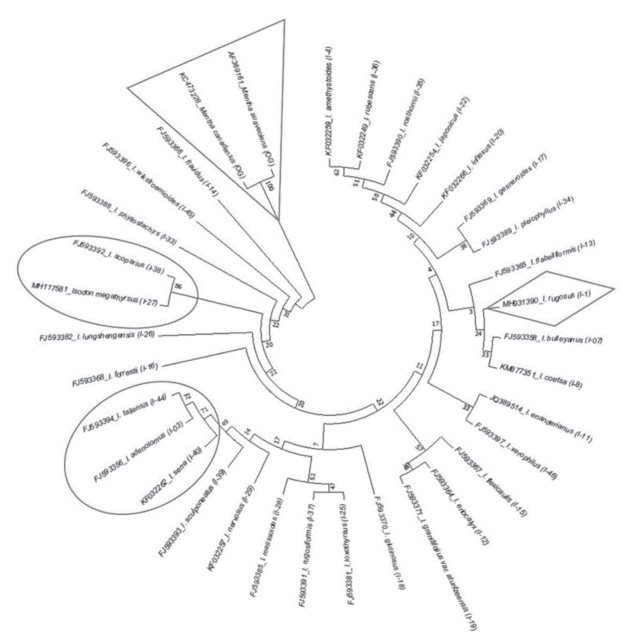


Fig. 1. Phylogenetic tree of *Isodon* species with two different out-group members (within a triangle), *Mentha suaveolens*-AF369161 and *M. canadensis*-KC473228, generated using *ITS* sequence according to the Maximum likelihood (ML) method. Numbers above lines are the bootstrap values in 1000 replicates, species within circles have bootstrap closeness above 70, query taxon *Isodon rugosus* within rhombus

in *I. flabelliformis* (I–13) and a minimum of 207 nt in four species, namely *I. rugosus* (I–01), *I. amethystoides* (I–04), *I. sculponeatus* (I–39) and *I. xerophilus* (I–46). Species *I. flabelliformis* (I–13) was reported with the lowest (66.03%) GC content. It was highest (68.72%) in *I. bulleyanus* (I–07), wherein 11 species, e.g. *I. adenolomus* (I–03), *I. flexicaulis* (I–15), *I. grandifolius var. atuntzeensis* (I–19), *I. loxothyrsus* (I–25), *I. lungshengensis* (I–26), *I. megathyrsus* (I–27), *I. melissoides* (I–28), *I. phyllostachys* (I–33), *I. rubescens* (I–36), *I. scoparius* (I–38) and *I. wikstroemioides* (I–45), were reported with same (67.79%) GC content.

The 32 species referred above with complete nrDNA-ITS regions (ITS-1 + 5.8S + ITS-2) were used to check the phylogenetic relationship of *I. rugosus*. Two different species of genus Mentha were used to confirm the out grouping, namely M. suaveolens (AF369161) and M. canadensis (KC473228). Nucleotide bases counts of nrDNA-ITS in Isodon species was observed with similar trends in their respective regions as C > G > A > T in ITS-1, G > C > A > T in 5.8S and C > G > T > A in ITS-2. The estimation of average evolutionary divergence in overall sequence pairs (i.e. overall mean distance in 34 sequences) was $d \pm S.E. = 0.023 \pm 0.002$ by the p-distance method and $d \pm S.E. = 0.024 \pm 0.002$ by both Jukes-Cantor's (Jukes & Cantor, 1969) and Kimura 2 parameters (KIMURA, 1980).

Phylogenetic tree generated by the maximum likelihood (ML) method (Fig. 1) placed the *I. rugo-sus* with a low bootstrap value near to the clade having species *I. coesta* (I–08) and *I. bulleyanus* (I–7). It indicated its divergence from the probable most recent common ancestor (*MRCA*) with missing linked members. The clades obtained above the bootstrap value 50 were informative. Two different clades (a clade having *I. scoparius – I. megathyrsus* and another having *I. adenolomus – I. taliensis*) show relatedness reasonably up to a good extend with their respective *MRCA* as in both clades bootstrap value was above 73.

The molecular signature of *ITS* marker in different medicinal plants has been studied by various researchers (ALICE & CAMPBELL, 1999; DOH et al., 2016; LIU et al., 2019; SRIVASTAVA & SAGGOO, 2014). In genus *Isodon*, 46 species were studied with *ITS* marker, and they are included in the present study

(Table 1). Similar to the above, species *I. rugosus* has been studied previously for *matK* marker by Shinwari et al. (2018) and Srivastava et al. (2020) and *psbA-trnH* intergenic spacers by Pekhna (2019).

Presently, taxon *I. rugosus* studied for its molecular (nrDNA-ITS) signature showed the presence of 32 informative (PI) sites together in the nrDNA-ITS regions (i.e. ITS-I + 5.8S + ITS-2 = 19 + 1 + 12). This information could be used as additional taxonomic knowledge for the species to confirm its presence in the local flora.

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MOLEKULINIS ISODON RUGOSUS (LAMIACEAE) nrDNA-ITS ŽYMENS PARAŠAS

Devendra Kumar SRIVASTAVA, Pekhna BANSAL, Pradeep Kumar SINGH, Manjit Inder Singh SAGGOO

Santrauka

Vaistinio augalo *Isodon rugosus* molekulinės nr-DNA-ITS sekos buvo įvertintos, naudojant universaliuosius ITS-1 ir ITS-2 pradmenis. Atlikus *in-silico* duomenų analizę buvo nustatyta, kad *Isodon* spp. 46 rūšių ITS sekos atitiko ITS-1, 5.8S ir ITS-2 regionus. Tačiau tik 32 rūšių sekos atitiko visus tris regionus, kitų rūšių sekos buvo dalinės. *Isodon rugosus* tyrimas atskleidė penkių genų regionus: 18S rRNR (da-

linė seka, > 1....38), ITS-1 (visa seka, > 39....236), 5.8S rRNR (visa seka, > 237....400), ITS-2 (visa seka, > 401....607) ir subgeno 26S rRNR (dalinė seka, > 608....672). Sekos regionai buvo nustatyti kintamose, singletoninėse ir rūšims būdingose informatyviose srityse. Informacija buvo pateikta filogenetiniame medyje, sudarytame naudojant maksimalios tikimybės metoda.