# PHYLOGENETIC ANALYSIS OF *ROSA DAMASCENA* L. FROM TAIF USING DNA BARCODING APPROACH

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#### Abstract

Little is known about the genome structure and the genetic origin of *Rosa damascene* L., widely distributed in Taif, Saudi Arabia for beauty and perfume production. For that, four specific barcodes; ITS2, mat*K*, rbcL and t-rnH were employed to genetically identify three *Rosa* species; *R. damascena*, *R. hybrida* and *R. damascena*Trigintipetala cultivated in Taif leading to the discrimination among them from one hand, and reviewing the previous suggestions of *R. damascena* origin from the other hand. rbcL, t-rnH and mat*K*revealed more effectiveness in sequence quality and species discrimination capability than ITS2 across the genome of *Rosa* depending on estimations of transition/transversion bias (R), Tajima relative evolutionary rate tests and the phylogenetic relationships using the Maximum Likelihood Tree method. *R. damascena* lineage exhibited accelerated evolutionary rates. The phylogenetic analyses proved the heterogeneous origin of *Rosa* cultivars from Taif doubting the validity of the previous suggestions about the genetic origin of *R. damascena*.

Key words: Rosa damascena, ITS2, matK, rbcL, t-rnH, barcoding genes.

#### Introduction

Roses nearly have 120 species and 20000 commercial cultivars worldwide that are popular for their beauty, medicinal purposes, decoration and perfume production (Rajapakse et al., 2001). Rose resources were categorized into native species, exotic species, exotic varieties and locally developed varieties, of these classified groups only 10-20 species have developed the modern cultivars of rose (Riaz et al., 2012). Despite their economic importance and for several reasons such as polyploidy, classical hybridization strategies, primitive selection, genetic mutations and insufficient funding for rose development projects, we have little information about the genome structure and subsequently the genetic origin of the three Rosa species; R. damascena, R. hybrida and R. damascena Trigintipetala cultivated in Taif. It was suggested that the summer Damask rose resulted from the recombination of Rosa phoenicia and Rosa gallica and Rosa moschata and R. gallica for the autumn Damask (Hurst, 1941), whereas Iwata et al., (2000) suggested that R. moschata, R. fedschenkoan and R. gallica were the parentsforthe old Damask roses. Based on the above, different molecular strategies for studying the genetic structure and variability in roses in different regions have been employed such as isozymes, RAPD, SSR, AFLP, ISSR, SCAR and the specific DNA barcoding genes (Baydar et al., 2004; Kiani et al., 2008; Jabbarzadeh et al., 2010; Schori & Showalter, 2011; Riaz et al., 2012; Ahmed et al., 2017).

The barcoding loci; ITS2, mat*K*, rbcL and t-rnH have been employed in the taxonomic and floristic studies as specific genes for plants. Their efficiency varied in quality and recovery and the discrimination ability among plant species (Tripathi *et al.*, 2013). Thus, many studies referred to the urgent need to use these loci

among and within plant species to get accurate data for their genomic structure and subsequently good identification leading to valid taxonomic relationships. Taifcity inSaudi Arabiais famous for the cultivation of some *Rosa* species especially *Rosa damascena* that is denominated "Ward Taifi", *R. hybrida* and *R. damascene* Trigintipetala for beauty and production of perfumes. There is scant available knowledge about the sequence data of gene regions of *Rosa* species cultivated in Taif, Therefore, the effectiveness of four barcodes; ITS2, mat*K*, rbcL and t-rnH, were investigated for improved identification of Taif*Rosa* species and to distinguish them from other *Rosa* species retrieved from the GenBank database confirming or denying the mentioned suggestions of *R. damascena* origin.

### **Materials and Methods**

**Plant materials:** The three *Rosa* species, *Rosa damascena* "Ward Taifi", *R. hybrida* and *R. damascene* Trigintipetala, utilized in the investigation were selected from Taif governorate, Saudi Arabia.

The extraction and amplification of DNA: The DNA of fresh leaves of Taif *Rosa* species was extracted using CTAB method (Doyle & Doyle, 1987). The purified DNA was used in PCR amplification of ITS2, rbcL, matK and trnH loci using universal primers as mentioned in Table 1. Each reaction included 1  $\mu$ l of each forward and reverse primer, 1  $\mu$ l genomic DNA, 12  $\mu$ l Master mix and 10  $\mu$ l of distilled water. After mixing the previous components, the reaction was set up in the PCR thermal-cycler. The first step was denaturation that was performed at 94°C for 5 min then followed by 35 cycles of another denaturation at 94°C for 1 min, annealing at 52°C for 30 sec and extension at 72°C for 2 min with a final extension step of 72°C for 8 min.

DNA locus	Primer name		Primer sequences (5'-3')	Ann. temp.	Reference
ITS2	ITS-S2F	F	ATGCGATACTTGGTGTGAAT	52°C	Chen et al., (2010)
11.52	ITS4	R	TCCTCCGCTTATTGATATGC	52 C	White et al., (1990)
rbcL	rbcLa	F	ATGTCACCACAAACAGAGACTAAAGC	52°C	Levin <i>et al.</i> , (2003) Kress & Erickson (2007)
IUCL	rbcLa	R	GTAAAATCAAGTCCACCRCG	52 C	Kress & Erickson (2007)
mat <i>K</i>	matK-KIM1	F	ACCCAGTCCATCTGGAAATCTTGGTTC	52°C	Fazekaset al., (2012)
man	matK-KIM3	R	CGTACAGTACTTTTGTGTTTACGAG	52 C	Fazekas <i>ei ul.</i> , (2012)
T-rnH	psbAF			52°C	Sang et al., (1997)
	t-rnH2			52 C	Sang <i>et al.</i> , (1997) Tate & Simpson (2003)

Table 1. List of primers used for the investigated DNA barcoding loci.

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Locus	ITS2	rbcL	mat <i>K</i>	T-rnH
Number of TaifRosa species examined	3	3	3	3
% PCR success	100	100	100	100
% Sequencing success	100	100	66.6	100
% Variable sites after alignment among Rosa species from Taif	15	39	40	25
Sequence length of <i>R. damascena</i>	241	558	825	360
Sequence length of <i>R. hybrida</i>	347	549	-	544
Sequence length of R. d. Trigintipetala	330	531	813	380
GC ratio in <i>R. damascena</i>	61	44	34	24
GC ratio in R. hybrida	59	44	-	32
GC ratio in R. d. Trigintipetala	59	44	38	26
Number of the retrieved Rosa species from the GenBank with lower E value	34	24	38	14
Mean query cover among R. damascena and the retrieved species	93	97	99	90
% Mean identity among R. damascena and the retrieved species	95	99	99.5	91

**The sequencing of PCR products:** The PCR products of Taif*Rosa* species for the four barcoding loci were sent for purification and sequencing process at Macrogen Inc., South Korea.

The sequences alignment and phylogenetic analysis: The sequences of ITS2, rbcL, matK and t-rnH of R. damascena were subjected to BLAST on http://blast.ncbi.nlm.nih.gov/Blast.cgi to confirm them from the other related Rosa species existed on database of the GenBank. Numbers of the retrieved Rosa species according to each barcode locus were shown in Table 2. MUSCLE algorithm was utilized for sequence alignments that were performed using the MEGA6 program (Edgar, 2004; Tamura et al., 2013). Sequence alignments were adjusted manually and gaps were eliminated in the phylogenetic analysis. The parameters of equality of evolutionary rate between sequences of R. damascena and R.hybrida, with sequence of R. damascena Trigintipetala used as an outgroup were estimated by Tajima's relative rate test (Tajima, 1993). Nucleotide substitution rates and Transition/Transversion bias (R) were calculated by Maximum Likelihood method using MEGA6 (Tamura et al., 2013). Means and levels of significance among values were determined by t-test using SPSS software (SPSS, 2011). The Maximum likelihood bootstrap (MLB) analysis were also performed for building the phylogenic trees. A total of 1000 bootstrap replicates were performed. Bootstrap support from maximum likelihood analysis  $\geq$ 

50% was detected on the branches of phylogenic trees and could be considered as good support.

#### **Results and Discussion**

DNA barcoding loci are modern important tools for developing the genetic relationships among plant species leading to rapid accurate approaches for crops improvement especially those possessing high economic value like roses. For that, ITS2, mat*K*, rbcL and t-rnH loci were investigated to identify and distinguish between *R. damascena* and two *Rosa* species from Taif, and its putative ancestors retrieved from the GenBank database.

Although the extraction of Rosa DNA was difficult, due to the large amounts of secondary products, the quantity and quality of DNA were sufficient for PCR amplification and sequencing. Statistics that derived from the amplification, sequencing and alignment processes of the four loci were shown in Table 2. The efficiency of both amplification (Fig. 1) and sequencing were 100% for the four candidate loci except sequencing of R. hybrida did not succeed for locus matK (66.6%). The sequences of R. damascena for ITS2, rbcL, matK and t-rnH were submitted to BLAST at the GenBank database for sequence symmetry and assertion of the amplified loci. Numbers of the retrieved Rosa species were 34, 24, 38 and 14 for ITS2, rbcL, matK and t-rnH respectively, that showed 95, 99, 99.5 and 91% similarity to sequences of *R. damascena* with lower E value ( $\leq 1e$ ).

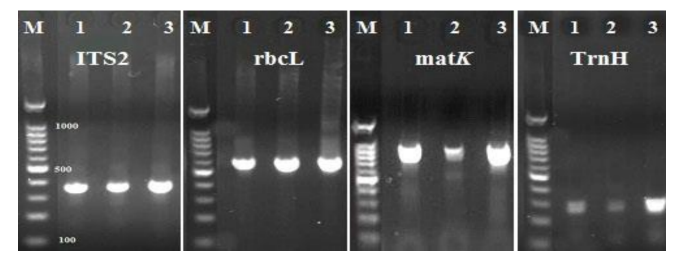


Fig. 1. The agarose gels showing PCR amplifications of ITS2, rbcL, matKand t-rnHloci of TaifRosa species; 1- R. damascena, 2- R. hybrida, 3- R. damascenaTrigintipetala. M- Marker.

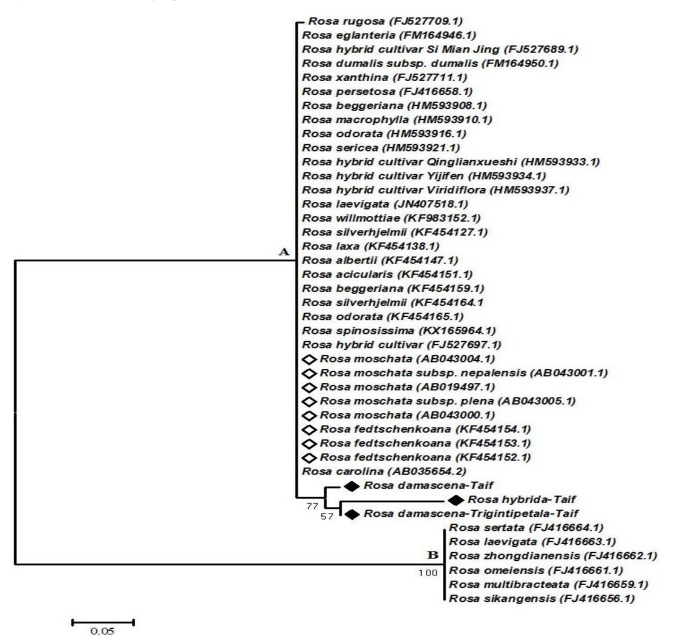


Fig. 2. Phylogram based on ITS2 locus for Taif roses and the retrieved species using Maximum Likelihood Tree method.

Current	1		Transition	ition					Transversion	rersion				é
opecies	TOCUS	A→G	G→A	T→C	C→T	A→T	T→A	A→C	C→A	T→G	G→T	C↑G	G→C	(K)
	ITS2	11.7	6.5	10.8	5.7	5.7	5.8	10.9	5.8	10.3	5.7	10.3	10.9	0.45
	rbcL	15.9	18.9	13.2	18.9	4.5	4.5	3.4	4.5	3.8	4.9	3.8	3.4	1.96
Overall	T-mH	6.5	16.0	7.3	29.0	7.9	7.6	2.0	7.6	3.1	7.9	3.1	2.0	0.99
	matK	9.4	16.7	10.2	21.1	7.5	6.6	3.6	6.6	3.7	7.5	3.7	3.6	1.17
	Mean	10.9*	14.5*	$10.4^{**}$	18.7*	6.4**	6.1**	5.0	6.1**	5.2	6.5**	5.2	5.0	1.14*
	ITS2	3.4	2.1	5.0	3.0	8.7	7.6	14.3	7.6	12.7	8.7	12.7	14.3	0.14
	rbcL	16.3	19.6	13.7	18.1	4.6	4.5	3.5	4.5	3.7	4.6	3.7	3.5	2.04
Taif cultivars	T-mH	7.8	22.1	9.3	21.9	6.8	7.3	2.9	7.3	2.6	6.8	2.6	2.9	1.07
	matK	,	•	,										•
	Mean	6.6	10.9	4.4	10.0	6.7*	6.5*	6.9	6.5*	6.3	6.7*	6.3	6.9	1.1
	ITS2	7.8	4.6	6.7	4.2	7.8	6.7	12.4	6.7	11.4	7.8	11.4	12.4	0.29
R. damascenaa and	rbcL			,									,	,
its supposed	T-mH	6.6	16.6	7.6	31.0	T.T	6.8	1.9	6.8	2.7	7.7	2.7	16.6	1.09
ancestors	matK	9.0	16.4	10.1	7.0	7.5	7.0	3.8	7.0	3.9	7.5	3.9	3.8	1.08
	Mean	7.8**	12.5	8.1*	14.1	7.7**	6.8**	6.0	6.8**	6.0	7.7**	6.0	10.9	0.82

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After alignment process, the sequence length range (from 544 to 360 bp) of t-rnH was higher than that of ITS2 (from 347 to 241 bp) and rbcL (from 558 to 531 bp), whereas, the percentages of variable sites were higher in matK (40%) and rbcL (39%) than those of ITS2 and t-rnH among Rosa species from Taif. The GC ratio scored in locus ITS2 was the highest and followed by those of rbcL, matKand t-mH loci as shown in Table 2. Knowledge of the rates of transitions to transversions for species sequences after alignment is important to explain the substitution mutations in plant genomes for studying the molecular evolution of these sequences of species under consideration (Kimura, 1981; Gojobori et al., 1982). Table 3 revealed that transitions occurred in general more than transversions. This is because the process of transition may be more easy than that of transversion supporting the point of view that nucleotide substitution bias prefer transitions more than transversions and some nucleotides mutate greater than others (Zhao & Boerwinkle, 2002; Zhang & Gerstein, 2003). It was also noticed that transition/transversion bias (R) of rbcL, t-rnH and matK was higher than that of ITS2. Luo et al., (2016) mentioned that the occurrence of transition-transversion bias differs based on genome region. Moreover, the Tajima relative evolutionary rate tests (Tajima, 1993) were estimated to investigate whether R. damascena and its relative species evolve at the same rate for the four specific loci (Table 4). The test showed that the evolutionary rates were not constant in R. damascena and its relative species. In spite of using different species as outgruops, R. damascena lineage exhibited accelerated evolutionary rates in all loci. Except in case of R. hybrida as an outgroup, R. damascena and R. damascena Trigintipetala showed a similar evolutionary rates because the P-values were higher than 0.05 in ITS2, rbcL and trnH loci. The null hypothesis of equal evolution rates between R. damascena and its supposed ancestors; R. moschata, R. gallica and R. fedschenkoana, was rejected because the P-values were lower than 0.05 in ITS2, rbcL and matKr evealing the accelerated evolutionary rate of R. damascena and subsequently reflecting the variance among them. The present results revealed that rbcL, t-rnH and matKwere more effective loci in sequence quality and also in species discrimination capability than ITS2 (Kress & Erickson, 2007; Piredda et al., 2011; De Vere et al., 2012) across the genome of Rosa.

For further clarification of the phylogenetic relationships of *Rosa* species under study, sequence data of the four loci was used to reconstruct the phylogenetic trees using the Maximum Likelihood Tree method (Figs 2, 3, 4 and 5). These loci have been used in many species for the development of the phylogenetic relationships due to their great discriminatory power (Dong *et al.*, 2014; Shinwari *et al.*, 2014; Al-Hemaid *et al.*, 2015; Bolson *et al.*, 2015; Tang *et al.*, 2016; Khan *et al.*, 2016; Zahra *et al.*, 2016; Shinwari *et al.*, 2018). Figure 2 demonstrated the phylogeny tree resulting from ITS2 data that split the *Rosa* species into two groups (A and B). Two subgroups were formed within group A showed a low level of diversity within the group. The

first subgroup had the putative ancestors; R. moschata and R. fedtschenkoana, of R. damascena within thirty three retrieved species from the GenBank, whereas, the three Rosa species from Taif were placed in the second subgroup showing low interspecific divergence among them. The phylogenetic tree derived from rbcL data categorized the three roses from Taif into single species, *R. hybrid* formed a separate operational taxonomic units (OTU) in cluster D and R. damascena and R. damascena Trigintipetala into two separate clades within cluster C (Fig. 3). The tree showed high divergence among the three species. Sequences of ancestors of R. damascena for rbcL gene were not represented in the GenBank database. t-rnH and matK trees showed complete divergence between R. damascena and its putative ancestors; R. moschata and R. gallica, into two distinct groups (Figs. 4 and 5). On the contraryt-rnH tree which grouped R. damascena and R. damascena Trigintipetala together in one group (F), the matK tree separated the two species into two different subgroups within group H. It could be concluded that, incontrastwith ITS2 tree, the phylogenetic trees depending upon rbcL, t-rnH and matK sequencing data revealed clear discrimination among the three Rosa species cultivated in Taif, on one hand, and between R. damascena and its supposed

ancestors on the other hand. These results confirmed those of Ahmed et al., (2017) depending on SSR and ISSR data. However, they were not in accordance with those of Iwata *et al.*, (2000) who suggested that *R*. moschata, R. gallicaand R. fedschenkoana were the parental species of Damask varieties depending on ITS and *psbA-t-rnH* sequences. This means that their suggestion may be uncertain and needs further studies or that R. damascenacultivated in Taif was originated from a hybridization process from other parents. The phylogenetic studies of the present Rosa cultivars are difficult and complex due to their heterogeneous origin resulting from free pollination in nature or the traditional strategies which permit the hybridization among cultivars or hybrids without declaring their parental form (Lidia & Irina, 2009). This confirmed the point of view that it is preferable to sequence more specific loci to obtain accurate results instead of one gene which is less reproducible to reconstruct the tree of evolution (Sandersson&Driskell, 2003). The obvious divergence of R. damascena from other species of Rosa may return to the geographic isolation, the traditional selection for the commercial traits and the adaptation to the environmental conditions in Taif through some mutational events.

T	Outgroup Testing group			ы	DD	DA	DD		Devalue
Locus	Outgroup	(A)	<b>(B)</b>	RI	RD	RA	RB	χ2	P value
	R. d. Trigintipetala	R. damascena	R. hybrida	201	3	7	19	5.54	< 0.05
ITCO	R. hybrida	R. damascena	R. d. Trigintipetala	201	3	7	7	0.00	>0.05
ITS2	R. fedtschenkoana	R. damascena	R. moschata	209	0	8	0	8.00	< 0.05
	R. moschata	R. damascena	R. fedtschenkoana	209	0	8	1	5.44	< 0.05
	R. d. Trigintipetala	R. damascena	R. hybrida	249	0	1	200	197	< 0.05
rbcL	R. hybrida	R. damascena	R. d. Trigintipetala	249	0	1	2	0.33	>0.05
IDCL	R.gallica	R. damascena	R. moschata	154	2	115	0	115	< 0.05
	R. moschata	R. damascena	R. gallica	154	2	115	1	112	< 0.05
T-rnH	R. d. Trigintipetala	R. damascena	R. hybrida	138	4	4	95	83.7	< 0.05
	R. hybrida	R. damascena	R. d. Trigintipetala	138	4	4	2	0.67	>0.05
maat V	R. gallica	R. damascena	R. moschata	417	0	276	0	276	< 0.05
mat <i>K</i>	R. moschata	R. damascena	R. gallica	417	0	276	1	273	< 0.05

Table 4. Tajima relative rate tests of the four loci for *Rosa* species.

The Tajima relative rate test was used to examine the equality of evolutionary rate between *R.damascena* and other relative species with different outgorups

RI is the identical sites in all three sequences

RD is the divergent sites in all three sequences

RA is the number of unique differences in the sequence A

RB is the number of unique differences in the sequence B

 $\chi^2$  test statistic more than 3.841 (p<0.05) indicates accelerated evolution

P value greater than 0.05 is often used to accept the null hypothesis of equal rates between lineages

#### Conclusion

This study is a pioneering attempt at producing DNA sequences from ITS2, matK, rbcL and t-rnH barcoding loci in three Rosa species cultivated in Taif. matK, rbcL and t-rnH were more applicable barcodes than ITS2 for the authentication and diversification of Rosa species. Like animal genomes, the plant genomes showed higher transition rates than the transversion rates. Mutation accumulation, little gene flow and geographic isolation encouraged R. damascenato gain accelerated evolutionary rates and subsequently considerable divergence from other related species to genus Rosa.

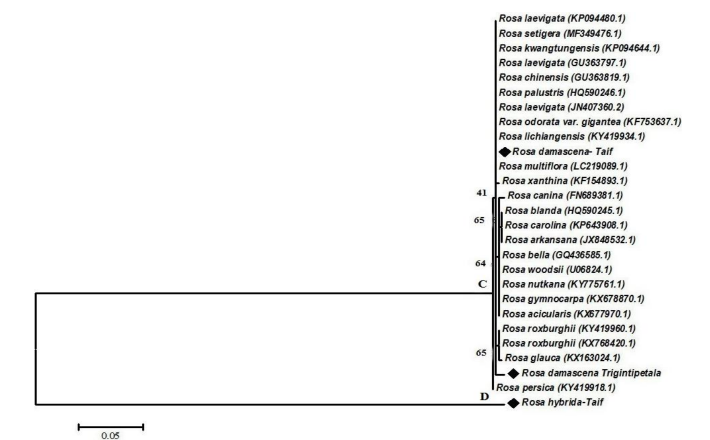


Fig. 3. Phylogram based on rbcL locus for Taif roses and the retrieved species using Maximum Likelihood Tree method.

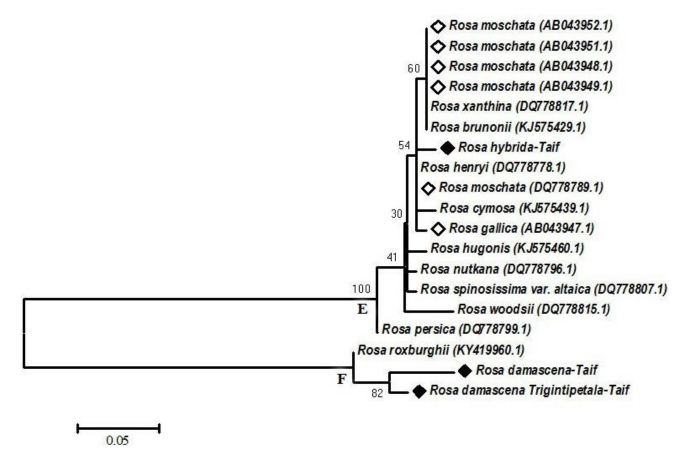


Fig. 4. Phylogram based on t-rnH locus for Taif roses and the retrieved species using Maximum Likelihood Tree method.

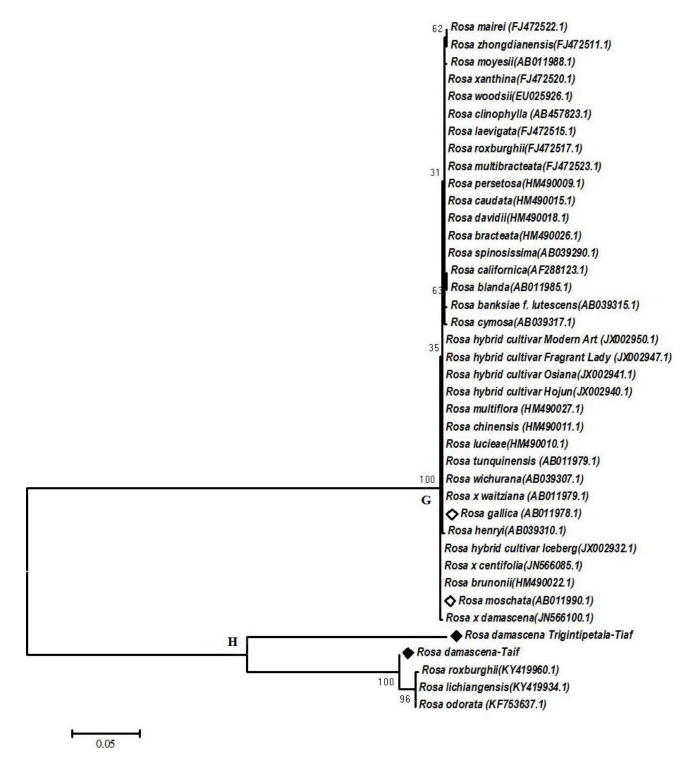


Fig. 5. Phylogram based on matK locus for Taif roses and the retrieved species using Maximum Likelihood Tree method.

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