

# Development of SSR Markers from Transcriptomes for Orinus (Poaceae), an Endemic of the Qinghai–Tibetan Plateau

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Source: Applications in Plant Sciences, 5(7)

Published By: Botanical Society of America

URL: https://doi.org/10.3732/apps.1700029

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PRIMER NOTE

## DEVELOPMENT OF SSR MARKERS FROM TRANSCRIPTOMES FOR ORINUS (POACEAE), AN ENDEMIC OF THE QINGHAI–TIBETAN PLATEAU<sup>1</sup>

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- *Premise of the study:* Transcriptomes were used to develop microsatellite markers for the plant genus *Orinus* (Poaceae), which comprises three species of grasses (*O. thoroldii*, *O. kokonoricus*, and *O. intermedius*) that are widely distributed in the Qinghai–Tibetan Plateau.
- *Methods and Results:* Primer pairs were developed for 16 high-quality simple sequence repeats (SSRs) using transcriptomes. SSRs were amplified in 248 individuals representing the three species of *Orinus*; the number of alleles per locus ranged from one to seven, with an average of 2.6. The expected and observed heterozygosity per locus varied from 0.00 to 0.83 and from 0.00 to 1.00, respectively, with respective mean values of 0.32 and 0.34.
- Conclusions: These newly developed SSR markers will be valuable for evaluating the population genetic structure of Orinus throughout its range.

Key words: next-generation sequencing; Orinus; orthologous gene; Poaceae; simple sequence repeat (SSR) marker.

Simple sequence repeats (SSRs) have been widely used for DNA fingerprinting, molecular-assisted breeding, detecting gene locations, genetic diversity analyses, and evolutionary studies because they are codominant, are highly polymorphic, can be amplified repeatably, and provide many informative sites distributed throughout the genome (Agarwal et al., 2008; Izzah et al., 2014). SSRs that are developed from transcribed RNA sequences, known as expressed sequence tag SSRs (EST-SSRs), can be developed cheaply and efficiently using next-generation sequencing technology (Simon et al., 2009). Previously, microsatellite markers developed from transcriptomes have primarily been for woody and medicinal plants (e.g., Liu et al., 2014;

<sup>1</sup>Manuscript received 28 March 2017; revision accepted 9 May 2017. The authors thank Prof. Jianquan Liu for providing laboratory space and funding and thank the reviewers for their comments on the manuscript. This study was supported by the National Natural Science Foundation of China (no. 31260052), the Natural Science Foundation of Qinghai Province (no. 2016-ZJ-937Q, 2014-ZJ-947Q), the State Scholarship Fund of China Scholarship Council (no. 201508635003), and the Young and Middle-Aged Research Foundation of Qinghai Normal University (no. 2015-19).

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doi:10.3732/apps.1700029

Kim et al., 2017). Few microsatellite markers have been developed to study endemic alpine grasses.

Here, we show the utility of SSR markers derived from transcriptome assemblies to detect genomic variation in the three species of Orinus Hitchc. (Poaceae), which are endemic grasses that occur at high elevations in xeric, alpine areas of the Qinghai-Tibetan Plateau (QTP) (Su et al., 2015). Orinus comprises O. thoroldii (Stapf ex Hemsl.) Bor, which is mainly distributed in the western QTP, O. kokonoricus (K. S. Hao) Tzvelev, which occurs in the eastern QTP, and O. intermedius X. Su & J. Quan Liu, which is a hybrid of O. thoroldii and O. kokonoricus and has a range in the southeastern QTP (Su et al., 2015, 2017). Orinus has ecological and conservation value for its role in soil stabilization in the QTP, especially due to its expansive root system (Su et al., 2013), and this genus may represent a good system for elucidating the timing and mechanisms of desertification in the OTP (Su et al., 2015). Therefore, developing SSR markers for Orinus is a first step toward utilizing the population history of its species to better understand the origins of the genus and to improve conservation efforts for desert habitats in the QTP. Here, we develop microsatellite markers in Orinus using transcriptomes obtained via Illumina paired-end sequencing.

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Locus	Primer sequences $(5'-3')$	Repeat motif	Allele size range (bp)	$T_{\rm a}$ (°C)	GenBank accession no.	GenBank accession of best BLAST hit	Organism of best BLAST hit	<i>E</i> -value
Oril	F: AGAAGATCATTGCTTTCAACC R. TTTTCCATCGCCACTACT	(CT) <sub>9</sub>	136–171	59	KY852238	KP856157.1	Quercus variabilis	$8.0 \times 10^{-7}$
Ori3	-	(GGA) <sub>9</sub>	160–178	60	KY705074	NM_001150415.1	Zea mays	$3.0 \times 10^{-12}$
Ori4	K: AAICGACIGCIGIAIGCAAAGG F: AAGAAACAAAAAGIGGIGAGG b. maaccommmemeencocomma	$(TGA)_9$	278–305	58	KY705075	EU976105.1	Zea mays	$5.0 imes10^{-28}$
Ori5		$(GAA)_7$	235–250	59	KY705076	NM_001151471.1	Zea mays	$4.0 \times 10^{-36}$
Ori12		$(TAA)_{12}$	243–267	60	KY705077	XM_004975213.3	Setaria italica	$1.0 \times 10^{-29}$
Ori13		(CCG) <sub>6</sub>	120–161	60	KY852239	XM_004973362.2	Setaria italica	$1.0 \times 10^{-10}$
Ori14		$(AT)_{10}$	155-204	58	KY852240	CR382128.1	Yarrowia lipolytica	$5.0 imes10^{-3}$
Ori15		$(AAT)_{10}$	204-228	59	KY705078	AC116411.7	Mus musculus	$3.0 \times 10^{-12}$
Ori17	• -	(TCC) <sub>12</sub>	297–321	09	KY705079	NM_001175683.1	Zea mays	$3.0 \times 10^{-43}$
Ori21		(CTC) <sub>12</sub>	249–270	60	KY705080	AC104200.12	Mus musculus	$1.0 \times 10^{-10}$
Ori31		$(TCT)_7$	182–221	69	KY852241	XM_004967825.1	Setaria italica	$1.0 \times 10^{-81}$
Ori32	K: CTCGAGGAGGAAGAGGACGA F: AGCAAGCATACCTAATGTTTTG R· CACCACGTTTCATATGGG	(TG) <sub>13</sub>	294–324	59	KY705081	XM_004975367.2	Setaria italica	$2.0 \times 10^{-21}$
Ori33		(TCC) <sub>6</sub>	184–222	60	KY852242	XM_004982769.1	Setaria italica	$8.0 \times 10^{-83}$
Ori36		$(AG)_{10}$	205–215	59	KY705082	CP018161.1	Oryza sativa	$2.0 \times 10^{-14}$
Ori38	-	$(CT)_{21}$	154-192	59	KY705083	AY486591.1	Hevea brasiliensis	$2.0 \times 10^{-15}$
Ori40		$(TC)_7$	141–188	60	KY852243	KF785779.1	Nandina domestica	$5.0  imes 10^{-3}$

TABLE 1. Characteristics of 16 polymorphic SSR markers developed in three species of Orinus.

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*Note*:  $T_a = optimal$  annealing temperature.

#### METHODS AND RESULTS

Twenty-five to 30 individuals were collected from three populations each of the three *Orinus* species (248 total collections) (Appendix 1). In addition, representative individuals of *O. thoroldii* and *O. kokonoricus* were collected (Appendix 1), from which fresh leaves were obtained; these were immediately frozen in liquid nitrogen in the field and later stored at  $-80^{\circ}$ C prior to RNA extraction. For all collections, voucher specimens were deposited at the Herbarium of the Northwest Plateau Institute of Biology (HNWP) (Appendix 1).

Total RNA and genomic DNA were extracted using a cetyltrimethylammonium bromide (CTAB) procedure (Ghangal et al., 2009). The RNA was quantified and its quality assessed using NanoDrop 2000 (Thermo Fisher Scientific, Waltham, Massachusetts, USA) and the Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, California, USA). The total RNA samples were purified to remove poly(A) tags using approximately 5  $\mu$ g of RNAs and (dT)-conjugated beads (Life Technologies, Carlsbad, California, USA), and purified RNAs were divided into 200-bp fragments using divalent cations at 75°C. The first strand of cDNA was synthesized with reverse transcriptase and random hexamer primers, and the second strand was synthesized by RNase H (Invitrogen, Ghent, Belgium) and *Taq* DNA polymerase I (New England BioLabs, Ipswich, Massachusetts, USA). Finally, the cDNAs representing the transcriptome were sequenced on an Illumina (Solexa) Genome Analyzer II (Illumina, San Diego, California, USA). All sequence information has been deposited in the National Center for Biotechnology Information (NCBI) Sequence Read Archive (Bioproject no. PRJNA385721).

SSR primers were designed using the transcriptomes of O. thoroldii and O. kokonoricus. To accomplish this, 79,111,257 and 67,617,602 raw reads were first filtered from O. thoroldii and O. kokonoricus. As a result, 60,124,556 and 50,037,026 high-quality reads, respectively, were obtained for de novo assembly. De novo assembly was performed in Trinity 2.2 with default parameters (Grabherr et al., 2011). From Trinity, 23,029 high-quality contigs were obtained from O. thoroldii and 24,086 from O. kokonoricus, representing total lengths of 20,808,832 and 22,281,570 bp, respectively, with an average size of 903 bp (N50 = 1188 bp) and 925 bp (N50 = 1203 bp). The assembly of each species was used to map the reads of the other using Bowtie 2 (Langmead and Salzberg, 2012) to identify an orthologous set of genes. Within the orthologous set, we searched for candidate SSRs using MISA 4.0 (http://pgrc.ipk-gatersleben.de/misa) and also identified single-nucleotide polymorphisms (SNPs) and insertion/deletion polymorphisms (indels) using SAMtools 1.4 (Li et al., 2009). Indels were selected as SSR candidates only if the indels appeared informative between the two species and could be confidently aligned at their 3' and 5' ends. In total, 58 polymorphic candidate loci were recovered from O. thoroldii and 52 from O. kokonoricus. For each locus, primers were designed in Primer3 (Rozen and Skaletsky, 2000), and primers with binding sites containing SNPs were rejected. Thus, a total of 50 primers suitable for both species of Orinus were found. Ten individuals were selected from each of three species of Orinus to test the 50 primers. Amplification was performed using a standard 25-µL PCR reaction containing 1.00 µL of template DNA, 0.20 mM MgCl<sub>2</sub>, 0.25 mM dNTPs, 2.00 µM of each primer, 2.50  $\mu$ L of 10× PCR buffer, 0.25  $\mu$ L of *Taq* DNA polymerase (5 U· $\mu$ L<sup>-1</sup>; TaKaRa Biotechnology Co., Dalian, Liaoning, China), and distilled water up to the final volume. The amplification was carried out under the following thermocycling protocol: enzyme initiation at 94°C for 5 min; followed by 36 cycles of denaturation at 94°C for 50 s, annealing at 49–58°C for 50 s (Table 1), and 72°C for 1 min; and a final extension at 72°C for 10 min. Amplification success was determined by viewing the results in a 1.5% (w/v) agarose gel. Sixteen primers showed successful amplification in all three Orinus species, and all of them possessed clear polymorphisms (Table 1).

Subsequently, the 16 primers were used to carry out fluorescence-based genotyping for all 248 sampled individuals of Orinus. Fluorescence-based genotyping was performed using a modification of the method presented in Hayden et al. (2008). In brief, the forward primers for the 16 SSRs were labeled with 6-FAM fluorescent tags (Applied Biosystems, Foster City, California, USA) and PCR reactions were performed as described above. The labeled products were detected on an ABI 3730XL sequencer with the GeneScan 500 LIZ Size Standard (Applied Biosystems). The profiles of the amplified loci were examined using GeneMapper 3.7 (Applied Biosystems), and peaks were scored manually by visual inspection. Each SSR marker was characterized by calculating three measures of genetic diversity in GENEPOP 4.2 (Rousset, 2008): number of alleles per locus, observed heterozygosity, and expected heterozygosity (Table 2). By these measures, the markers were highly polymorphic, with the number of alleles ranging from one to seven within this genus, with an average of 2.6 alleles per locus. The expected and observed heterozygosity ranged from 0.00 to 0.83 and 0.00 to 1.00, respectively, with respective mean values of 0.32 and 0.34 (Table 2).

					O. thoroldii	oldii							0.	O. kokonoricus	ricus								0. intermedius	nedius			
	Ŭ	Ge'er $(N = 30)$	= 30)	R	Renbu $(N = 28)$	= 28)	Z	Zhongba (N =	V=28)	Gol	Gonghe $(N = 27)$	= 27)	Nan	Nangqian $(N = 25)$	<i>l</i> = 25)	Bia	Bianba $(N = 26)$	= 26)		Aba $(N =$	30)	Rai	Rangtang $(N = 26)$	V = 26)	Ŵ	Mangkang $(N = 28)$	N = 28)
Locus	A	$H_{\rm o}$	$H_{\rm e}^{\rm b}$	A	$H_{ m o}$	$H_{\rm e}^{\rm b}$	A	$H_{ m o}$	$H_{\rm e}^{\rm b}$	A	$H_{\rm o}$	$H_{\rm e}^{\rm b}$	A	$H_{ m o}$	$H_{\rm e}^{\rm b}$	A	$H_{ m o}$	$H_{\rm e}^{\rm b}$	A	$H_{ m o}$	$H_{\rm e}^{\rm b}$	A	$H_{ m o}$	$H_{\rm e}^{\rm b}$	A	$H_{ m o}$	$H_{ m e}^{ m b}$
Ori1	_	0.000	0.000	З	0.600	0.540	9	0.533	0.584	4	0.571	0.498	-	0.000	0.000	ŝ	0.200	0.416	2	0.100	0.089	-	0.000	0.000	0	1.000	0.500
Ori3	2	0.143	0.133	0	0.111	0.278	0	0.400	0.320	1	0.000	0.000	2	0.125	0.117	2	0.250	0.219	-	0.000	0.000	0	0.500	0.375	0	0.143	0.133
Ori4	2	0.333	0.278	0	0.222	0.346	0	0.600	0.420	0	0.429	0.337	1	0.000	0.000	б	1.000	0.580	0	0.500	0.375	0	0.500	0.375	С	0.143	$0.255^{**}$
Ori5	З	0.667	0.486	4	0.778	0.543	4	0.600	0.665	0	0.143	0.133	-	0.000	0.000	0	0.400	0.320	б	0.333	0.569	-	0.000	0.000	0	0.143	0.133
Ori12	З	0.333	0.292	0	0.111	0.105	-	0.000	$0.320^{**}$	0	0.286	0.490	1	0.000	0.000	ŝ	0.800	0.660	0	0.500	0.375	С	0.250	0.219	0	0.143	0.133
Ori13	0	0.400	0.320	0	0.111	0.105	-	0.000	0.000	0	0.200	0.480	1	0.000	$0.564^{*}$	ŝ	0.467	0.520	0	0.400	0.320	-	0.000	0.000	З	0.500	0.457
Ori14	0	1.000	0.500	0	0.000	0.444	-	0.000	0.000	0	0.100	0.092	б	0.333	0.287	0	0.200	0.320	-	0.000	0.000	0	0.067	0.180	9	0.067	0.447
Ori15	-	0.000	0.000	0	0.222	0.198	0	0.100	0.095	1	0.000	0.000	1	0.000	0.320	0	0.200	0.180	0	0.500	0.375	С	0.000	0.625*	9	0.857	0.776
Ori17	Э	0.333	0.569	0	0.111	0.105	0	0.300	0.455	-	0.000	0.000	0	0.000	0.124	ŝ	0.800	0.540	-	0.000	0.000	С	0.250	0.219	0	0.286	0.245
Ori21	0	0.500	0.486	4	0.667	0.623	0	0.400	0.320	0	0.429	0.337	4	0.625	0.578	0	1.000	0.500*	0	1.000	0.500	4	0.750	0.656	4	0.286	0.622
Ori31	З	0.500	0.467	4	0.300	0.610	0	0.100	0.095	4	0.571	0.497	9	0.067	0.447	0	0.100	0.095	Э	0.500	0.580	ŝ	0.500	0.395	С	0.500	0.457
Ori32	0	0.500	0.375	С	0.111	0.105	-	0.000	0.000	0	0.286	0.490	2	0.750	0.828*	ŝ	0.600	0.580	-	0.000	0.000	С	0.067	0.064	0	0.000	0.245**
Ori33	0	0.100	0.095	4	0.500	0.665	-	0.000	0.000	0	0.200	0.480	ŝ	0.643	0.676	2	0.267	0.320	-	0.000	0.000	0	0.000	0.124	З	0.067	0.127
Ori36	2	0.667	0.444	0	0.111	0.105	0	0.100	0.095	0	0.143	0.133	2	0.500	0.375	-	0.000	0.444	-	0.000	0.000	0	0.000	0.375*	2	0.000	$0.245^{**}$
Ori38	ŝ	0.833	0.667	9	0.667	$0.784^{*}$	ŝ	0.700	0.775	ŝ	0.714	0.622	5	0.500	0.500	2	0.800	0.480	Э	1.000	0.625	4	0.750	0.656	9	1.000	0.776
Ori40	1	0.000	0.000	4	0.571	0.497	0	0.400	0.320	4	0.267	0.333	1	0.000	0.000	0	0.111	0.105	-	0.000	0.000	0	0.067	0.064	4	0.200	0.218
Note:	= A	numbei	r of alle	les:	$H_{a} = \exp \left( \frac{1}{2} + \frac$	<i>Note:</i> $A =$ number of alleles: $H_{\circ} =$ expected heterozy gosity: $H_{\circ}$	teroz	v gosity:	$H_{c} = obs$	erved	= observed heterozygosity	vgosity.															
<sup>a</sup> Loc:	dity :	and vol	Icher in	form	nation fo	<sup>a</sup> Locality and voucher information for the populations are prov	Julati	ons are	provided 1	n Api	rided in Appendix 1																
<sup>b</sup> Sign	ifica	int devia	ation fro	m F.	Iardy-W	<sup>b</sup> Significant deviation from Hardy-Weinberg equilibrium after	equil	librium ɛ		ction	for mult	tiple test	s (*I	<sup>9</sup> < 0.05,	correction for multiple tests (* $P < 0.05$ , ** $P < 0.01$ )	.01).											

TABLE 2. Genetic diversity statistics for each sampled population of the three Orinus species based on 16 pairs of SSR primers.<sup>a</sup>

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#### CONCLUSIONS

We have developed 16 polymorphic SSR markers from two cDNA libraries for investigating population structure in *Orinus*. These markers amplified easily and showed considerable polymorphisms for 248 individuals from three populations each of the three species of *Orinus*. These markers represent valuable new tools that will facilitate the development of *Orinus* as a model for understanding the origins and phylogeographic processes of the alpine desert of the QTP.

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APPENDIX 1. Locality information for the populations of Orinus used in this study.

Species	Ν	Population	Geographic coordinates	Altitude (m)	Voucher no. <sup>a</sup>
O. thoroldii	30	Ge'er, Xizang, China	31°36'13.7"N, 80°22'15.6"E	4570	X. Su 2011055
	28	Renbu, Xizang, China	29°18'0.4"N, 89°46'7.3"E	3767	X. Su 2011018
	28	Zhongba, Xizang, China	29°41'7.9"N, 84°8'48.1"E	4563	X. Su 2011044
	1	Gongga, Xizang, China	29°0'27.0"N, 85°26'48.8"E	4687	X. Su 2011078 <sup>b</sup>
O. kokonoricus	27	Gonghe, Qinghai, China	36°11'3.0"N, 100°59'16.9"E	2826	X. Su 2012040
	25	Nangqian, Qinghai, China	32°32'50.6"N, 96°11'45.2"E	4119	X. Su 2011074
	26	Bianba, Xizang, China	30°58'40.3"N, 94°43'35.3"E	3597	X. Su 2013083
	1	Gonghe, Qinghai, China	36°21'26.3"N, 100°43'5.8"E	3130	X. Su 2013008 <sup>b</sup>
O. intermedius	30	Aba, Sichuan, China	32°45′26.7″N, 102°33′3.8″E	3319	X. Su 2012003
	26	Rangtang, Sichuan, China	31°46'16.2"N, 100°58'57.1"E	3478	X. Su 2012007
	28	Mangkang, Xizang, China	29°32′27.2″N, 98°15′3.3″E	3507	X. Su 2012016

*Note*: *N* = number of individuals sampled.

<sup>a</sup>All voucher specimens were deposited at the Herbarium of the Northwest Plateau Institute of Biology (HNWP), Chinese Academy of Sciences, Xining, Qinghai Province, China.

<sup>b</sup>These representative individuals of Orinus thoroldii and O. kokonoricus were only used for RNA extraction.