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PROTOCOL NOTE

A LONG PCR-based approach for DNA enrichment prior to next-generation sequencing for systematic studies¹

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- Premise of the study: We present an alternative approach for molecular systematic studies that combines long PCR and next-generation sequencing. Our approach can be used to generate templates from any DNA source for next-generation sequencing. Here we test our approach by amplifying complete chloroplast genomes, and we present a set of 58 potentially universal primers for angiosperms to do so. Additionally, this approach is likely to be particularly useful for nuclear and mitochondrial regions.
- Methods and Results: Chloroplast genomes of 30 species across angiosperms were amplified to test our approach. Amplification success varied depending on whether PCR conditions were optimized for a given taxon. To further test our approach, some amplicons were sequenced on an Illumina HiSeq 2000.
- *Conclusions:* Although here we tested this approach by sequencing plastomes, long PCR amplicons could be generated using DNA from any genome, expanding the possibilities of this approach for molecular systematic studies.

Key words: angiosperms; chloroplast enrichment; long PCR; next-generation sequencing; plastome; universal chloroplast PCR primers.

Advancements in next-generation sequencing (NGS) technologies have permitted the assembly of large, genome-scale data sets that have shed light on the evolutionary history of many taxa (e.g., Parks et al., 2009; Moore et al., 2010; Xi et al., 2012; Eaton and Ree, 2013; Tennessen et al., 2013). For plant phylogenetics, there has been a major focus on methods for chloroplast phylogenomics (e.g., Parks et al., 2009; Moore et al., 2010), although methods for collecting phylogenomic data sets from the nuclear and mitochondrial genomes have also been developed (e.g., Straub et al., 2012; Eaton and Ree, 2013). Stull et al. (2013) developed a custom RNA probe set designed to capture angiosperm plastomes via solution-based hybridization. While their capture system was broadly successful, Stull et al. (2013) found that the most variable spacer regions were often captured at much-reduced coverage compared to more conserved regions, and were sometimes missed entirely if the target taxon was phylogenetically divergent from one of the 22 plastomes used in the bait design. Moreover, the current cost of the

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capture probes makes this method most efficient for projects dealing with hundreds of species. Another commonly employed method for plant phylogenomic studies is genome skimming (Straub et al., 2012), which takes advantage of the fact that organellar DNA and nuclear ribosomal DNA are present at high copy numbers in genomic DNA. However, a significant limitation of this method for systematic studies is that only high-copy number regions are recovered consistently across all samples, whereas regions with lower representation are only recovered in some samples and missed completely in others (Straub et al., 2011). This can be problematic for molecular systematic studies where missing data may result in misleading phylogenetic results (Lemmon et al., 2009). Moreover, being limited to highcopy regions in the genome becomes restrictive for experimental design as it excludes putatively highly informative regions in the genome such as single-copy nuclear genes (e.g., the single-copy orthologous genes [COSII] and the pentatricopeptide repeat [PPR] gene family; Wu et al., 2006, and Yuan et al., 2009, respectively).

As an alternative, we present an NGS approach that combines long PCR and Illumina sequencing to strategically compile phylogenomic data sets for molecular systematic studies. Long PCR, or long-range PCR, uses a combination of two polymerases—a nonproofreading polymerase at high concentration and a proofreading polymerase at a lower concentration—to amplify DNA fragments that range between 3 and 15 kilobases (kb), although cases of extremely large fragments (22–42 kb) have been reported (e.g., Cheng et al., 1994). Long PCR has been used extensively in human genome projects (e.g., Craig et al., 2008) and to sequence complete mitochondrial genomes (e.g., Knaus et al., 2011; Alexander et al., 2013), using both

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TABLE 1. List of :	species included in this	study, with vo	oucher inf	formation, t	issue sourc	es, and	NGS assemb	ly statisti	cs whe	en available. ^a							
Species	Order/Family	Collection no.	Herbarium	Type of tissue	Collection date	No. of amplified regions	Region no. 1 not 1 amplified ^b s	Base pairs equenced ^e e	No. of contigs	CAL bp a (min-max)	Ave. ssembly depth	No. of masked bp ^d	% of masked bp	N50	% called ar bases ^e	No. of nbiguous bases	% of ambiguous bases
Bartsia inaequalis	Lamiales/Orobanchaceae	Uribe-Convers	D	Silica	5 July 2010	16	n/a	125,283	25	5011	656	2126	1.7	19,294	99.9729	34	0.02714
Benth. Castilleja covilleana	Lamiales/Orobanchaceae	2010-22 Tank 1046	Ð	gel-dried Silica	13 July 2009	16	n/a	133,595	10	(204–28,257) 13,360	641	101	0.08	37,107	99.9948	7	0.00524
L. F. Hend. Castilleja elmeri	Lamiales/Orobanchaceae	Olmstead	WTU	gel-dried Silica	4 July 2001	16	n/a	122,614	Ξ	(1222–48,767) 11,147 (464–24,603)	664	440	0.36	33,049	92.9976	3	0.00245
remala Castilleja linariifolia Banth	Lamiales/Orobanchaceae	2001-78 Tank 2001-49	WTU	gel-dried Silica	21 July 2001	16	n/a	122,046	~	(404-34,0U2) 15,256 (810 50.680)	642	260	0.21	28,529	99.9984	7	0.00164
Castilleja miniata Douoloo ov Hoole	Lamiales/Orobanchaceae	Tank 1048-b	D	Silica	13 July 2009	16	n/a	134,704	4	(000,00-210) 33,676 (6157 75 123)	844	35	0.03	75,123	0266.66	4	0.00297
Douglas ex Hook. Castilleja pallescens	Lamiales/Orobanchaceae	Tank 2009-8	Ð	gel-dried Silica	6 June 2009	16	n/a	125,490	4	(621,67-7610) 31,372 7002 27 0002	764	29	0.02	73,629	99.9984	5	0.00159
(A. Uray) Ureenin. Bartsia stricta (Kunth)	Lamiales/Orobanchaceae	Uribe-Convers	Ð	gel-dried Silica	7 July 2010	15	13	119,828	14	(2029-12,029) 3559 (175 27 105)	707	1045	0.87	67,195	7966.66	4	0.00334
Denui. Castilleja applegatei Femold	Lamiales/Orobanchaceae	z010-24 Tank 2001-35	WTU	Silica	24 June 2001	15	10	119,647	41	(423-01,193) 8546 (201-28,550)	642	394	0.33	18,856	99.9983	5	0.00167
Castilleja virgata (Domb. ex Wedd.)	Lamiales/Orobanchaceae	Olmstead 2009-22	WTU	Silica gel-dried	5 Mar. 2009	15	Γ	113,650	21	5412 (178–39,914)	869	1525	1.34	14,541	99.9938	٢	0.00616
Castilleja ortegae	Lamiales/Orobanchaceae	Egger 1213	WTU	Silica	22 Feb. 2002	15	13	108,071	ŝ	36,024	925	198	0.18	97,615	1666.66	1	0.00093
Castilleja lineariloba (Benth.) T. I. Chuang	Lamiales/Orobanchaceae	Tank 2002-04	WTU	gel-uneu Silica gel-dried	27 Apr. 2004	14	9,10	122,182	23	(C10,12–202) 5312 (179–36,972)	540	810	0.66	11,656	99.9844	19	0.01555
& Heckard Castilleja victoriae Fairbarns & J. M.	Lamiales/Orobanchaceae	Fairbarns s.n.	WTU	Silica gel-dried	21 July 2005	14	10, 14	111,371	10	11,137 (616–44,011)	688	547	0.49	18,398	99.9982	7	0.00180
Egger Lamourouxia virgata Viinth	Lamiales/Orobanchaceae	Zak & Jaramillo 3307	. Е	Herbarium	16 Jan. 1988	14	9, 10	108,767	30	3626 7714 36 8500	652	2255	2.07	11,012	6996.66	36	0.03310
Castilleja oresbia	Lamiales/Orobanchaceae	Tank 2001-27	WTU	Silica	19 June 2001	10	6, 9, 10, 13,	83,384	20	(0000000000000000000000000000000000000	717	1544	1.85	9866	9676	27	0.03238
Cham. & Schltdl.	Lamiales/Orobanchaceae	Tank 2005-27	MTU	ger-ured Silica gel-dried	16 Apr. 2005	9	4, 6, 7, 8, 9, 10, 13, 14, 10, 15, 16	73,378	15	(186–36,621) (186–36,621)	701	1187	1.62	9803	99.9877	6	0.01227
Penstemon montanus Greene var. idahoensis (D. D. Vaal) Conor	Lamiales/Plantaginaceae	Brunsfeld 4159	Ð	Herbarium	14 June 2001	16	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Balsamorhiza sagittata	Asterales/Asteraceae	Willard	D	Silica	3 July 2013	15	5	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
(Pursh) Nutt. Lomatium dissectum (Nutt.) Mathias &	Apiales/Apiaceae	2013-42 Poor 21	₽	gel–dried Herbarium	27 May 2004	15	14	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Constance Nuphar polysepala	Nymphaeales/Nymphaeaceae	Morales-Briones	Ð	Silica	8 July 2013	15	5	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Engelm. Salix scouleriana Barratt ev Hook	Malpighiales/Salicaeae	412 Brunsfeld 7213	Ð	gel-dried Herbarium	11 June 2008	15	6	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Crataegus columbiana Howell	Rosales/Rosaceae	Hetrick 1005	Ð	Herbarium	10 Apr. 1996	13	9, 14, 17	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Polygonum douglasii Greene	Caryophyllales/Polygonaceae	Smith 8040	Ð	Herbarium	23 June 2005	12	5, 6, 9, 15	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Umbellularia californica (Hook. & Am.) Nutt.	t Laurales/Lauraceae	Halse 6901	Ð	Herbarium	28 Mar. 2002	12	6, 8, 9, 10	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Bromus tectorum L. Alnus rhombifolia Nutt.	Poales/Poaceae Fagales/Betulaceae	Clippinger 2 Gray 52	ΒΩ	Herbarium Herbarium	1 May 2004 7Aug. 1989	11 10	5, 6, 9, 11, 17 5, 6, 8, 9,	n/a n/a	n/a n/a	n/a n/a	n/a n/a	n/a n/a	n/a n/a	n/a n/a	n/a n/a	n/a n/a	n/a n/a
Poa bulbosa L.	Poales/Poaceae	Willard 2013-26	D	Silica orl_dried	3 July 2013	10	5, 6, 9, 12, 14 13 14	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Senecio integerrimus Nutt. var. exaltatus (Nutt.) Cronq.	Asterales/Asteraceae	Willard 2013-21	Ð	Silica gel-dried	3 July 2013	10	3, 5, 6, 8, 9, 11	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a

						No. of	Region no.				Ave.	No. of				No. of	% of
				Type of	Collection	amplified	not	Base pairs	No. of	CAL bp	assembly	masked	% of		% called a	mbiguous a	umbiguous
Species	Order/Family	Collection no.	Herbarium	tissue	date	regions	amplified ^b	sequenced ^c	contigs	(min-max)	depth	pb^q	masked bp	N50	$bases^e$	bases	bases
Abies amabilis Douglas Pinak	ss/Pinaceae	1419-46	WA Park 3	Silica	24 May 2009	6	4, 6, 7, 9,	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
ex J. Forbes			Arb.	gel-dried			10, 11, 12										
Capsella bursa-pastoris Brass	icales/Brassicaceae	Brunsfeld	Ð	Herbarium	1 June 2005	~	4, 6, 8, 9, 10,	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
(L.) Medik.		6313					13, 14, 17										
Lupinus leucophyllus Fabal	es/Fabaceae	Willard	Ð	Silica	3 July 2013	%	1, 6, 8, 9, 10,	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Douglas ex Lindl.		2013-03		gel-dried			12, 13, 14										
Abies fraseri (Pursh) Pinak	es/Pinaceae	1005-47	WA Park	Silica	24 May 2009	2	4, 5, 6, 7, 8, 9,	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
Poir.			Arb.	gel-dried			10, 11, 12										
Balsamorhiza Aster	ales/Asteraceae	Smith 9421	Ð	Herbarium	4 June 2007	Г	4, 5, 6, 7, 8, 9,	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
hookeri Nutt.							10, 11, 13										
Abies grandis (Douglas Pinak	es/Pinaceae	1084-49	WA Park	Silica	24 May 2009	9	1, 3, 4, 6, 7,	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a	n/a
ex D. Don) Lindl.			Arb.	gel-dried			8, 9, 10,										
							11, 12										
Average								11,493	14.13	3,166.60	698.73	833.0′	0.79	35,052.87	66.66	10.6	0.01
Note: CAL = contig average	e length; F = Field Mus	seum of Natural Hi	story Herbariu	ım; ID = Uni	iversity of Idaho	Stilling	r Herbarium; W	A Park Arb.	= Washir	gton Park Arbo	retum; WT	U = Uni	versity of W	ashington l	Herbarium.		
^a All data from the 16 chose	in primer combinations																
^b The number of the regions	s is the same as the orde	er in Fig. 1.															
c Base pairs (bp) sequenced	is the sum of all contig	gs when including o	only one copy	of the inverte	ed repeat.												
^d Number of bases masked l	because the minimum s	sequencing depth of	f 5× was not a	chieved.													

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Percentage of unambiguously called bases

Sanger sequencing and NGS technologies. Here, we use long PCR to generate chloroplast DNA templates for systematic studies using NGS. While we focus on whole chloroplast amplification, this approach is directly translatable to targeted studies where only particular regions of the plastome are of interest (e.g., the inverted repeat or the small single-copy region). In addition, long PCR could also be very useful for the enrichment of mitochondrial and/or nuclear regions where intron sizes are large or unknown, as well as for regions that are difficult to assemble bioinformatically, such as repetitive regions.

Our focus on the chloroplast genome is driven by its phylogenetic informativeness at essentially all taxonomic scales and its relative ease of amplification (e.g., Downie and Palmer, 1992; Graham and Olmstead, 2000; Moore et al., 2007; Parks et al., 2009; Moore et al., 2010), which have made the chloroplast the workhorse of molecular plant systematics since the beginning of the field. Moreover, the availability of a large number of angiosperm plastome sequences had facilitated the design of potentially universal PCR primers. To test this approach, we amplified the chloroplast genomes of 30 species (17 genera) across angiosperms using a set of 58 chloroplast PCR primers that were designed to potentially be universal in angiosperms and that may work in some gymnosperm lineages.

METHODS AND RESULTS

Representatives of 17 different genera (30 spp.) spanning 12 orders of angiosperms sensu APG III (Angiosperm Phylogeny Group, 2009) were chosen to test this approach (Table 1). Special focus was given to three genera in Orobanchaceae: Lamourouxia Kunth (one species), Bartsia L. (two species), and Castilleja Mutis ex L. f. (12 species). High-quality genomic DNA was extracted from ca. 0.02 g of silica gel-dried or herbarium tissue using a modified 2× cetyltrimethylammonium bromide (CTAB) method (Doyle and Doyle, 1987), yielding 30–70 ng/ μL of DNA per sample. Using the 83 plastid gene angiosperm alignments of Moore et al. (2010; Appendix S1), we developed 58 primers with a goal of maximizing universality across angiosperms (Table 2). Conserved regions for primer design were identified by eye, and the primers were tested with IDT OligoAnalyzer tools (Integrated DNA Technologies, Coralville, Iowa, USA) to ensure that melting temperatures (T_m) were greater than 50°C and that there were no significant hairpins or self-dimerization problems. From these, 16 overlapping primer combinations were chosen to amplify the entire chloroplast genome in appropriately sized, overlapping fragments, making sure to allow at least 100 bp of overlap between regions (Fig. 1, Table 2) to minimize the decrease in sequencing depth usually associated with the ~30 bp immediately adjacent to the primer sites (Cronn et al., 2008; Harismendy and Frazer, 2009; Cronn et al., 2012).

PCRs were performed using a combination of two high-quality Taq polymerases-QIAGEN Taq DNA Polymerase (5 units/µL) and QIAGEN HotStar HiFidelity DNA Polymerase (2.5 units/µL) (QIAGEN, Valencia, California, USA)-to obtain amplification of fragments between 5 kb and 12 kb. The QIAGEN HotStar HiFidelity DNA Polymerase was diluted to 0.2 units/µL by combining 0.1 µL of 5× QIAGEN HotStar HiFidelity PCR buffer, 0.36 µL of double-deionized water (ddH2O), and 0.04 µL of QIAGEN HotStar HiFidelity DNA Polymerase (2.5 units/ μ L). Each PCR had a total volume of 25 μ L, was prepared on ice, and contained the following reagents: 2.5 µL of 10× PCR buffer (QIAGEN CoralLoad or colorless, with 15 mM MgCl₂), 1.0 µL MgCl₂ (QIAGEN 25 mM), 0.75 µL of deoxyribonucleotide triphosphates (dNTPs, each at 10 mM), 5.0 µL of 5× QIAGEN Q solution, 2.5 µL of both forward and reverse primers (each at 5 µM), 0.25 µL (1.25 units) of QIAGEN Taq DNA Polymerase, 0.5 µL of the diluted QIAGEN HotStar HiFidelity DNA Polymerase solution, 9 µL of ddH2O, and 1.0 µL of DNA template. Long PCR profiles were as follows: preheat at 93°C, initial denaturation at 93°C for 3 min followed by 35 cycles of denaturation at 93°C for 15 s, annealing at 48-68°C (depending on the primer pair) for 30 s, and extension at 68°C for 5-12 min (1 min/kb of target). To assess amplification, 2 µL of the final reactions were examined on a 1% agarose gel with appropriate size standards and the final products were kept at 4°C. The complete, step-by-step long PCR protocol can be found in Appendix 1.

TABLE 2. Universal angiosperm primers used for chloroplast genome amplifications. The 16 primer combinations chosen for this study are in bold with approximate amplicon sizes in kilobases (kb) indicated.^a

Region no.	Approx. size (kb)	Primer (F/R)	Primer sequence (5'–3')	Overlap between regions in bp ^b
1	8	trnH.GUG.6R	CCTTRATCCACTTGGCTACAT	Regions 1 & $2 = 542$
1	Ū.	nsbK.195R	ACTTACAGCAGCTTGCCAAAC	Regions 1 & $2a = 542$
2/2a	10 3/6 3	trnO.UUG.50R	GGACGGAAGGATTCGAACC	Regions $2a \& 2b = 627$
2a	1010/010	atpH.17F	CTGCYGCTTCYGTTATTGCT	Regions 2b & $3 = 2059$
2h	4	atpF.65R	CGGTATTAAACCCGAAACTCC	Regions 2 & $3 = 2059$
2/2h		rpoC2.4805F	GYCGTATYGATTGGTTRAAAGG	Regions 3 & $4 = 1274$
3	7	atpl.705R	CRGCTAAAGTTGCAAAAATAAGAGCT	Regions 4 & $5 = 860$
3		rpoC1.1670F	GRGATCAAATGGCTGTTCAT	Regions 5 & $6 = 618$
4	9	rpoC2.520R	GTTCGTACAGCAGTATCYACAAC	Regions 6 & $7 = 764$
4	· · · · · · · · · · · · · · · · · · ·	netN.3R	GCCCAAGCBAGACTTACTATATCC	Regions 7 & $8 = 153$
5	10.5	trnC.GCA.47F	СССАСТТСАААТСССССТ	Regions 8 & $9 = 1216$
5	1010	nsaB.2170F	GCBGCTTTCTTGATTGCYTC	Regions 9 & $10 = 135$
6	10	trnfM.CAU.21R	GGTTATGAGCCTTGCGAGCTA	Regions 10 & $11 = 771$
6	10	trnT.UGU.17F	GGTTAGAGCATCGCATTTGTAATG	Regions 11 & $12 = 2781$
7	10.3	rns4.380R	GGTTTGCARCGATAACTTGGKATATC	Regions 12 & $13 = 142$
7	10.0	rbcL 178R	GTCCATGTACCAGTAGARGATTC	Regions 13 & $14 = 392$
8	9.2	rbcL.2F	TGTCACCACACACAGARACTAAAG	Regions 14 & $15 = 1911$
8	7.2	nsh L 3F	GCCYCATACTACTGGAAGRAT	Regions 16 & $1 = 840$
9	9.8	netA.920F	CTTCAAGAYCCATTACGTGTHCAAG	
9	2.0	nshB.160R	TRCCYTGTCTCCACATTGGAT	
10	10.9	nshB 3F	GCGTTTRCCTTCGTATCGTGT	
10	10.9	rns3 17F new		
10	87	rnl16 3R		
11/16	0.7	vcf2 5100R		
12	10	vcf2 2300F		
12	10	rns12.190F		
12	11	rps12.1901		
13	11	vcf1.59R	CCGACCACAACGACCGAAT	
14/15	11.2	trnN.GUU.7R	CCGCTCTACCACTGAGCTAC	
14		ndhA.535F	GCTGCTCAATCDATTAGTTATGAA	
15	10.5	ndhL194R	CGAACRCATACTTCACAAGCAA	
16	8.2	nsbA.640F	GCTATGCATGGTTCYTTGGTAAC	
		rps16.50R	CGAACATCAATTGCAACGATTCGATA	
		rps16.50F	TATCGAATCGTTGCAATTGATGTTCG	
		psbK.200F	GGCAAGCTGCTGTAAGTTTTCGA	
		atpE.70F	GGGTTTAATACCGATATTTTAGCAAC	
		trnR.UCU.45F	GGTATAGGTTCAAATCCTATTGGAC	
		trnO.UUG.47F	CGGAGGTTCGAATCCTTCC	
		trnK.UUUU.3R	GAGATGGCAACTCAATCGTTG	
		trnK.UUU.3F	CAACGATTGAGTTGCCATCTC	
		atpA.430F	CGTTCYGTATATGARCCTCTTCAAAC	
		atpA.820F	ATCGMCAAATGTCTCTTCTATTAMG	
		ccsA.890R	TCCAAGTAATAAANGCCCAAGTTTC	
		trnR.ACG.15F	GAGGATTAGAGCACGTGG	
		vcf1.70F	GTGGTCGGACTCTATTATGGAT	
		trnL.UAG.18F	GGTAGACACGCTGCTCTTAGG	
		trnL.UAG.19F	GTAGACACGCTGCTCTTAGGAAG	
		rps12.320R	GGGTTCCTCGAACAATGTGATATC	
		rpl2.550F	GTGCTGTAGCGAAACTGATTG	
		rp12.640F	TCAGCAACAGTCGGACARGT	
		psbT.3F	TGGAAGCATTGGTTTATACATTYCT	
		atpB.1290R	ARGGTTGTGATAAGAAACGYTCAA	
		trnT.UGU.42F	GATGGTCATCGGTTCGATTC	
		psbC.3R	AGTTCCATTAAAGAGCGTTTCC	
		psbD.860F	CYGGTTTATGGATGAGYGCT	
		rpoB.900R	CGTCGACCAATCYTTCCTAATTC	
		rpoB.470R	CCRGGRCTTTGCAATATTTGATTG	
		rpoC2.430R	ATRGGTAAATCAATCATTTGYCCTTG	

^aAll primers are shown in the 5' to 3' direction; the name of each primer consists of three parts: the gene in which the primer is anchored, the approximate position of the primer within that gene, and either an "F" or an "R." It is important to note that the F and R designations do not indicate that the primer should be used as a forward or reverse primer; rather, they indicate the 5' to 3' orientation of the primer with respect to the gene—i.e., a primer that is designated as an "F" primer has its 5' to 3' orientation in the same orientation as the gene (i.e., on the forward strand), whereas an "R" primer is oriented in the direction opposite to the 5' to 3' orientation of the gene (i.e., on the reverse strand).

^bOverlap between regions is given in number of base pairs (bp), without taking the length of the primers into consideration.

For the three genera of Orobanchaceae in which PCR optimization was performed, amplification of the fragments was straightforward and had an average success rate of 89.7% (range = 73-100%). The most difficult regions to

amplify were regions 2 ($trnQ^{(UUG)}$ -rpoC2), 9 (petA-psbB), 10 (psbB-rps3), and 14 ($trnN^{(GUU)}$ -ndhA), which are among the largest fragments (10.3 kb, 9.8 kb, 10.9 kb, and 11.2 kb, respectively; Table 2). It was possible to split



Fig. 1. The final annotated chloroplast genome assembly of *Bartsia inaequalis* with the 16 overlapping primer combinations indicated. Note that the primer combinations for regions 11, 12, 13, and 16 amplify both inverted repeat A and B in a single reaction. Photos by Simon Uribe-Convers.

region 2 into two smaller fragments, 2a ($trnQ^{(UUG)}$ -atpH: 6.3 kb) and 2b (atpF-rpoC2: 4 kb), which facilitated its amplification in several taxa. This was not the case for regions 9, 10, and 14, for which multiple long PCR experiments using varying amounts of DNA template were necessary to obtain successful amplifications. Amplification outside of Orobanchaceae was highly variable, with an average success rate of 70.8% (range = 22–100%) with regions 5, 6, 9, 10, and 11 showing the lowest success. Importantly, the results for these taxa were obtained after just two rounds of PCR where the annealing temperatures were changed to either 48°C or 55°C. Although we did not optimize the long PCRs for each group, we are confident that optimization on a per group basis (e.g., increasing template volume, altering annealing temperatures, and/or long PCR profiles) and/or the use of fresh

tissue for DNA extractions would improve success rates. Furthermore, if genomic rearrangements and/or primer mismatches are present in certain groups, primer combinations other than the 16 that were used here could be tested (Table 2). Nevertheless, we successfully amplified all 16 regions in seven species, whereas in the remaining 23 species it was only possible to amplify between six (1 sp.) and 15 (8 spp.) regions (Table 1). These results translate to 21 species having at least 12 regions amplified (114.7 kb based on potential amplicon size), representing ca. 74% of the chloroplast genome when considering only one copy of the inverted repeat. Even the species with the smallest number of amplified fragments (*Castilleja arvensis* Cham. & Schltdl.) was represented by ~73 kb of data, exemplifying the effectiveness of this approach.

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It is notable that many of the DNAs that were tested were extracted from herbarium tissues that ranged from five to 25 yr old when isolated. In addition, we tested these primers in several species of *Abies* Mill. (Pinaceae; Table 1) with surprising success, amplifying between six and nine regions without any PCR optimization. We caution that our long PCR protocol works best using recent DNA extractions that have not been through multiple freeze-thaw cycles. Ideally, long PCR should be conducted using new DNA extractions that are stored at 4°C while performing experiments. Additionally, discrete PCR bands were only obtained using high-quality *Taq* polymerases. When conventional polymerases were used (e.g., Go*Taq* [Promega Corporation, Madison, Wisconsin, USA] or Top*Taq* [QIAGEN]), the resulting PCR products were smears rather than discrete bands and were not used for sequencing.

To confirm that our long PCR approach was compatible with NGS and that our primers would yield complete chloroplast genomes, the amplicons from each of the 15 Orobanchaceae taxa were purified by precipitation in a 20% polyethylene glycol 8000 (PEG)/2.5 M NaCl solution and washed in 70% ethanol. The amplicons were sheared by nebulization at 30 psi for 70 s, yielding an average shear size of 500 bp as measured by a Bioanalyzer High-Sensitivity Chip (Agilent Technologies, Santa Clara, California, USA). DNA normalization is a critical step when pooling samples for multiplexing in NGS; however, due to the large number of plastomes per cell and the very few samples that were being sequenced in such a high-throughput sequencing platform, no DNA quantification was made and the sheared amplicons were pooled by species at equal volume ratios. Sequencing libraries were constructed using the Illumina TruSeq library preparation kit and protocol (Illumina, San Diego, California, USA) and were standardized at 2 nM prior to sequencing. Library concentrations were determined using the KAPA qPCR kit (KK4835; Kapa Biosystems, Woburn, Massachusetts, USA) on an ABI StepOnePlus Real-Time PCR System (Life Technologies, Grand Island, New York, USA). The resulting libraries were multiplexed in one Illumina HiSeq 2000 lane (~187.5 million reads per lane [Glenn, 2011]) at the Vincent J. Coates Genomics Sequencing Laboratory at the University of California, Berkeley, yielding ~12.5 million 100-bp singleend reads for each taxon (GenBank Sequence Read Archive accessions: SRR1023085, SRR1023089, SRR1023095, SRR1023112, SRR1023113, SRR1023126, SRR1023128-SRR1023136). Average depth of coverage of our sequencing experiment was ~8333× (taking 150 kb as the average plastome size). The results obtained here clearly do not maximize the potential of the Illumina HiSeq 2000 for plastome sequencing. To take full advantage of the large amount of data produced by a HiSeq 2000 for plastome sequencing, it would be theoretically possible to sequence ~4170 samples per lane and still reach the 30× minimum threshold generally regarded as ideal for plastome sequencing (Straub et al., 2012). However, high-level multiplexing in NGS with this or any other high-throughput method requires careful normalization of DNA concentrations across samples and sufficient adapter barcodes; commonly used commercial kits currently offer either 96 (NEXTflex DNA Barcode kit; Bioo Scientific, Austin, Texas, USA) or 386 (Fluidigm, San Francisco, California, USA). Alternatively, one could choose to perform this type of experiment on an NGS platform that yielded a lesser amount of data, e.g., 1 million 250-bp paired-end reads on an Illumina MiSeq Reagent Nano Kit version 2, which would allow a 30× sequencing depth for 96 samples (or 50× sequencing depth for 64 samples).

Because of the high depth of coverage of our sequencing experiment, reads were cleaned at high stringency (minimum quality = 30/40, maximum number of low-quality bases per read = 5, maximum number of duplicate reads = 10, minimum number of duplicate reads = 2) and assembled against a reference genome (Sesamum indicum L., GenBank accession no. JN637766) using the Alignreads pipeline version 2.25 (Straub et al., 2011) with the following options: percent identity = medium, minimum coverage depth = 5, and single nucleotide polymorphism (SNP) minimum coverage depth = 25 with 80% of those reads supporting the SNP. The resulting assemblies had an average depth of ~700×, an average of 0.79% bases that were masked for not reaching the minimum sequencing depth of 5×, and an average N50 of 35,053 bp (Table 1; contigs and ACE files deposited in the Dryad Digital Repository: http://doi.org/10.5061/dryad.kc75n; Uribe-Convers et al., 2014). We noticed a small decrease in sequencing depth in regions immediately adjacent to some primer sites, which is a phenomenon that has been reported in the past (Whittall et al., 2010; Knaus et al., 2011; reviewed in Cronn et al., 2012). Given that our shortest overlap between amplicons is 135 bp (between regions 9 and 10; Table 2), with the rest spanning hundreds of base pairs (Table 2), and that our experiment yielded a high sequencing depth, we had no problems calling bases unambiguously (99.99% on average, Table 1). The Bartsia inaequalis Benth. assembly (Fig. 1; GenBank accession no. KF922718) was annotated using DOGMA (Wyman et al., 2004) and visualized in GenomeVx (Conant and Wolfe, 2008).

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We present an alternative approach for systematic studies that combines long PCR and NGS to strategically compile phylogenomic data sets for molecular systematic studies. This approach is on par with genome skimming in terms of costs, but it has the advantage of being a targeted approach and has the potential to produce data more uniformly across samples, i.e., minimizing missing data across taxa. Although this approach was only tested with chloroplast data, we emphasize that the long PCR amplicons can be generated using DNA from any genome, expanding the possibilities of long PCR and NGS for molecular systematic studies. This last point is important for studies targeting the mitochondrion or low-copy regions of the genome that otherwise might be missed or not shared across all samples using genome skimming approaches. For example, this approach may be particularly useful for the enrichment of nuclear regions, where intron sizes are large or unknown.

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APPENDIX 1. Protocol for long PCR for amplification of 4–20-kb targets. Developed by the Tank Laboratory, University of Idaho; published January 2014.

Product	Contents	Catalog no.
QIAGEN Taq DNA Polymerase ¹	250 units Taq DNA Polymerase, 10× PCR Buffer, [†] 5× Q-Solution, 25 mM MgCl ₂	201205
QIAGEN HotStar HiFidelity DNA Polymerase ²	100 units HotStar HiFidelity DNA Polymerase, ² 10× HotStar PCR Buffer, 5× Q-solution, 25 mM MgSO ₄	202602

¹Almost any high-quality *Taq* polymerase should work; however, cheap *Taq* polymerases (e.g., QIAGEN Top*Taq* or Promega Go*Taq*) do not work and result in large smears, rather than discrete bands.

²QIAGEN HotStar HiFidelity DNA Polymerase was the only high-fidelity polymerase used in this study.

[†]Q-solution does seem to be an important additive, thus the use of QIAGEN *Taq*. However, this does work using Q-solution with other high-quality *Taq* polymerases such as Promega's or New England Biolab's standard *Taq* (i.e., if you have a stock of Q-solution, but no QIAGEN *Taq*).

Genomic DNA must be high quality. Run a 0.8% or 1% gel to check. Standard CTAB extractions from silica gel-dried or herbarium material work well if they (1) are recent (extraction and tissue), and (2) contain high-molecular-weight DNA. Most important, we have found that recent DNA extractions that have not been through numerous freeze-thaw cycles work best. For best results, long PCR should be done using new DNA extractions stored at 4°C while performing long PCR experiments.

All preparations should be done on ice.

1. Number tubes or prepare plate. Make sure to include appropriate negative controls.

2. Prepare QIAGEN HotStar HiFidelity DNA polymerase dilution:

Reagents to prepare the HotStar Taq dilution	Volumes for 25 reactions (total 12.5 μ L)	Volumes for 50 reactions (total 25 $\mu L)$	Volumes for 100 reactions (total 50 µL)
5× HotStar HiFidelity PCR buffer	2.5 μL	5.0 μL	10 μL
H ₂ O	9.0 μL	18 μL	36 μL
QIAGEN HotStar <i>Taq</i>	1.0 μL	2.0 μL	4.0 μL

3. Prepare cocktail:

Cocktail	$\times 1$ (25 µL reaction)
10× PCR buffer (QIAGEN CoralLoad PCR Buffer or colorless, 15 mM MgCl ₂)	2.5 μ L
MgCl ₂ (25 mM)	1.0 μ L (3 mM final conc.; adjustable)
dNTR (10 mM each)	0.75 μ L (2 μ L of 2.5 mM coch)
Q solution (5×) 5' primer (5 μ M)	$5.0 \mu\text{L}$ (5 μL of 2.5 mW each) 5.0 μL 2.5 μL (0.5 μ M final conc.)
3' primer (5 μM)	2.5 μ L (0.5 μ M final conc.)
<i>Taq</i> DNA polymerase (QIAGEN)	0.25 μ L (1.25 units) ¹
QIAGEN HotStar DNA polymerase (diluted)	0.50 μL
H ₂ O	to 25 μL (9 μL if using 1.0 μL DNA)

 1 The success rate was lower when a smaller quantity was used, but the best DNAs work with ${\geq}0.125~\mu L.$

4. Add $1-2 \mu L$ of template to each of the tubes.

5. While the tubes/plate with template are on ice, add $24 \,\mu\text{L}$ of cocktail to each tube, being careful not to cross contaminate. Spin down to bring all liquid to the bottom of the tube.

6. Run appropriate long PCR profile. Generic temperatures and times are:

i. 93°C infinity (important to go directly from ice to hot block) ii. 93°C for 3 min (initial denaturation) iii. 93°C for 15 s iv. 48–68°C for 30 s (T_a should be ~5°C below T_m of primers) v. 68°C for 5–20 min (1 min/kb of target) vi. go to step 3, 34× vii. 4°C infinity

7. Check reactions by running $2 \,\mu$ L on 1% agarose gel with appropriate size standards.

Primer	combinations	for long	PCR am	plification	of the	chloroplast	genome. ^{1,2}
							0

Region no.	Approx. size (kb)	Primers (F/R) ³	Primer sequence $(5'-3')$
1	8	trnH.GUG.6R	CCTTRATCCACTTGGCTACAT
		psbK.195R	ACTTACAGCAGCTTGCCAAAC
2	10.3	trnQ.UUG.50R	GGACGGAAGGATTCGAACC
		rpoC2.4805F	GYCGTATYGATTGGTTRAAAGG
$2a^{4}$	6.3	trnQ.UUG.50R	GGACGGAAGGATTCGAACC
		atpH.17F	CTGCYGCTTCYGTTATTGCT
$2b^{4}$	4	atpF.65R	CGGTATTAAACCCGAAACTCC
		rpoC2.4805F	GYCGTATYGATTGGTTRAAAGG
3	7	atpI.705R	CRGCTAAAGTTGCAAAAATAAGAGCT
		rpoC1.1670F	GRGATCAAATGGCTGTTCAT
4	9	rpoC2.520R	GTTCGTACAGCAGTATCYACAAC
		petN.3R	GCCCAAGCRAGACTTACTATATCC
5	10.5	trnC.GCA.47F	CCCAGTTCAAATCCGGGT
		psaB.2170F	GCRGCTTTCTTGATTGCYTC
6	10	trnfM.CAU.21R	GGTTATGAGCCTTGCGAGCTA
		trnT.UGU.17F	GGTTAGAGCATCGCATTTGTAATG
7	10.3	rps4.380R	GGTTTGCARCGATAACTTGGKATATC
		rbcL.178R	GTCCATGTACCAGTAGARGATTC
8	9.2	rbcL.2F	TGTCACCACAAACAGARACTAAAG
		psbJ.3F	GGCYGATACTACTGGAAGRAT
9	9.8	petA.920F	CTTCAAGAYCCATTACGTGTHCAAG
		psbB.160R	TRCCYTGTCTCCACATTGGAT
10	10.9	psbB.3F	GGGTTTRCCTTGGTATCGTGT
		rps3.17F.new	ATCCACTTGGTTTYMGACTTGG
11	8.7	rpl16.3R	AACCAACGAGTCACACACTAAGC
		ycf2.5100R	CAGATCATGAATGTTTGGAATCCAT
12	10	ycf2.2300F	TCGGGATCCTRATGCATATAGATAC
		rps12.190F	GTTGCCAGAGTACGMTTAACCT
13 ⁵	11	rps12.360R	CCCTTGTTGACGATCCTTTACTC
		ycf1.59R	CCGACCACAACGACCGAAT
14	11.2	trnN.GUU.7R	CCGCTCTACCACTGAGCTAC
		ndhA.535F	GCTGCTCAATCDATTAGTTATGAA
14'6	7	trnR.ACG.15F	GAGGATTAGAGCACGTGG
		ccsA.890R	TCCAAGTAATAAANGCCCAAGTTTC
15	10.5	ndhI.194R	CGAACRCATACTTCACAAGCAA
		trnN.GUU.7R	CCGCTCTACCACTGAGCTAC
16	8.2	psbA.640F	GCTATGCATGGTTCYTTGGTAAC
		ycf2.5100R	CAGATCATGAATGTTTGGAATCCAT

¹Universal primers designed by M.J.M.; compiled and tested by D.C.T. and S.U.C.

 ${}^{2}T_{a}$ should be ${}^{-5}$ °C below T_{m} of primers; however, temperatures of 55°C have worked for all primer combinations.

³The name of each primer consists of three parts: (1) the gene in which the primer is anchored in, (2) the approximate position of the primer within that gene (based on all-angiosperm alignment per Moore et al., 2007), and (3) either an "F" or an "R." The F and R designations do not indicate that the primer should be used as a forward or reverse primer; rather, they indicate the 5′ to 3′ orientation of the primer with respect to the gene. In other words, a primer that is designated as an F primer has its 5′ to 3′ orientation in the same orientation as the gene (i.e., on the forward strand, or from start to stop), whereas an R primer is oriented in the direction opposite to the 5′ to 3′ orientation of the gene (i.e., on the reverse strand).

⁴Regions 2a and 2b can be used to amplify region 2 in two pieces.

⁵Regions 11, 12, and 13 represent a large portion of the inverted repeat (IR), thus, one amplification for both IRa and IRb.

⁶Region 14' amplifies ca. 2/3 of region 14.