



Developing molecular tools and insights into the *Penstemon* genome using genomic reduction and next-generation sequencing

Dockter *et al.*

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Developing molecular tools and insights into the *Penstemon* genome using genomic reduction and next-generation sequencing

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Abstract

Background: *Penstemon*'s unique phenotypic diversity, hardiness, and drought-tolerance give it great potential for the xeric landscaping industry. Molecular markers will accelerate the breeding and domestication of drought tolerant *Penstemon* cultivars by, creating genetic maps, and clarifying of phylogenetic relationships. Our objectives were to identify and validate interspecific molecular markers from four diverse *Penstemon* species in order to gain specific insights into the *Penstemon* genome.

Results: We used a 454 pyrosequencing and GR-RSC (genome reduction using restriction site conservation) to identify homologous loci across four *Penstemon* species (*P. cyananthus*, *P. davidsonii*, *P. dissectus*, and *P. fruticosus*) representing three diverse subgenera with considerable genome size variation. From these genomic data, we identified 133 unique interspecific markers containing SSRs and INDELs of which 51 produced viable PCR-based markers. These markers produced simple banding patterns in 90% of the species × marker interactions (~84% were polymorphic). Twelve of the markers were tested across 93, mostly xeric, *Penstemon* taxa (72 species), of which ~98% produced reproducible marker data. Additionally, we identified an average of one SNP per 2,890 bp per species and one per 97 bp between any two apparent homologous sequences from the four source species. We selected 192 homologous sequences, meeting stringent parameters, to create SNP markers. Of these, 75 demonstrated repeatable polymorphic marker functionality across the four sequence source species. Finally, sequence analysis indicated that repetitive elements were approximately 70% more prevalent in the *P. cyananthus* genome, the largest genome in the study, than in the smallest genome surveyed (*P. dissectus*).

Conclusions: We demonstrated the utility of GR-RSC to identify homologous loci across related *Penstemon* taxa. Though PCR primer regions were conserved across a broadly sampled survey of *Penstemon* species (93 taxa), DNA sequence within these amplicons (12 SSR/INDEL markers) was highly diverse. With the continued decline in next-generation sequencing costs, it will soon be feasible to use genomic reduction techniques to simultaneously sequence thousands of homologous loci across dozens of *Penstemon* species. Such efforts will greatly facilitate our understanding of the phylogenetic structure within this important drought tolerant genus. In the interim, this study identified thousands of SNPs and over 50 SSRs/INDELs which should provide a foundation for future *Penstemon* phylogenetic studies and breeding efforts.

Keywords: Breeding domesticated *Penstemon*, Genome reduction, Homologous sequences, LTR retroelements, Plantaginaceae, Pyrosequencing, Repetitive elements

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Background

Interest is increasing in drought tolerant landscape plants due to water shortages experienced by many municipalities, especially in the Southwestern US [1,2]. However, the increased use of drought tolerant species also carries concerns regarding the introduction of non-native and potentially invasive species [3,4]. One way to address both issues is to landscape with native xeric flora [3]. *Penstemon* Mitchell (Plantaginaceae) has excellent potential for xeric landscapes and some *Penstemon* cultivars, adapted to mild climates, are already used throughout Europe as landscape plants [5-10]. Despite its potential, few *Penstemon* cultivars are used in xeric landscapes and there has been little to no drought or cold tolerant cultivar development for such landscapes [6-8,10-12]. *Penstemon*, with over 270 species, is one of the largest and most diverse plant genera of those that are strictly indigenous to North and Central America. This genus features a deep diversity in morphology, including a broad assortment of colors, flowers, and leaf structures. *Penstemon*'s putative center of origin is the arid Intermountain West of the United States [13,14] and has frequently been discussed as an untapped resource for xeric landscape cultivar development [5-7,9-11,15-17]. Because domestication and cultivar development, of any species, is slow, costly, and time consuming, few in the landscape industry have invested in native species breeding. However, given the recent and dramatic decrease in costs and relative ease of genotyping, we anticipate the wider utilization of marker assisted selection to accelerate breeding programs of native species, including drought tolerant *Penstemon* [18-20].

PCR-based markers are now essential tools to facilitate plant domestication, plant breeding, germplasm conservation, phylogenetics, and genetic mapping studies [19-22]. Not surprisingly, little molecular or traditional genetic work has been reported for *Penstemon* [23]. To achieve broad resolution of the genome with three of the most efficient markers, SSRs (simple sequence repeats or microsatellites), INDELs (insertions/deletions), and SNPs (single nucleotide polymorphisms), vast amounts of DNA sequence are needed, particularly for SNPs where sufficient read depth is needed to distinguish true polymorphisms from sequence noise [24-26]. With the development of next-generation sequencing (e.g., Roche 454-pyrosequencing) the cost of high-throughput marker discovery has been dramatically reduced [18]. Additionally, Maughan et al. [25] described a simple genome reduction method, known as GR-RSC (genome reduction using restriction site conservation), which reduces the genome by > 90% thereby, making it feasible to redundantly sequence the remaining genome with next-generation sequencing technologies. This process is repeated across multiple cultivars or species, with comparisons identifying

many inter- and intraspecific homologous loci. Genomic reduction techniques consistently identify homologous loci between related species [20,27], and GR-RSC has enabled the identification and development of interspecific homologous SNPs [20].

We utilized GR-RSC to identify homologous sequences in four diploid ($2n = 2x = 16$) *Penstemon* species chosen to represent a range of taxonomic and genome size diversity [5,14]. Included in our analysis are two closely related species from the subgenus *Dasanthera* (*P. davidsonii* Greene and *P. fruticosus* (Pursh) Greene var. *fruticosus*), one from the subgenus *Habroanthus* (*P. cyananthus* Hook. var. *cyananthus*), and one (*P. dissectus* Elliot) from the monophyletic subgenus *Dissecti*, which is phenotypically divergent from all other *Penstemon* species. This experimental design allowed us to make broad inter- and intra-subgenera comparisons in *Penstemon*. The objectives of our study were three-fold: First, identify homologous SSR and INDEL markers from the four diverse species and test their conservation across 93, mostly xerophilic, *Penstemon* taxa. Second, identify conserved homologous sequences for SNPs for use in future interspecific studies. Third, assess observed variation in the GR-RSC sequences to gain insights into the *Penstemon* genome and possible reasons for the large size variation previously identified among the diploid taxa [5].

Methods

Plant material and DNA extraction

DNA from *P. cyananthus*, *P. davidsonii*, *P. dissectus*, and *P. fruticosus* leaf tissue was extracted using the CTAB purification method [28] with modifications [29] for the GR-RSC technique. The source localities and identification of these plants have been reported previously [5]. A single sample from each species with the highest quality and DNA concentration, as determined using a ND-1000 spectrophotometer (NanoDrop Technologies Inc., Montchanin, DE), was selected to provide the 500 ng of DNA necessary for the genome reduction protocol.

For the molecular marker experiments, we used 93 *Penstemon* taxa. Leaf tissue was collected mostly from wild populations in the United States Intermountain West (Table 1). Each field-collected sample was identified to species and (or) variety using taxonomic keys specific to the area [30,31]. We extracted DNA using Qiagen DNeasy Plant Mini Kit (Qiagen Inc., Valencia, CA), and concentrations were diluted to 25–35 ng/μL.

Genome reduction, barcode addition and 454 pyrosequencing

Genome reduction followed Maughan et al. [25]. Briefly, for each sample, *EcoRI* and *BfaI* were used for the initial restriction digest, after which a biotin-labeled adapter was ligated to the *EcoRI* restriction site and a non-labeled adapter was ligated independently to the *BfaI* restriction

Table 1 *Penstemon* taxa (with collection counties) utilized in the 12 marker analysis with respective marker sizes

Species	County ¹	Marker sizes in bp											
		PS004	PS011	PS012	PS014	PS017	PS032	PS034	PS035	PS048	PS052	PS053	PS075
Subgenus <i>Dasanthera</i>													
<i>P. davidsonii</i>	Purchased ²	460	500	360	370	700	370	320, 950	520	440	220	320	140
<i>P. fruticosus</i> v. <i>fruticosus</i>	Purchased	460	500	360	410	700	340	360	520	420	220	320	140
<i>P. montanus</i> v. <i>montanus</i>	Utah	480	430	390	370	450	310	340, 310	470	430	200	390	115
Subgenus <i>Dissecti</i>													
<i>P. dissectus</i>	Purchased	440	860	370	380	750	370	320	920	380	220	320, 450	140
Subgenus <i>Habroanthus</i>													
<i>P. ammophilus</i>	Kane	480	800	400	430	470	300	1250, 340	470	420	230	360	125
<i>P. barbatus</i> v. <i>torreyi</i>	Garfield	420	800	400	490	500	320	340	500	410	200	360	110
<i>P. barbatus</i> v. <i>trichander</i>	San Juan	650, 480	850	420	500, 490	520	310	310	500	450, 410	200	370	130
<i>P. comarrhenus</i>	Garfield	650, 480	850	420	490	470	330, 310	310	500	430	200	360	125
<i>P. compactus</i>	Cache	440	850	400	500	490	300	300	480	410	210	390, 360	125
<i>P. cyananthus</i> v. <i>cyananthus</i>	Wasatch	420	860	400	410	750	370	310, 340	630	420	220	160, 320	160
<i>P. cyananthus</i> v. <i>subglaber</i>	Box Elder	440, 420	850	400	490, 470	500, 450	340, 310	310, 280	520	410	210	360	120
<i>P. cyanocalis</i>	Emery	440	310	420	490, 470	520	330, 320	320	480	420	210	350	120
<i>P. eatonii</i> v. <i>eatonii</i>	Utah	420	800	420	490	450	320	300	500	NM ³	210	350	135, 125
<i>P. eatonii</i> v. <i>undosus</i>	Washington	420	850	420	470	420	320	290	650, 500	410	210	340	125
<i>P. fremontii</i>	Uintah	480	850	400	430	490	320, 310	340	500	420	220	370	130
<i>P. gibbensii</i>	Daggett	480, 440	850	420	490	420	320	300	480	430	220	360	130
<i>P. idahoensis</i>	Box Elder	440	800	400	410	470	310	340	500	430	250	340	130
<i>P. laevis</i>	Kane	440	850	400	470	470	310	350, 320	500	420	220	360	125
<i>P. leiophyllus</i> v. <i>leiophyllus</i>	Iron	480	850, 490	420	430	450	310	340	480	430	220	350	120
<i>P. longiflorus</i>	Beaver	440	800	420, 400	470	470, 450	330, 310	310	500	450	230	350, 220	125
<i>P. navajoa</i>	San Juan	480	800	400	490	550	330, 300	360, 340	500	450	230	410	135, 130
<i>P. parvus</i>	Garfield	480	850	450	500, 490	490	320	300	500	430	210	380, 360	130
<i>P. pseudoputus</i>	Garfield	480	800	420, 400	430	490, 420	320	340	480	450	230, 220	350	130
<i>P. scariosus</i> v. <i>albifluvis</i>	Uintah	440	850	400	490	490	310	320	480	410	210	370	115
<i>P. scariosus</i> v. <i>cyanomontanus</i>	Uintah	440	850	400	490	490	330	310	500	420	210	360	115
<i>P. scariosus</i> v. <i>garrettii</i>	Duchesne	490, 480	850	420	430	490	320	420, 340	520	430	230	360	125
<i>P. scariosus</i> v. <i>scariosus</i>	Sevier	480	1500, 1300	400	470	520	340, 310	310	500	430	210	360	130
<i>P. speciosus</i>	Box Elder	440	800	420	500, 490	490	320	340, 310	520	310	210	360	120

Table 1 *Penstemon* taxa (with collection counties) utilized in the 12 marker analysis with respective marker sizes (Continued)

<i>P. strictiformis</i>	San Juan	480	850	400	470	500, 470	370, 310	350	500	410	220	370	125
<i>P. strictus</i>	Wasatch	480	850	400	410	450	310	350	520, 500	430	230	340	110
<i>P. subglaber</i>	Sevier	480	850	420, 400	470	490	310	350	500	430	220	350	115
<i>P. tidestromii</i>	Juab	390	850	420	470	490	310	300	480	400	190	360	140, 120
<i>P. uintahensis</i>	Duchesne	480	850	420	490	450	340	300	520	410	220	380	120
<i>P. wardii</i>	Sevier	480	800	420	430	490, 450	310	310	520	420	220	340	135, 120
Subgenus <i>Penstemon</i>													
<i>P. abietinus</i>	Sevier	440	AD ⁴	390	400	520	320	340	500	430	230	350	125
<i>P. acaulis</i>	Daggett	570	490	420	430	470	320	350	480	420	220	340	120
<i>P. ambiguus</i> v. <i>laevissimus</i>	Washington	520	850	390	490	470	320	1250, 340	500	400	220	AD	120
<i>P. angustifolius</i> v. <i>dulcis</i>	Millard	440	850, 600	400	490	520	370, 150	310	520	420	220	360	125
<i>P. angustifolius</i> v. <i>venosus</i>	San Juan	480	310	390	470	470	320, 150	340	550	450	220	360	135
<i>P. angustifolius</i> v. <i>vernalensis</i>	Daggett	480	800	390	430	470	370, 150	350	500	420	220	380	125
<i>P. atwoodii</i>	Kane	440	800	420	490, 390	470	300	310	480	400	180	360, 280	115
<i>P. bracteatus</i>	Garfield	440	850	400	500	AD	330, 310	320	520	420	230	380	125
<i>P. breviculus</i>	San Juan	650, 480	190	400	AD	470	500, 220	320	480	430	210	350	125
<i>P. caespitosus</i> v. <i>caespitosus</i>	Uintah	440	850	390	390	490	320	NM	190	NM	210	390, 370	115
<i>P. caespitosus</i> v. <i>desertipicti</i>	Washington	440	230	390	470, 370	470, 360	330	350	1000, 300	430	210	400, 380	130
<i>P. caespitosus</i> v. <i>perbrevis</i>	Wasatch	420	490	390	430, 400	470	320	350	520	380	220	340	120
<i>P. carnosus</i>	Emery	440	850	420	490	490	330, 300	310	500	430	220	350	130, 120
<i>P. concinnus</i>	Beaver	440	800	420	430, 400	500	480	350	480	420	190	700, 360	120
<i>P. confusus</i>	Washington	480	850	420	490	520	300	320	480	450	220	350	125
<i>P. crandallii</i> v. <i>atratus</i>	San Juan	420	490	390	500	450	370	320	280	400	190	350	120
<i>P. crandallii</i> v. <i>crandallii</i>	San Juan	420	340, 190	390	500	450	370, 340	310	280	380	190	350	115
<i>P. deustus</i> v. <i>pedicellatus</i>	Teton	420	850	420	430	550	340	320	550	340	230	370	130
<i>P. dolius</i> v. <i>dolius</i>	Millard	NM	710	400	400	490, 320	530, 300	320	480	450, 420	180	340	105
<i>P. dolius</i> v. <i>duchesnensis</i>	Duchesne	420	AD	420	400	500	340	320	480	410	180	360, 340	140, 120
<i>P. eriantherus</i> v. <i>cleburnei</i>	Daggett	420	850	420	410	450	480	300	500	490, 420	190	390, 360	140, 130
<i>P. flowersii</i>	Uintah	480	AD	420, 400	490, 430	470	300	350	520	420	220	360	125
<i>P. franklinii</i>	Iron	480	800	400	430	470	320, 300	350	520	420	240	380	125
<i>P. goodrichii</i>	Uintah	420	650	390	400	490	480	310	480	400	200	370, 350	135
<i>P. grahamii</i>	Uintah	420	850	400	390	470	530, 320	350	500	420	230	500, 370	120
<i>P. humilis</i> v. <i>brevifolius</i>	Cache	390	850	370	500	450	340, 320	320	480	500	220	280	115

Table 1 *Penstemon* taxa (with collection counties) utilized in the 12 marker analysis with respective marker sizes (Continued)

<i>P. humilis</i> v. <i>humilis</i>	Box Elder	420	850	390	410	520	330, 310	360	500	500, 470	220	350	120
<i>P. humilis</i> v. <i>obtusifolius</i>	Washington	420	800	390	520, 490	AD	330	340	480	470	200	350	120
<i>P. immanifestus</i>	Millard	480	710	420	490	380	300	320	480	410, 380	220	400, 360	120
<i>P. lentus</i> v. <i>albiflorus</i>	San Juan	NM	430	390	430	450	320	300	500	400	210	470, 370	140
<i>P. lentus</i> v. <i>lentus</i>	San Juan	480	850	400	430	470	300	310	500	410	210	400, 370	145
<i>P. linarioides</i> v. <i>sileri</i>	Washington	420	850	370	490, 390	470	330, 310	350	470	400	210	370	125
<i>P. marcusii</i>	Emery	390	800	450	370	490	310	340, 320	500	NM	200	390, 360	120
<i>P. moffatii</i>	Grand	390	800	420	390	490	330	340	480	430	290, 200	380, 350	140
<i>P. nanus</i>	Millard	480	800	420	390	470	280	320	470	NM	180	360	120
<i>P. ophianthus</i>	Sevier	520	850	420	370	900, 750	330, 310	310	480	420	190	AD	115
<i>P. pachyphyllus</i> v. <i>congestus</i>	Kane	480	850	400	430	470, 380	320	310	520	410	250	370	170
<i>P. pachyphyllus</i> v. <i>mucronatus</i>	Daggett	440	800	390, 370	430	500	320	300, 280	520, 500	430	220	350	120
<i>P. pachyphyllus</i> v. <i>pachyphyllus</i>	Duchesne	480	850	390	410	490	370, 330	340, 240	400, 190	500, 430	290, 230	380, 220	125
<i>P. palmeri</i> v. <i>palmeri</i>	Washington	440	850	400	500, 490	520, 490	330	310	500	430	210	380	125
<i>P. petiolatus</i>	Washington	420	1000	400	500, 490	500	330	300	480	420	210	380	145
<i>P. pinorum</i>	Washington	480	800	420	610	500	480	310	480	470	200	390	125
<i>P. procerus</i> v. <i>aberrans</i>	Garfield	440	1000, 850	450, 370	520	520	330	360	480	410	220	370	115
<i>P. procerus</i> v. <i>procerus</i>	Iron	420	850, 550	370	490	470	340, 310	360	470	470	220	340	120
<i>P. radicosus</i>	Daggett	420	AD	420	490	470	330, 310	310	500	450	200	360	125
<i>P. rydbergii</i> v. <i>aggregatus</i>	Box Elder	420	850	400	520	500	340	360	520	470	210	380	115
<i>P. rydbergii</i> v. <i>rydbergii</i>	Rich	420	710	400	520	500	370	320	500	470, 430	AD	390	115
<i>P. thompsoniae</i>	Kane	420	AD	370	500	450	340, 320	340	500	410	220	390, 370	130
<i>P. tusharensis</i>	Beaver	420	1300, 230	370	430	450	320	320	500, 300	410	230	340	120
<i>P. utahensis</i>	San Juan	480	410	420	430	490	300	310	500	410	220	370	125
<i>P. watsonii</i>	Sevier	420	AD	370	490	470	320	350	480	490	220	350	120
<i>P. whippleanus</i>	Iron	420	800	400	370	450	310	350	500	430	210	370	105
<i>P. yampaensis</i>	Daggett	570	710	400	430, 390	490	500, 320	310	480	410	260, 230	340	120
Subgenus <i>Saccanthera</i>													
<i>P. leonardii</i> v. <i>higginsii</i>	Washington	390	1300	420, 400	490, 430	550, 520	320	310	480	470	250	AD	125
<i>P. leonardii</i> v. <i>leonardii</i>	Utah	440	800	420	430	490	320	340, 320	480	500	240	370	120
<i>P. leonardii</i> v. <i>patricus</i>	Tooele	440	850	370	470	AD	370	310	550, 520	470	230	380	115
<i>P. platyphyllus</i>	Salt Lake	420	800	400	430	470	330	310	520	430	240	AD	135
<i>P. rostriflorus</i>	Washington	420	1100, 430	400	410	420	320, 300	290	500	490, 430	470	370	120
<i>P. sepalulus</i>	Utah	420	800	400	470	450	330	310	500	430	230	390	130

Table 1 *Penstemon* taxa (with collection counties) utilized in the 12 marker analysis with respective marker sizes (Continued)

Total unique molecular weight bands	9	18	6	10	14	12	12	13	12	11	17	11
Total pairs of dual molecular weight bands	6	7	7	16	11	28	14	7	8	4	20	7
Total monomorphic markers	85	80	86	76	80	65	78	86	81	88	69	86
Total NM	2	0	0	0	0	0	1	0	4	0	0	0
Total AD	0	6	0	1	2	0	0	0	0	1	4	0

¹ All counties are in Utah except Teton, Co. which is in Wyoming.

² Purchased = *P. davidsonii* and *P. fruticosus* were purchased from nurseries in Utah Co., Utah while *P. dissectus* was purchased from a nursery in Aiken Co., South Carolina.

³ NM = no marker.

⁴ AD = ambiguous data (usually multiple bands and or smearing).

site. Next, a non-labeled size exclusion step using Chroma Spin + TE-400 columns (Clontech Laboratories, Inc., Mountain View, CA) and magnetic biotin-streptavidin separation (Dynabeads M-280 Streptavidin, Invitrogen Life Science Corporation, Carlsbad, CA) was performed. Unique multiplex identifiers (MID) barcodes were added independently to each species using primers complementary to the adapter and cut sites (Table 2). Preliminary amplification was performed using 95°C for 1 min., 22 cycles of 95°C for 15 s, 65°C for 30 s, and 68°C for 2 min. PCR products were loaded into a 1.2% agarose Flashgel DNA Cassette (Lonza Corporation; Rockland, ME) to verify smearing and adequate amplification in preparation for pyrosequencing.

After the initial PCR, concentrations of each of the four species samples were determined fluorometrically using PicoGreen® dye (Invitrogen, Carlsbad, CA). Samples were then pooled using approximately equal molar concentrations of each species except for *P. cyananthus* (genome size = 1C = 893 Mbp), where the molar concentration was doubled to maintain a similar genomic representation compared to the other three species with smaller genome sizes (*P. dissectus*, 1C = 462 Mbp; *P. davidsonii*, 1C = 483 Mbp; and *P. fruticosus*, 1C = 476 Mbp; [5]). DNA fragments between 500–600 bp were selected following Maughan et al. [25]. Sequencing was performed by the Brigham Young University DNA Sequencing Center (Provo, UT) using a half 454-pyrosequencing plate, Roche-454 GS GLX instrument, and Titanium reagents (Brandford, CT).

Sequence assembly

Sequence data were sorted by species using their unique MID species barcode (Table 2) by means of the software package CLC Bio Workbench (v. 2.6.1; Katrinebjerg, Aarhus N, Denmark). Following sorting (Table 2), assemblies were performed using Roche's de novo assembler, Newbler (v. 2.6), which yields consensus sequences (contigs) of all individual reads, from each independent species, for use in subsequent analyses.

A full assembly (all individual reads of all four species pooled together) was performed by Newbler with "complex genome" parameter set and a trim file with MID barcodes specified; all other parameters were left to their defaults. For all subsequent species assemblies (all individual reads of one species), these same parameters were used with a few added conservative options selected: an expected depth of '10' (20 default), a minimum overlap length of '50' (40 default), and a minimum overlap identity of 95% (90% default).

Repeat element identification

Assembled sequences from all four species were masked for possible genome wide repetitive elements using a combination of RepeatModeler and RepeatMasker [32]. RepeatModeler is a de novo repeat element family identification and modeling algorithm that implements RECON [33] and RepeatScout [34]. RepeatModeler scanned all contigs from the four *Penstemon* species assemblies and produced a predicted repeat element library of predictive models to find repeat elements. Using this reference library, RepeatMasker then scanned the four species to filter out repetitive elements. Singletons were omitted from the analysis. To assess possible repetitive element biases with RepeatMasker when implementing a denovo library from RepeatModeler, we analyzed the GR-RSC data from *Arabidopsis* RILs (recombinant inbred lines) Ler-O and Col-4 from Maughan et al's. [35] study, compared to the *Arabidopsis* non-reduced genome downloaded from TAIR (The Arabidopsis Information Resource) [36].

Marker development, verification, and use

To identify SSRs, INDELS, and SNPs, we used software MISA and SNP_Finder_Plus (custom Perl-script), respectively [25,37,38]. RepeatMasker was used to identify and mask transposable elements. MISA parameters were set as follows: di-nucleotide motifs had a minimum of eight repeats, tri-nucleotide motifs had a minimum of six repeats, tetra-nucleotide motifs had a minimum of five repeats, and 100 bp was set as the interruption

Table 2 The four multiplex identifiers (MID) barcodes (adapter) primers used for the genomes of *Penstemon cyananthus*, *P. dissectus*, *P. davidsonii*, and *P. fruticosus*

Species	MID ID #	EcoR1 MID primer ¹	Bfa1 MID primer ²
<i>P. cyananthus</i>	MID 1	5'- ACGAGTGC GTGACTGCGTACCAATTC	5'- ACGAGTGC GTGATGAGTCCTGAGTA
<i>P. dissectus</i>	MID 2	5'- ACGCTCGACAGACTGCGTACCAATTC	5'- ACGCTCGACAGATGAGTCCTGAGTA
<i>P. davidsonii</i>	MID 3	5'- AGACGCACTCGACTGCGTACCAATTC	5'- AGACGCACTCGATGAGTCCTGAGTA
<i>P. fruticosus</i>	MID 4	5'- AGCACTGTAGGACTGCGTACCAATTC	5'- AGCACTGTAGGATGAGTCCTGAGTA

¹ The "AATTC" at the 3' end the primer was where adapters complement the enzyme *EcoR1* cut site and the preceding "C" is where the base was changed to avoid further enzymatic cleavage of the fragment.

² The "TA" at the 3' end of the primer was where adapters complement the enzyme *Bfa1* cut site and the preceding "G" is where the base was changed to avoid further enzymatic cleavage of the fragment.

(max difference between two purported SSR alleles). For the comparison of SSR frequency and repeat motifs across species, “unmasked” assembly files were used to remove bias caused by masking low complexity reads. The following parameters were used to define the heuristic thresholds for SNP_Finder_Plus: 8× minimum read depth for the SNP, 30% proportion of the reads representing the minor allele and 90% identity (an indication of homozygosity within a single species used in a dual-species assembly) required for each SNP locus. These parameters also helped compensate for sequencing and assembly errors, which allow greater confidence in calling base pair discrepancies as actual SNPs in the dual-species assemblies and the confident identification of heterozygosity in the individual assemblies. For both individual assemblies and dual species assemblies SNPs reported are those conforming to the aforementioned parameters.

All genomic sequences matching the above criteria were used for marker development. Primer3 v2.0 [39] was used to identify primers for amplifying these markers, with the following parameters: optimal primer size = 20 (range = 18–27); product size range = 100–500 base bp; T_m range = 50–60°C with 55°C optimum; and maximum polynucleotide = 3. Allowing PCR products greater than 200 bp greatly increased the possibility of INDELs in the PCR products.

The PCR (SSR/INDEL) markers were validated using the original four species as template DNA. Each 10 μ l PCR reaction had ~30 ng genomic DNA, 0.05 mM dNTPs, 0.1 mM cresol red, 1.0 μ l of 10X PCR buffer (Sigma-Aldrich, St. Louis, MO), 0.5 units of JumpStart™ Taq DNA Polymerase (Sigma-Aldrich, St. Louis, MO) and 0.5 μ M (each) of the forward and reverse primers. The thermal cycler (Mastercycler® Pro; Eppendorf International; Hamburg, Germany) was set as follows: 94°C for 30 s, 45 cycles of 92°C for 20 s, (primer specific annealing temperature)°C for 1 min. 30 s, 72°C for 2 min., and 72°C for 7 min. (final extension). Following PCR reactions, DNA was loaded into 3% Metaphor® agarose (Lonza Corporation; Rockland, ME) gels and run using a gel electrophoresis box at 100 V for 2 h. Optimal annealing temperatures for each SSR/INDEL marker were selected based on clarity of bands produced over varying annealing temperatures. Only SSR/INDEL markers with one or two reproducible bands are reported in the marker studies (Tables 1 and 3). The same conditions used for marker validation were used in the SSR/INDEL marker studies, except gel electrophoresis times were increased to 4 h at 100 V.

The gels were evaluated and scored as: 1 = marker present; 0 = marker absent based upon molecular weight. The results were then analyzed to assess the strength of hierarchical signal in these data using 10,000

replications of fast bootstrapping as implemented in PAUP* v. 4.0b10 [40].

Our interspecific SNP genotyping was accomplished using Fluidigm (Fluidigm Corp., South San Francisco, CA) nanofluidic Dynamic Array Integrated Fluidic Circuit (IFC) Chips [40] on the EP-1TM System (Fluidigm Corp., South San Francisco, CA) and competitive allele-specific PCR KASPar chemistry (KBioscience Ltd., Hoddesdon, UK). A 5 μ L sample mix, consisting of 2.25 μ L genomic DNA (20 ng μ L⁻¹), 2.5 μ L of 2x KBiosciences Allele Specific PCR (KASP) reagent Mix (KBioscience Ltd.), and 0.25 μ L of 20x GT sample loading reagent (Fluidigm Corp., South San Francisco, CA) was prepared for each DNA sample. Similarly, a 4 μ L 10x KASP Assay, containing 0.56 μ L of the KASP assay primer mix (allele specific primers at 12 μ M and the common reverse primer at 30 μ M), 2 μ L of 2x Assay Loading Reagent (Fluidigm Corp., South San Francisco, CA), and 1.44 μ L DNase-free water was prepared for each SNP assay.

The two assay mixes were added to the dynamic array chip, mixed, and then thermal cycled using an integrated fluidic circuit Controller HX and FC1 thermal cycler (Fluidigm Corp., South San Francisco, CA). The thermo cycler was set as follows: 70°C for 30 min; 25°C for 10 min for thermo mixing of components followed by hot-start Taq polymerase activation at 94°C 15 min then a touchdown amplification protocol consisting of 10 cycles for 94°C for 20 sec, 65°C for 1 min (decreasing 0.8°C per cycle), 26 cycles of 94°C for 20 sec, 57°C for 1 min, and then hold at 20°C for 30 sec. Five end-point fluorescent images of the chip were acquired using the EP-1TM imager (Fluidigm Corp., South San Francisco, CA), once after the initial touchdown cycles were complete and then after each additional run on “additional touchdown cycles.” The extra cycles were run four times, with an analysis of the chip after each run.

The determination of each SNP allele was based on a minimum of at least two of three SNP genotyping experiments. The primers were then analyzed for functionality using the results from each of the five stops for each chip, which were compared to determine the most accurate call. Functionality was determined by number of calls verses no calls, and consistency.

Cross species sequencing verification

To evaluate the DNA sequence homology and polymorphism type (SSR or INDEL) at specific marker amplicons (Table 1) across the *Penstemon* genus, DNA samples from each of five species (*P. cyananthus*, *P. davidsonii*, *P. dissectus*, *P. fruticosus*, and *P. pachyphyllus*) were amplified and Sanger sequenced. We accomplished the PCR amplification using Qiagen HotStarTaq Plus Master Mix (Valencia, CA, USA) according to the manufacturer’s recommendations. The amplification protocol consisted

Table 3 Summary of marker characteristics including the primary SSR motif identified in the original GR-RSC (genome reduction using restriction site conservation) sequence, primer sequences, EFL (expected fragment length), total bands, and fragment sizes

Marker name ¹	Primary motif	Forward primer (5'-3') Reverse primer (5'-3')	GenBank accession ID	EFL	Total unique bands	Fragment size			
						<i>P. cyananthus</i>	<i>P. davidsonii</i>	<i>P. dissectus</i>	<i>P. fruticosus</i>
PS003 ^(di,f)	(AT) ₈	TGCCTCTGTCTTTACATTCCAA CATGAAGCACTGCAAATCCA	JQ966997	217	3	360	260	250	260
PS004 ^(da,f)	(ATT) ₆	TGTTTCAATTGCTGTCCACAT TTGTCTGTCCAAACGGTAGGT	JQ951613	476	3	420	460	440	460
PS005 ^(c,di,f)	(GAA) ₆	GCCCAACTTCCGTAATTGAA AACTGCTTGCCACTCGACTC	JQ966998	303	3	260, 300	260	280	280
PS009 ^(c,da,f)	(TGA) ₆	ACCTCGAATTGACGGTCC TTCTGAGGAGAAACCAAGGG	JQ966999	466	4	370, 650	540	650	600
PS011 ^(da,f)	(GA) ₈	AAGTGCACACTGGATGTCTT GCAGCTTCAGCTCCAGAAAT	JQ951614	435	2	860	500	860	500
PS012 ^(c)	(TA) ₈	TCCATATTGTAACCAACAATGACTG TGAATGGCAAACCGTAATCA	JQ951615	402	3	400	360	370	360
PS013 ^(f)	(TA) ₈	GAAGAATTGATTTAAACAAGATGCAA TCAGTACGTGAGAAACTTGATCAATAA	JQ967000	399	2	400	650	650	400
PS014 ^(c)	(TGA) ₆	CGATTTGGTATAGTTGGATTACGA CCTTCATCACCCGGTACTTG	JQ951616	409	3	410	370	380	410
PS015 ^(di)	(TCG) ₆	GCCGAGTTTCAAGAAAGCAA AATTACGACCTGCCACGC	JQ967001	409	2	490	500	490	490
PS016 ^(c,di)	(CT) ₈	CATGGCCCTTCTTCACACT GACGCGGTTGGCTATACAGT	JQ967002	447	3	NM ²	1,100	1,060	1,030
PS017 ^(da,di)	(AG) ₉	GAAGGCTTAGCATAAATCCTCAAA ATTAGGCTCCCACGAACAAA	JQ951617	455	2	750	700	750	700
PS019 ^(c,di)	(AG) ₈	AATCCCACAGCCATACAAA TGAATTGAGTCTATACCCTATTCAA	JQ967003	473	1	380	380	380	380
PS021 ^(f)	(CT) ₈	CTTTAGCTTAGCTGGAATACACGTT AGATTCTTGATCACAGTTCAATTA	JQ967004	386	3	350	450	450	420
PS023 ^(da)	(AG) ₈	GCTGGAGAATAACATGGCG CCATCTTGCAAGTCCATACG	JQ967005	469	4	310	480	120, 740	480
PS024 ^(da,f)	(CTG) ₆	CTTCTTGCCCTGTGCTCT CCACCACCAACAACAAC	JQ967006	403	2	430	430	400	430
PS025 ^(c,di)	(TC) ₉	GCACATGAATGAAGGAATGC ACGATCTGTGAAGGAACCCA	JQ967007	440	3	440	410	440	400

Table 3 Summary of marker characteristics including the primary SSR motif identified in the original GR-RSC (genome reduction using restriction site conservation) sequence, primer sequences, EFL (expected fragment length), total bands, and fragment sizes (Continued)

PS026 ^(c,da,di,f)	(CTT) ₆	<u>ACTTAATAATGCCTCCTTGTGTCA</u> TTCCGCAACGTTGTATTTGA	JQ967008	465	1	460	NM	460	460
PS028 ^(di)	(AC) ₉	<u>GGGAGGCAGGTAACAACAAA</u> TACCTCTGCCGAACTGGATT	JQ967009	316	4	950	400, 460	320	400, 460
PS029 ^(di)	(TA) ₈	<u>ACCAAGTTGTTGGATGTTTGG</u> GGTTTGAATGAGACTTAGAAGGA	JQ967010	440	3	840	500	500	420
PS032 ^(c,di)	(GT) ₉	<u>ACAAAGTCTCCTCAATCGCC</u> GCATGTACCGTGACACACT	JQ951618	328	2	370	370	370	340
PS034 ^(c)	(AC) ₉	<u>CCAAACAATCAAACAGCACTC</u> CATGCGAATCAGTGTGCTAA	JQ951619	322	5	310, 340	320, 950	320	360
PS035 ^(da,f)	(TC) ₉	<u>TTGCACAGCTACTTTGGCAT</u> ATCTGTCCAAGGCATGGAAT	JQ951620	486	3	630	520	920	520
PS036 ^(c,di)	(TA) ₈	<u>TTCCTAATTTGGTAGCTGCAATC</u> TCCGAGGAACTATTGCCATT	JQ967011	405	3	770	770, 820	590	770
PS038 ^(c,da)	(TA) ₈	<u>GTAATTACTTCGGCAGTTTGTAATTT</u> GGTGCGACCTAATTACGTTTCTAT	JQ967012	100	1	NM	100	100	NM
PS040 ^(da)	(CA) ₉	<u>TAAAGAGGCTTAAGCGCGG</u> ACCTGAAGAGCTGCGGAGTA	JQ967013	399	3	380	390	410	390
PS041 ^(c,da,di,f)	(AT) ₈	<u>TTCCGCAAGAGAAGAGCAT</u> CTTGTCACGATTCCATTGT	JQ967014	249	3	270	670	270	240
PS045 ^(c,da)	(CT) ₈	<u>GCCACATACATGAAACGTGAA</u> CGAACTCTTGTGTTTCTCCC	JQ967015	366	4	460	NM	440	120, 400
PS047 ^(c,di,f)	(AC) ₈	<u>ACACGACATCGTTTCAGCAA</u> GCGTATGGAGAGATTTGGGA	JQ967016	428	3	470, 510	440	470	470
PS048 ^(c,di)	(CA) ₉	<u>GCATTAGATGCCGAAATATCTACAA</u> TGCCTGTAGGTTGATTTCTTTT	JQ951621	436	3	420	440	380	420
PS049 ^(c,da,di,f)	(AG) ₈	<u>CCCATCAATAAAGAAAGAAAGAAAGA</u> GGTGAACCCTGTCCTAAACC	JQ967017	436	2	460	460	1,000	460
PS050 ^(c,di)	(AT) ₉	<u>GTGTAACCTCTGAACAAGTTTACTGAA</u> TGCACTGAGCCATGCTATTC	JQ967018	434	2	480	460	480	460
PS051 ^(c,di,f)	(TG) ₈	<u>TGTAACACGACAATTTAACTCTTTCA</u> CGAGAACTCTTCCGAGAACC	JQ967019	352	1	280	NM	280	280

Table 3 Summary of marker characteristics including the primary SSR motif identified in the original GR-RSC (genome reduction using restriction site conservation) sequence, primer sequences, EFL (expected fragment length), total bands, and fragment sizes (Continued)

PS052 ^(c,da,di,f)	(AC) ₉	<u>CGCGGTCAATCTTGAATCT</u> TGACTTCTCTCTCTCTCACAC	JQ951622	206	1	220	220	220	220
PS053 ^(di)	(AC) ₈	<u>AATCATAGTCTCGAGCGCGT</u> GAGATAAATTAGATCAGCGCATCA	JQ951623	410	3	160, 320	320	320, 450	320
PS054 ^(c,da,f)	(GA) ₈	<u>TCGTTAAGCAATCTCGGAGC</u> TCGACTGGAGAGCAAAGCA	JQ967020	192	3	200	200	180	190
PS055 ^(f)	(AG) ₈	<u>TGTGGTCCGGTTCATAAAC</u> TTTGTCTCCCTAATATGTGTGATGAT	JQ967021	412	4	960	500	1,040	470
PS056 ^(da,di)	(TG) ₈	<u>CATGTTTCAGGATTGGGCTT</u> CGGTTACACACAGTTGTTGA	JQ967022	319	4	690	450	230	340
PS057 ^(da,f)	(AT) ₈	<u>TGCCTAATGGACCTGATCCT</u> CCCAATTGTTGAAGAAAGAACA	JQ967023	402	2	570	440	570	440
PS058 ^(da)	(AT) ₉	<u>GTGCAACCAATGCAACTAATTC</u> TCTCTCATTCCAATGATTCTCA	JQ967024	469	1	NM	720	NM	720
PS059 ^(di)	(CT) ₈	<u>CATCAATTGACACACAAGCAGA</u> TCGAATCTTAAAGAAACACATCCA	JQ967025	312	2	930	340	340	340
PS060 ^(c,di)	(AC) ₉	<u>CCATGAGAAGTAGATGACTGGGA</u> TTGTAATTATGATTAACCTCCCTCGTT	JQ967026	484	2	560	560	560	540
PS061 ^(da,f)	(TA) ₈	<u>CGACCAATCATCAACCAACA</u> GACGGGCAGATAATTGGAA	JQ967027	453	3	480	480, 530	450, 480	NM
PS062 ^(c,di)	(TA) ₉	<u>TGGAGAGGGTACGAAAGTGC</u> CAACGATCGATTATTAGCACCA	JQ967028	320	2	350	290	350	290
PS064 ^(c,da,f)	(AG) ₈	<u>ATGGATGCCCTATGGGTACA</u> TGAAATGGAGGGAGTAATATAAACAA	JQ967029	437	4	490	500, 680	470	470
PS066 ^(di)	(GA) ₉	<u>CAAGGATGCAGGCTCTCATT</u> CTCTGCTCGTCGTAGTGCAA	JQ967030	434	2	250	480	250, 480	480
PS068 ^(c,da,di,f)	(GA) ₈	<u>TTTGGGATGCATTCTCCAC</u> TCAAAGTGACATCTCCAACAAA	JQ967031	463	2	500	500	480	480
PS069 ^(di)	(GT) ₈	<u>CATTGGGTCAGATTTGGCTT</u> GCTTTCAGTTTGTATATTTGTGCC	JQ967032	309	4	220	210	390	350
PS071 ^(c,di)	(AT) ₈	<u>AAGATGGCCCTGATCTGTTG</u> TTCGTGGGAGTTGCAAATTA	JQ967033	446	1	NM	NM	490	NM

Table 3 Summary of marker characteristics including the primary SSR motif identified in the original GR-RSC (genome reduction using restriction site conservation) sequence, primer sequences, EFL (expected fragment length), total bands, and fragment sizes (Continued)

PS074 ^(c,da,di,f)	(AAG) ₆	<u>AGAAATCTCGCTCCACGA</u> CGACAACCTTAGTGATCGCTTT	JQ967034	168	1	170	170	170	170
PS075 ^(c,da,di,f)	(TA) ₈	<u>CACCACTTTCGCAGCATTTA</u> CAAATTACATTATTGTATGGAAACACG	JQ951624	120	2	160	140	140	140
PS076 ^(c,di)	(GTG) ₆	<u>CTGACAGCAACATGAACATGAA</u> CAATCTTGCCAATTTCCCA	JQ967035	161	1	170	170	170	170

¹ Parentheses indicates the species possessing sequence from which primers were designed (c = *P. cyananthus*, da = *P. davidsonii*, di = *P. dissectus*, f = *P. fruticosus*).

² NM = no marker.

of an initial denaturation step of 5 min at 95°C, followed by 40 cycles of amplification consisting of 30 sec denaturation at 94°C, 30 sec for primer annealing at 55°C and 1 min of extension at 72°C. PCR products were separated on 1% agarose gels run in 0.5X TBE and visualized by ethidium bromide staining and UV transillumination. PCR products were purified using a standard ExoSAP (Exonuclease I/Shrimp Alkaline Phosphatase) protocol and sequenced directly as PCR products. DNA sequencing was performed at the Brigham Young University DNA Sequencing Center (Provo, UT, USA) using standard ABI Prism Taq dye-terminator cycle-sequencing methodology. DNA sequences were analyzed, assembled and aligned using Geneious software (Biomatters, Auckland, New Zealand).

Gene ontology

We used BLASTX [41] on assembled sequences of all four species to compare with the GenBank refseq-protein database [42] with a threshold of $<1.0e^{-15}$. Blast2GO (v2.4.2) was used to map the blast hits and annotate them to putative cellular components, biological processes, and molecular functions found in the blast database [43]. For species comparisons, the GO level 3 was used for cellular components and level 2 was used for both biological processes and molecular functions.

Assembled sequences of all four species were also compared to all available *Antirrhinum* and *Mimulus* (genera more or less related to *Penstemon*) genes on GenBank (downloaded 23 June 2011). Comparisons were made using BLASTN [41] with an e-value threshold of $<1.0e^{-13}$.

Results and discussion

Genome reduction, pyrosequencing and species assemblies

Given that a full 454 pyrosequencing plate using Titanium reagents is capable of producing 1.3 million reads averaging ~400 bp each [25], we expected a half plate to produce approximately 250 Mbp from 650,000 reads. Our reaction produced 287 Mbp from 733,413 reads, 20% more than expected, with an average read length of 392 bp. In total, 93.8, 46.4, 48.8, and 53.3 Mbp were sequenced from *P. cyananthus*, *P. dissectus*, *P. davidsonii* and *P. fruticosus*, respectively, closely resembling the 2:1:1:1 ratio of DNA pooled from each species for sequencing (Table 4). Likewise, from our de novo assemblies, we identified nearly twice as many contigs, 9,714 in *P. cyananthus* than the 4,777 found in *P. fruticosus*, for example, which was expected because we sequenced approximately twice as much DNA from *P. cyananthus* than the other three species. There was 0.6% of *P. cyananthus* genome represented compared to 0.5% average coverage of the other three species (Table 4); thus,

essentially an equal genome representation from each species was realized using the GR-RSC technique by pooling approximately equal genome molar concentrations in the sequencing reaction. The contigs of this study have been deposited at DDBJ/EMBL/GenBank as a Whole Genome Shotgun project under the accessions AKKG00000000 (*P. cyananthus*), AKKH00000000 (*P. dissectus*), AKKI00000000 (*P. davidsonii*), and AKKJ00000000 (*P. fruticosus*). The version described in this paper is the first version for each accession, XXXX01000000.

DNA sequences produced by the GR-RSC technique represent a broad sample of the genome. With this sample, we can begin to estimate genome-wide characteristics, such as GC content, frequency of repeat elements, and so forth. From the genome reduction, GC content was measured to be 36.4%, 34.5%, 35.3%, and 35.15% for *P. cyananthus*, *P. dissectus*, *P. davidsonii* and *P. fruticosus*, respectively (Table 4), matching the average 35% GC content reported for dicots [44]. Using the dicot average GC content a priori, we estimated a theoretical frequency of the *Bfa*I and *Eco*RI recognition sites. The theoretical GC content in combination with estimated genome sizes of the four species [5] suggested the GR-RSC should have rendered a 104 fold reduction of the genome of each species. With a reduced genome of these species, the 650,000 reads that were sequence suggest an average of 11× coverage; however the observed read depth was 8.5×, 22.7% less than expected (Table 4). This lighter coverage is partly due to the lower than expected specificity of reads. An average of 48.2% of the reads were matched to contigs with the other 51.8% either too short or lacking in homology to successfully match to a contig (Table 4).

The full assembly of all four *Penstemon*, using the Newbler de novo assembler, produced a total of 44,966 contigs, representing 16.4 Mbp, or 5.7% of our total sequence. In the individual species assemblies of *P. cyananthus*, *P. dissectus*, *P. davidsonii*, and *P. fruticosus*, a total of 9,714, 5,364, 4,882, and 4,777 contigs were created representing 4.6, 2.6, 2.4, and 2.3 Mbp of assembled bases respectively. These contigs represent, on average, 0.5% of the total genomes being sequenced (Table 4).

Marker analysis

We utilized assembly contigs from genomic sequence of all four species with “masked” multiple repeats, such as transposons, to identify SSRs. *Penstemon cyananthus*, *P. dissectus*, *P. davidsonii*, and *P. fruticosus* had 97, 113, 49, and 58 SSRs identified respectively (Table 5). There were more SSRs identified in *P. dissectus* than *P. cyananthus*, which has a 1.9 times larger genome and a higher representation of sequence than *P. dissectus* (Table 5). This inverse relationship between genome size and SSRs content agrees with observations in other plant genomes [45]. Some SSRs were found as putative homologs in

Table 4 Summary data from 454-pyrosequencing and Newbler de-novo assembly (v.2.0.01) of *Penstemon cyananthus*, *P. dissectus*, *P. davidsonii*, and *P. fruticosus*

Assembly	Genome size (Mbp) ¹	GC content	Reads	Bases ²	% Reads assembled	% Bases assembled	Contigs created	Bases in assembly	% Genome represented	Average coverage	Bases shared between assemblies
<i>P. cyananthus</i>	893	36.4%	199,329	87,753,792	53.1%	50.0%	9,714	4,623,755	0.5%	7.7X	
<i>P. dissectus</i>	462	34.5%	98,868	43,304,550	52.8%	50.9%	5,364	2,629,819	0.6%	8.2X	
<i>P. davidsonii</i>	483	35.3%	103,963	45,599,742	45.8%	43.5%	4,882	2,376,141	0.5%	9.1X	
<i>P. fruticosus</i>	476	35.2%	113,146	49,786,980	41.0%	38.8%	4,777	2,322,606	0.5%	8.9X	
<i>P. cyananthus</i> × <i>P. dissectus</i>			298,197	131,058,342	53.0%	50.1%	14,523	6,915,079			338,495
<i>P. cyananthus</i> × <i>P. davidsonii</i>			303,292	133,353,534	49.9%	46.9%	14,254	6,757,023			242,873
<i>P. cyananthus</i> × <i>P. fruticosus</i>			312,475	137,540,772	47.8%	44.9%	14,134	6,705,536			240,825
<i>P. dissectus</i> × <i>P. davidsonii</i>			202,831	88,904,292	48.3%	46.1%	10,053	4,855,491			150,469
<i>P. dissectus</i> × <i>P. fruticosus</i>			212,014	93,091,530	45.7%	43.5%	9,873	4,774,539			177,886
<i>P. davidsonii</i> × <i>P. fruticosus</i>			217,109	95,386,722	44.0%	41.7%	9,184	4,442,194			256,553
Full <i>Penstemon</i> Assembly			730,215	265,987,500	47.9%	46.4%	44,966	16,363,589			

¹ The diploid (2n = 2x = 16) genome size as reported by Broderick et al. [5].

² Bases denotes the total number of bases used to create the assembly and not the total number of bases sequenced.

Table 5 Data obtained from MISA (SSR), Blast2GO (GO) and RepeatMasker (RM)

		Penstemon species				
		<i>P. cyananthus</i>	<i>P. dissectus</i>	<i>P. davidsonii</i>	<i>P. fruticosus</i>	
SSR	Total SSRs ¹	97	113	49	58	
	SSRs/Assembly Length	2.1E-05 (~1/48000)	4.3E-05 (~1/23000)	2.1E-05 (~1/48000)	2.5E-05 (~1/40000)	
	Repeat Type	di-	44.3%	40.7%	46.9%	48.3%
		tri-	45.4%	43.4%	44.9%	41.4%
tetra-		10.3%	15.9%	8.2%	10.3%	
GO	Contigs Analyzed	9,714	5,364	4,882	4,777	
	Blast Hits Found ²	1,899	1,125	1,121	1,091	
	Annotated Hits	1,430	844	388	826	
	% Blast Hits	19.5%	21.0%	23.0%	22.8%	
	% Annotated	14.7%	15.7%	7.9%	17.3%	
RM	Masked Repeat Elements	28.5%	16.8%	17.4%	16.1%	
	Retroelements (LTR)	7.8%	3.0%	4.9%	4.6%	
	DNA Transposons	0.3%	0.9%	1.0%	1.0%	
	Other Repeats ³	20.4%	12.9%	11.6%	10.5%	

¹ For MISA, "unmasked" individual species assemblies were used.

² Sequence compared to the GenBank refseq-protein database e-value threshold of <1.0e⁻¹⁵.

³ Other Repeats includes: lines, unclassified repeats, satellites, simple repeats, and low complexity sequence.

multiple species; after eliminating redundancies, we tallied 133 unique SSRs (Table 3). We generated primer pairs surrounding 77 of these SSRs large enough to potentially capture INDELs, of these, 51 produced 1 or 2 reproducible bands with no or few faint superfluous bands. From those 51, there was an overall success rate of 94% with 42 (82%) being polymorphic between the four species (Table 3).

To assess the possibility of utilizing these markers in interspecific plant improvement studies, 12 of the 51 SSR/INDEL markers (Table 3) were tested on 93 mostly xeric *Penstemon* taxa (72 species [Table 1]) representing five of six subgenera recognized in the genus [14]. The overall success rate of the markers was 98% with 100% being polymorphic across the 93 taxa. Without sequencing each band and/or doing inheritance studies on each marker it is not possible to clearly determine if a polymorphism of a given marker is a variant of an allele or a new locus. However, we did amplify and sequence the amplicon produced at 11 of these markers in five *Penstemon* species

(*P. cyananthus*, *P. davidsonii*, *P. dissectus*, *P. fruticosus*, and *P. pachyphyllus*). *P. pachyphyllus* var. *pachyphyllus* represents the largest subgenus (*Penstemon*) in the genus. These five species represented four of the presently classified six *Penstemon* subgenera. Of the 55 attempted sequences, 60% produced high quality sequences results which could be compared to the original 454 contigs containing the microsatellites. Using BLASTN (v2.2.25+) [41] we found that 33 sequences matched the respective microsatellite-containing contigs from which the SSR/INDEL markers were derived with an e-value of no more than 1.0e⁻³⁶. An example of the types of polymorphism (SSRs and INDEL) found at these loci across the various species is represented graphically for the marker PS035 (Figure 1). For 22 (40%) of the 55 attempted sequences, we were unable to obtain high quality sequence information. In the majority of these cases (94%) the lack of high quality data was clearly due to the amplification of multiple amplicons (seen as multiple bands in gel electrophoresis)

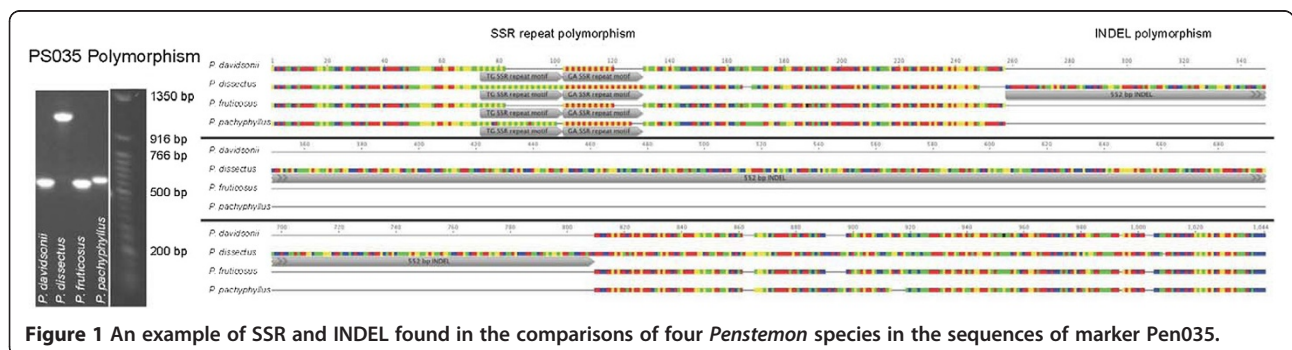


Figure 1 An example of SSR and INDEL found in the comparisons of four *Penstemon* species in the sequences of marker Pen035.

which impeded the sequencing of the PCR reaction. The source of the multiple amplicons may be from heterozygosity at the locus or from the amplification of paralogous loci.

Both the sequence data (Figure 1) as well as the marker size data (Tables 1 and 3) are clear evidence of sequence conservation, and probable homologous loci, in many of the SSR/INDEL markers. Marker PS012, the apparent most conserved marker, had six unique molecular weight bands and was present in all 93 taxa. The marker with the most diversity in its molecular weights was PS011 which had 18 variants and was not readable in seven of the 93 taxa. Of the 1,116 possible marker \times taxa interactions, 22 (2.0%) did not produce reliable data. Seven of those 22 (0.5%) were absent of any product with the remaining 15 producing multiple bands (reported as ambiguous data). Clearly readable double bands were found in 135 of the 1,116 (12.1%) marker \times taxa interactions (Table 1).

Our data suggest a high degree of sequence conservation across the genus, favoring the present hypothesis of a recent and rather rapid evolutionary radiation of the genus [13,14]. Furthermore, our data agree with Morgante et al. [45] who suggest that SSR presence in non-coding sequence are highly conserved and predate recent genome expansions of many plants. Some of our markers differed in length by as much as 570 bp (Tables 1 and 3) suggesting the presence of INDELs and possibly additional SSRs (Table 3). We confirmed the presence of INDELs in the sample of 11 markers which we sequenced (Figure 1). In some instances, these large fragment length variances may be amplifying a different locus, which is a recognized concern when using SSR based markers above the species level [46,47]. INDELs are useful as PCR based markers since they, like SSRs, are codominant and abundant in the genome and are commonly used in genetic mapping [26]. By combining the SSRs we identified in the source sequence for each of these markers with potential INDELs, alleles will be easily and inexpensively identified by gel electrophoresis.

To assess the possibility of phylogenetic (i.e., hierarchical) structure of the variation within these SSR/INDEL data at the broad taxonomic scale of our survey, we analyzed the 12 marker data set (Table 1) with PAUP*. Fast bootstrapping recovered a largely unresolved topology suggesting rampant homoplasy. Or one or more of these markers represent more than one locus. These results are similar to what others have reported about SSR type markers. SSRs have demonstrated utility for population and intraspecific relationships, such as cultivar differentiation; however, they can be problematic when used to reconstruct relationships above the species level where length differences are expected to poorly reflect homology [47,48]. Nonetheless, with over 96% of these SSR/INDEL regions being

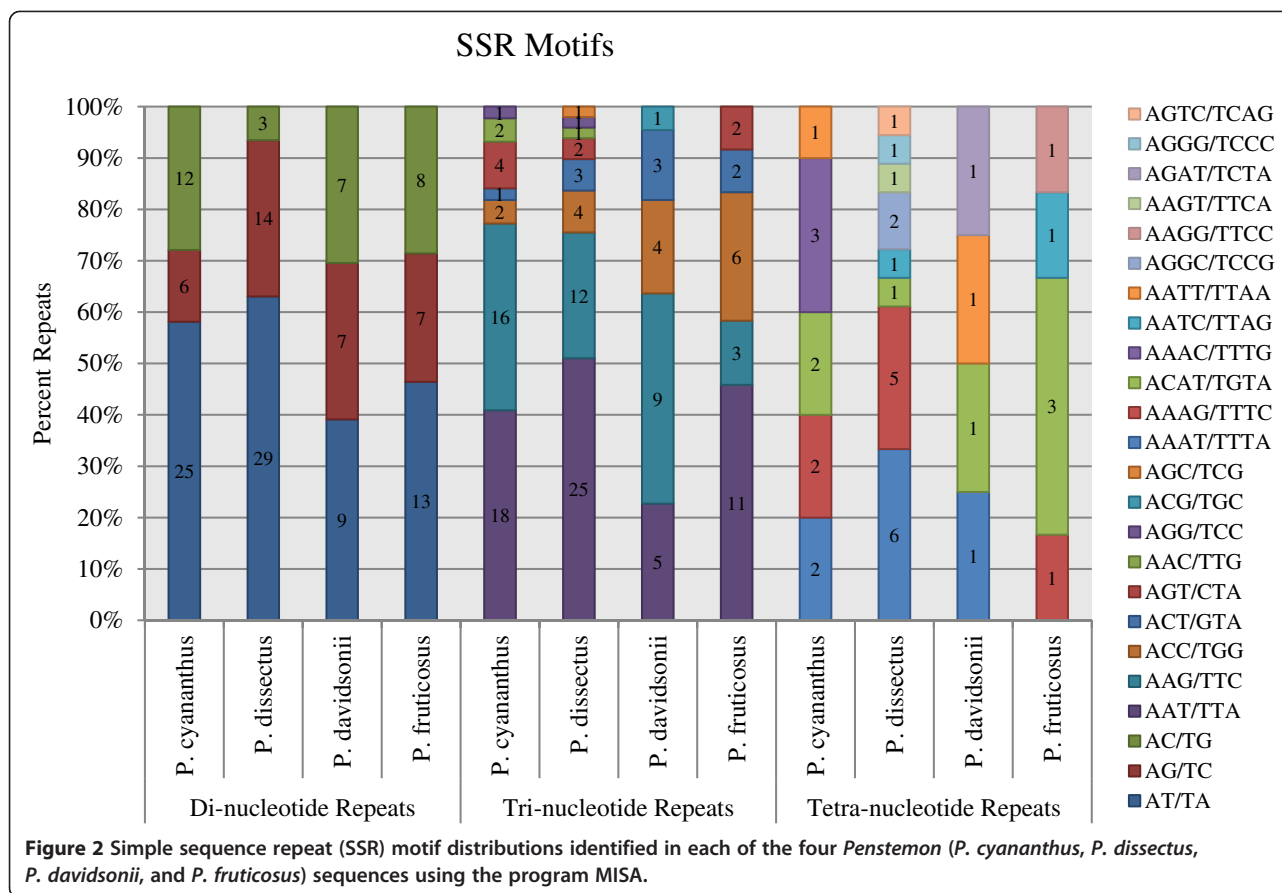
conserved across *Penstemon*, these markers have potential for studies of interspecific hybridization and cultivar development.

Interspecific *Penstemon* breeding is complex [7,11,15,49]; thus, having a set of inexpensive and easily used SSR/INDEL markers, which amplify across the genus, will have utility in understanding the results of some wide crosses. Empirical studies of various *Penstemon* interspecific crosses have ranged from a clearly recognizable intermediate phenotype of the two parents, to the F¹ essentially mimicking one of the two parents, usually mirroring the female parent. Furthermore, in some instances the F²'s and additional generations continue to mimic the female parent to the point that Viehmeyer [49] began to question if apomixis was involved. An example of this phenomenon was a 'Flathead Lake' \times *P. cobaea* interspecific cross. It was not until the hybrid progeny of this cross was crossed with other interspecific hybrids when the progeny gave a much wider range of phenotypes [49]. A probable reason for this phenomenon is "unequal segregation" which has been described in other wide crosses [50,51]. Thus through the use of these SSR/INDEL markers, regions of the genome can be identified which are unusual genotypic combinations, for that specific cross, and selections made accordingly [51-54]. Thus increasing the number of unique genotype/phenotype plants to be grown out to maturity from thousands of seedlings. Since many *Penstemon* require two years before their first anthesis, using markers to identify the greatest number of genotypic diverse plants is potentially very useful in the breeding of this crop.

Beyond amplification ability, we also assessed the composition and trends of all SSRs identified. On average, adenine and thymine rich repeat motifs were the most common repeat type in the di-, tri-, and tetra-nucleotide repeat motifs (Figure 2). In general, AT motifs are the most common motifs in noncoding regions of most plant genomes [45]. More variation was observed in the repeat motifs in the tetra-nucleotide repeats across the four species. Even closely related *P. fruticosus* and *P. davidsonii* had completely distinct tetra-nucleotide repeat motifs (Figure 2). This is likely due, in part, to the rarity of the motifs and high number of possible nucleotide combinations. Several studies have found that the hypothetical origins of some SSRs are retrotransposition events [48,55,56] and, as such, may be useful in developing part of a unique "fingerprint" for a given species.

SNP analysis

Using our SNP discovery parameters of an 8 \times minimum coverage, and 30% representation of the minor allele, we identified an average of one SNP per 2,890 bp across the four species ranging from *P. cyananthus* (1/1,855 bp) to



P. fruticosus (1/3,777 bp). The three species with similar genome sizes all had similar SNP frequencies (Table 6). As reported in other plant species [57,58], we found that the frequencies of bp transitions (A↔G or C↔T) were more common compared to transversions (A↔T, A↔C, G↔C, G↔T) in *Penstemon* by an average factor of 1.5

(Table 6). This is close to the 1.4 factor in *Arabidopsis* [35]. In the dual species assemblies, using the same parameters and a 90% SNP identity, the average transition to transversion mutation rate was lower at 1.2 (Table 6).

In the dual species assembly, we found an average of 1 SNP/97 bp between homologous sequence assemblies

Table 6 SNP type and distributions along with SNP comparisons of sequences found within and between species (homologous sequence comparisons) using SNP_Finder_Plus (8X min. coverage, 30% min. minor allele, 90% min. identity)

Species assembly	SNP	Average coverage	SNPs/assembly length ¹	SNP distribution						
				A↔C	A↔G	A↔T	C↔G	C↔T	G↔T	
<i>P. cyananthus</i>	2,493	16.4	0.000539 (~1/1855 bp)	10.7%	29.5%	13.9%	4.3%	30.2%	9.5%	
<i>P. dissectus</i>	737	14.3	0.000280 (1/3568 bp)	9.8%	30.7%	15.6%	4.6%	27.4%	9.8%	
<i>P. davidsonii</i>	713	14.4	0.000300 (~1/3333 bp)	11.9%	26.4%	15.2%	3.9%	28.3%	11.8%	
<i>P. fruticosus</i>	615	12.4	0.000265 (~1/3777 bp)	11.7%	27.2%	17.9%	4.2%	25.4%	12.0%	
Homologous sequence comparisons										
<i>P. cyananthus</i> × <i>P. dissectus</i>	3,253	10.6	0.009610 (~1/104 bp)	11.7%	27.5%	16.0%	7.1%	27.1%	10.6%	
<i>P. cyananthus</i> × <i>P. davidsonii</i>	1,958	10.7	0.008062 (~1/124 bp)	11.1%	27.6%	15.8%	7.1%	28.5%	9.9%	
<i>P. cyananthus</i> × <i>P. fruticosus</i>	2,015	10.6	0.008367 (~1/119 bp)	10.6%	27.2%	16.7%	6.8%	28.7%	10.1%	
<i>P. dissectus</i> × <i>P. davidsonii</i>	2,348	10.8	0.015605 (~1/64 bp)	12.6%	26.7%	15.5%	7.5%	27.3%	10.4%	
<i>P. dissectus</i> × <i>P. fruticosus</i>	2,133	10.0	0.011991 (~1/83 bp)	12.0%	26.4%	16.5%	7.6%	27.2%	10.4%	
<i>P. davidsonii</i> × <i>P. fruticosus</i>	2,156	10.1	0.008404 (~1/119 bp)	12.8%	28.2%	14.5%	7.2%	27.2%	10.1%	

¹ Assembly length is bases shared between assemblies (see Table 4).

of any two of the four species. The frequency of SNPs between homologous sequences of *P. dissectus* and *P. davidsonii* was the highest at 1/64 bp, with the lowest being between *P. cyananthus* and *P. davidsonii* at 1/119 bp. These results are in line with previous molecular based studies [5,14]. *Penstemon davidsonii* and *P. fruticosus* both belong to subgenus *Dasanthera*, while *P. cyananthus* and either *P. davidsonii* or *P. fruticosus* homologous sequences had fewer SNPs at 1/124 and 1/119, respectively. All homologous sequence comparison involving *P. dissectus* had the highest density of SNPs (Table 6) suggesting that *P. dissectus* is the most evolutionary distant of the four species.

It is important, for a high degree of confidence in the results, when the “SNP identity” parameter in SNP_Finder_Plus to have two or more independent samples from the same species. This requirement was not met for each of the species assemblies, thus, introducing a weakness in our interspecific SNP comparisons. Although with the parameters of a minimum 8× coverage and minor allele frequency set at least 30%, a putative SNP must be present in at least three of the eight contig reads, thus providing some protection from mislabeling a sequencing and/or assembly error as a SNP. Furthermore, when doing across species comparisons the average SNP coverage was actually 14.4× (Table 6). Therefore, on average, five identical putative SNPs represented the minor allele.

To understand the viability of our interspecific SNP as markers, we utilized the 1,958 *P. davidsonii* × *P. cyananthus* and 2,348 *P. davidsonii* × *P. dissectus* SNPs identified in the 14,254 and 10,053 respective homologous contig pairings (Tables 4 and 6). After removing contigs absent of identifiable SNPs, putative repetitive elements, and nonnuclear plastid DNA, 431 remained. Of these contigs, 99 were homologous across all three species (*P. cyananthus*, *P. davidsonii* and *P. dissectus*) another 164 were only in the *P. davidsonii* × *P. cyananthus* comparisons while the remaining 168 were in the *P. davidsonii* × *P. dissectus* contigs. Of those 431 contigs, we selected the first 192 for SNP marker development, 86 from each of the species comparisons. These contigs were utilized for competitive allele-specific PCR SNP primer design using PrimerPicker (KBioscience Ltd., Hoddesdon, UK).

Of the 192 SNP markers tested, using KASPar genotyping chemistry, 75 (39%) of produced consistent results for *P. cyananthus*, *P. davidsonii*, *P. dissectus*, and *P. fruticosus* (Table 7). All 75 SNP markers indicated polymorphisms between *P. cyananthus*, *P. davidsonii*, and *P. dissectus*, where only 16 (21% of the 75) produced results in *P. fruticosus* (Table 7). These results suggest that it is possible to develop intrageneric SNPs for *Penstemon*. However, it is unclear as to how viable these markers will be for use

across all the species of the genome since only 21% worked on all the species used in this GR-RSC study.

Repetitive elements

We identified 28.5%, 16.8%, 17.4% and 16.1% of the respective sequence from *P. cyananthus*, *P. dissectus*, *P. davidsonii*, and *P. fruticosus* as repeat elements using RepeatModeler and RepeatMasker. Of these elements, 3.0-7.8% were identified as LTR (long terminal repeat) retroelements, 0.3-1.0% transposons and the remainder were unclassified (Table 5). Since RepeatModeler utilizes RECON and RepeatScout to create a de novo model in RepeatMasker in place of the *Arabidopsis* model, details about the subcategories of LTRs and transposons which are included in the model could not be addressed. Maughan et al. [35] utilized GR-RSC on the *Arabidopsis* lines Ler-0 and Col-4. Utilizing RepeatModeler, then RepeatMasker on their sequence data from these lines, we found an average of 6.2% were identified as repetitive elements, of which 4.4% were identified as LTR retroelements and 0.4% were transposons. By way of comparison, the downloaded full “non-genome reduced” sequence of *Arabidopsis* line TAIR10 had a similar 7.4% of the sequence identified as repeat elements of which 3.0% were LTR retroelements and 0.2% were transposons (Table 5 and Figures 3 and 4). These data suggest that the GR-RSC method reflects, at least for repetitive elements, similar proportions as to that found in the full sequence of *Arabidopsis*.

Broderick et al. [5] hypothesized that the broad range found in *Penstemon* genome sizes, of the same ploidy, may be explained by retrotransposons. Lynch [60] detailed a relationship between genome size and repeat elements suggesting a linear relationship between the number of elements and genomes size [60-62]. The four *Penstemon* species used in this study provide insufficient evidence to establish a linear relationship between genome size and repeat elements in *Penstemon*. However, the three smaller, similar sized, *Penstemon* genomes possess comparable quantities of repetitive elements whereas *P. cyananthus* (the largest genome) has nearly double the number of repeat elements compared to the other three species (Figure 3).

Not only do repetitive elements largely influence genome size, but they are also likely to evolve more rapidly than do low-copy sequence [62,63]. Thus, repetitive elements of a species take on unique “fingerprints” which become valuable in phylogenetic relationship studies [64,65]. Thus, our limited four *Penstemon* species genomic data set suggest agreement with the two hypotheses that firstly, repetitive elements are a major component of the genome size variation identified by Broderick et al. [5]. Secondly, these elements are variable between the species we tested suggesting the possibility of identifying species

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays

Name	Contig Source ¹	SNP Allele GenBank Accession # ²	SNP Type	Allele Specific A1 Forward (5'→3') ³ Allele Specific A2 Forward (5'→3') ³	Common Allele Specific Reverse 5'→3')	<i>P. davidsonii</i>	<i>P. cyananthus</i>	<i>P. dissectus</i>	<i>P. fruticosus</i>	<i>P. pachyphyllus</i>	<i>P. cyananthus</i> + <i>P. davidsonii</i>	<i>P. dissectus</i> + <i>P. davidsonii</i>
PenSNP00001	00336CD	JX649978	A/G	AAGATTGCA TGGAGAGGA AATGGATT AGATTGCAT GGAGAGGAA ATGGATC	CGATCCAAA TGGCAGATC CGAGAAA	X ⁴	Y	X		X	Y	Y
PenSNP00002	00405CD	JX649979	C/T	ACGCGAGTA ATAAGTTGG TTTTCTTC GACGCGAGT AATAAGTTG GTTTTCTTT	CCAACACTT CCGCAGAAG CTCTTAA	Y	X	Y	X	Y	H	H
PenSNP00003	02625CD	JX649980	A/T	AAAAGCTCC CAAACATGA CTATGAACT AAAAGCTCC CAAACATGA CTATGAACA	AATTCCTCGA CACTTGAAGA GAGCGTAA	Y	X	Y		Y	H	H
PenSNP00004	02857CD	JX649981	A/C	ATCAAATGA ACTTGCTC ATGAGCCT CAAATGAAC TTGTCTCATG AGCCG	GCAACAAGGT GCAAAAAATT GTAGCGTAA	X	Y	X		X	H	H
PenSNP00005	03943CD	JX649982	A/G	ACTACCAA ACTACCCT CCCTTA ACTACCAA ACTACCCTC CCITG	GGGGTACAGA GTTGAGAAGA AGGAA	X	Y	X		X	H	H
PenSNP00006	04420CD	JX649983	A/C	TGTCTTAA ATCGATATG ATGAGGCT GTCTTAA TCGATATGAT GAGGCG	GTGGTTCTC CCCTTTAGA GGACTT	Y	X	Y	X	Y	H	H

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00007	08446CD	JX649984	A/T	GGCAACATC CTCAGCAGA GACA	CCGACTCCCT TAGCAATCTT AGCAT	Y	X	X		X	H	X
				GGCAACAT CCTCAGCA GAGACT								
PenSNP00008	11303CD	JX649985	C/G	GGGTGGTA TTGGTTAC TTTTATGGG	CGGTATAAGA GCAACTAAGC TAAATGACTT	Y	X	Y		Y	H	H
				GGGTGGTAT TGGTACTT TTATGGC								
PenSNP00009	11357CD	JX649986	C/T	ACAATATTTG ATAATTCATT CTCAAGTGCG	AAGCATGCAG TGAGACAAAA GCTAAGAT	X	Y	X	Y	X	H	H
				CACAATATT GATAATTCAT TCTCAAGTGCA								
PenSNP00010	11935CD	JX649987	A/C	AGCCTGATTA TCCCTTAAAC CCAATT	GAATCACGG CGGGGGAG CAAAT	X	Y	X		X	Y	
				GCCTGATTAT CCCTTAAAC CCAATG								
PenSNP00011	12047CD	JX649988	C/T	TTTGCCACT GCAGTGAC CATC	TGCTCCAGT CCGAAGGA AGTTGAAT	X	Y	Y		X	Y	Y
				CTTTGGCAC TGCAAGTAC CATT								
PenSNP00012	12119CD	JX649989	A/G	AAGATAGAC GTGGTATTTT TTCAGCA	GCAATTAG TCACAGAC CATAGTGG	X	Y	X		X	H	H
				AGATAGACG TGGTATTCT TCAGCG								
PenSNP00013	12398CD	JX649990	A/T	TATTTTCCTT TCTGCAATC TCAACATTGA	GTTGAGTGTG ATTTTAGAGT GCAITTAGTT	X	Y	X		X	Y	Y
				ATTTTCCTT CTGCAATCT CAACATTGT								

Table 7 Penstemon SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00014	13398CD	JX649991	A/C	AGGCCTGTGG CTGACTTGTC GGCCTGTGG CTGACTTGTC	GGCATATCT TTGCCCGTT TCCACAA	X	Y	X	X	X	H	H
PenSNP00015	13752CD	JX649992	A/C	AAATGCTC CCTCATTG ACCATATGA ATGCTCCCT CATTTGAC CATATGC	GTC AACGG ATTTGTGGA AGTCGGTA	Y	X	Y	Y	H	H	
PenSNP00016	14394CD	JX649993	C/G	TGAAAATTTC AGATTTAATG AACAAACAGTC GAAAATTTC GATTTAATGAA CAAACAGTG	AGACTTGTA CAAATTCCTT GGGTCCAAA	X	Y	X	Y	H	H	
PenSNP00017	14661CD	JX649994	A/G	TGACCAAGGA ATCTGTCAAG AACTT GACCAAGGA ATCTGTCAA GAACTC	CTTCTACTGTG GCTGTTCCACC TCTA	Y	X	Y	H	H		
PenSNP00018	15226CD	JX649995	G/T	TACCTCCAAT TGTGATGCA ACATTAG CTTACCTCCA ATTGTGATGC AACATTAT	CTAAGTGA GAAGCACA AGGA	X	Y	X	X	H	H	
PenSNP00019	17421CD	JX649996	G/T	ATCCTCCTC CTTTGCATC AAAGC CATCCTCC TCCTTGC ATCAAAGA	GAGCCAA CCTCGACT GCTTCTATTT	Y	X	Y	Y	X	H	
PenSNP00020	17816CD	JX649997	A/G	AAGGACTG AGTACCAA GACAGATCT GGACTGAG TACCAAGA CAGATCC	GCCAGGGTA CTGAACCTG TCITTTA	X	Y	X	X	H	H	

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00021	18745CD	JX649998	C/T	AGCATATTG AAAAGATC AGTCGCATAG AAAGCATAT TGAAAAGAT CAGTCGCATAA	CAGCTGCTCC TATCCAATC TTCGAA	Y	X	Y	Y	H	H	
PenSNP00022	19267CD	JX649999	A/G	AAATACCT GAGCTTCT GCCTCTTGT ACCTGAG CTTCTGCC TCTTGC	GATGCTCGT CATCTTGCT CAACGAT	X	Y	X	Y	X	H	H
PenSNP00023	21409CD	JX650000	C/G	ACCATTTCAG GTAATATTT CCAAAGGC ACCATTTCAG GTAATATTT CCAAAGGG	AGCGGTTCT AGAACCGT CAATGCTT	Y	X	Y	Y	Y	X	H
PenSNP00024	22934CD	JX650001	A/G	GTACAATTGT CAAGTGTGTA TTTTCTTACATA ACAATTGTCA AGTGTGTATT TTCTTACATG	GCACTGCAC CATTTCATGC CCTAAAA	Y	X	Y	Y	H	H	
PenSNP00025	22942CD	JX650002	A/T	ATCCGATTCT TCGTCTACTA TGCCA ATCCGATTCT TCGTCTACTA TGCCT	AGAAAAGCA CAAGCTGAA ATCAGGGAA	X	Y	X	Y	H	H	
PenSNP00026	27992CD	JX650003	A/G	TCCTCCTCG TCTCTTCT CTT CCTCCTCG TCTCTTCC TCTC	CTTGACCCTG CCAAAGAAG GAAAGAA	Y	X	Y	Y	H	H	
PenSNP00027	01179DD	JX650004	A/G	TCGACCC CAACCTG TCACA CTTCGACC CCAACCTG TCACG	CTTGCTTGTT TCGGAAAGAG	Y	X	Y	Y	H	H	

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00028	01235DD	JX650005	C/T	TGTGATCTT TGGTTTGAA CTTTGTC <hr/> CACTTGTGAT CTTTGGTTTG AACTTTGTT	CTACCAAAC TCACTCTAAC ATCCGGAT	X	Y	X	X	H	H
PenSNP00029	01600DD	JX650006	A/G	TGGTCTTGTT CTTTACCATT ACGCAT <hr/> GGTCTTGTT CTTTACCAT TACGCAC	GAAGTAGCTG CCATGGAAA AGGAAGTT	X	X	Y	X	X	H
PenSNP00030	04630DD	JX650007	A/G	AGTAGTACA GAATACTTAA AACTATCACCA <hr/> GTAGTACAGA ATACTTAAAA CTATCACCG	GTTGGGGGA GTTGCCTTCT TGAAAT	X	X	Y	X	X	H
PenSNP00031	05304DD	JX650008	C/T	AGTTTTCTTT TGTCCTTATG TGCAG <hr/> CAGTTTTCT TTTGCCTTA TGTGCAA	AAGGCTTAGC TTGGATGATA TCCTACAA	Y	Y	X	Y	Y	Y
PenSNP00032	05884DD	JX650009	A/T	GTCACCGCC TCCGATTGA GATT <hr/> GTCACCGCC TCCGATTGA GATA	CGGCTTTTGA CGCCGCCGT AAA	X	Y	X	X	H	H
PenSNP00033	06956DD	JX650010	G/T	GTTGATTCTA CAGATCTTAA TTCTTGATTG <hr/> AGTTGATTCTA CAGATCTTAA TTCTTGATTG	TACTACAAA GGGTAAAAAG TGCAATTCATA	X	Y	X	X	H	H
PenSNP00034	08307DD	JX650011	C/G	ACATTAAGG GTCCACCAA AAATCCG <hr/> ACATTAAGG GTCCACCAA AAATCCC	GCGCAATTAA AATCTCTTAAA TCACCTGGT	Y	Y	X		Y	H

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00035	08352DD	JX650012	A/T	AGTACAAGGA AAAACCTTTTA TTAGTAAGTATA	CTGACACAA ACCCATTCTA ATATGACCAA	X	Y	X		X	H	H
				AGTACAAGGA AAAACCTTTTAT TAGTAAGTATT								
PenSNP00036	08488DD	JX650013	A/T	GTGTTGGAG AGCCAGGT GCGA	GTATTGAGGAT CATTCTGACAA AAAACATA	Y	X	Y		Y	H	H
				GTGTTGGAG AGCCAGGTG CGT								
PenSNP00037	08608DD	JX650014	C/T	GTAGATAAG TTGATTGCGA GAGGC	CCAAACAAAT GCACCACATT CTCCTT	X	Y	X	Y	X	Y	H
				GGTAGATAA GTTGATTGC GAGAGGT								
PenSNP00038	08831DD	JX650015	A/T	TTTGAAGTGC CATGTAAAGT TGTTTTAGA	ATTTTGAACCA AGGAGCTATC AGAGG	X	Y	X		X	Y	H
				TTGAAGTGC ATGTAAAGTT GTTTTAGT								
PenSNP00039	08947DD	JX650016	A/T	GGGATCGTAA AACTCAGGAA AAATGA	TCAGATACTC GTGGGGTCTT CGATT	X	Y	X	H	X	Y	H
				GGGATCGTAA AACTCAGGAA AAATGT								
PenSNP00040	08959DD	JX650017	A/G	AGAGAATGAAG AAGGAGAAGGA AGAAA	CTCCTACGG TTGCATTATC GGTAGTA	Y	X	Y		Y	Y	Y
				GAGAATGAAGA AGGAGAAGGA AGAAG								
PenSNP00041	09272DD	JX650018	A/T	TTCTACAAAAC AATCAGCAGTC ATCATT	TCGACACCTT TTGCCTTATC TTGAA	X	Y	X		X	Y	H
				TCTACAAAAC AATCAGCAGT CATCATA								

Table 7 Penstemon SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00042	09369DD	JX650019	C/T	GTTTTATACG CATCCATATAC ATAATAATAG	GGTTCACCTCT CCAGAAATAA AATCTTATAT	Y	X	Y	Y	H	X	
				GTTTTATACGC ATCCATATACAT AATAATAA								
PenSNP00043	09764DD	JX650020	A/G	AATTCAACGTC AAATTGCAAG GTTGCA	TTCACTATAC CGGCTGAGT TGGCAT	Y	X	Y	Y	H	H	
				CAACGTCAAATT GCAAGGTTGCG								
PenSNP00044	10765DD	JX650021	A/G	TTTTTAATAAAT ATCCTGGTGGAT AATTTAT	AAATTGAGT GGATGGCTA GGAAGACTAA	X	Y	X	Y	X	H	H
				TTTTTAATAAAT ATCCTGGTGGGA TAATTTAC								
PenSNP00045	10870DD	JX650022	A/T	AGATCTGGAG ACTAAAT	CGAAGAGTT TGGGTGGGC GGAT	X	Y	X	X	Y	Y	
				AGATCTGGAG ACTAAAA								
PenSNP00046	11107DD	JX650023	C/T	GTCCGACGTG ACAATGCAGC	CGCCGTCAA AGAGACTTT GTTGGAT	Y	X	Y	Y	H	H	
				CTGTCCGACGT GACAATGCAGT								
PenSNP00047	11531DD	JX650024	C/T	AGAAGATTCTT CGGCTGGGAGC	TCTTCACATG ATTACGACAA TGGCTGAAT	X	Y	X	X	H	H	
				AAGAAGATTCTT CGGCTGGGAGT								
PenSNP00048	11655DD	JX650025	A/G	ACGTCCATGGA GGACCATAAA	GCTGTCITCC TGCAAGGAA CTTCTT	X	Y	X	X	H	H	
				CTACGTCCATGG AGGACCATAAG								
PenSNP00049	11974DD	JX650026	G/T	AAAATGCATGTA GTTTGGTTTACG	CACACCCCC AAAGGAAG AATAGCAT	Y	X	Y	Y	H	H	
				AAAATGCATGTA GTTTGGTTTACT								

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00050	13159DD	JX650027	C/T	TGAATGTACTTT TCATTGATAGA GAACG GTTGAATGTAC TTTTCATTGATA GAGAACA	AACAATAGT ACAACACAAC TAAAGCAGAGA	Y	Y	X	Y	Y	H	
PenSNP00051	13463DD	JX650028	G/T	GCCTTTGACG GCCAAGGAT TTC CGCCTTTGA CGGCGAAG GATTTA	GCAAGCACGG CACTAAGCCCTT	X	Y	X	X	H	H	
PenSNP00052	14334DD	JX650029	A/G	AGAAACAAC AAATACGAA TAAATCACCCA GAAACAACA AATACGAATA AATCACCCG	TTCGAAAATTG TGCTTGAATCA CGCAGT	X	X	Y	H	X	X	Y
PenSNP00053	00290DD03373CD	JX650030	C/T	TGCCTTTGCG TCGCCACAATC CTTGCCTTTG CGTCGCCAC AATT	AGCTAAGAGA TGGGCAGACT TTACAAAAT	Y	X	Y	Y	H	H	
PenSNP00054	00354DD04637CD	JX650031	A/G	GCAAAAGG GAACCTCA TTTCGTT CAAAAGGGA ACCCTCATT CGTC	TACTTGTCTGG GACTTTTCTT TCTCTT	X	X	Y	X	X	H	
PenSNP00055	01161DD11697CD	JX650032	A/G	ACTGGTAAA TACTACTACG TTCACAGT CTGGTAAA TACTACTAC GTTACAGC	GAAACACAGCA GCCCAACGACA TAT	Y	Y	X	X	Y	Y	H
PenSNP00056	01323DD15501CD	JX650033	A/G	ACCTGAAGA ATTGTTCAC TACTTCGT CCTGAAGAA TTGTTCAC ACTTCGC	GGATCGGGTGG ACGATTTGTGTT	X	Y	X	X	H	H	

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00057	01541DD02481CD	JX650034	A/G	AATTAGAAC CACATCCAC TGATTCCAA	GGAGCCCAAA CCTTTTACATT CTTTTCTA	Y	X	Y	Y	H	H
				AGAACCACA TCCACTGAT TCCAG							
PenSNP00058	02019DD03127CD	JX650035	A/G	GTGATTGTTA AATCTGAATA TATAATTTCTTTT	GTACGAGGCT TCGAAAAAGA CCAGAT	X	Y	X	X	H	H
				GTGATTGTTA AATCTGAATA TATAATTTCTTTC							
PenSNP00059	02851DD17191CD	JX650036	A/G	AAGAGTTGA TCCTAAGTTA TCGAGA	GAAGAAAATC ATTGTCCAC ATCTCGTGTA	X	Y	X	X	Y	H
				AGAGGTTGAT CCTAAGTTAT CGAGG							
PenSNP00060	03089DD14703CD	JX650037	C/T	TTTCAGAGTC ACTAATGTTT TCACG	GCATTTCTTG TCCATCTCTT CAAGATGTA	X	X	Y	Y	X	H
				GTTTCAGAG TCACTAATGT TTCACA							
PenSNP00061	03423DD25897CD	JX650038	A/C	AATTCCTTA CGTCCATTTG ATCGGAT	TATTCCTAGA CATGGACAT GGAAATTGAGA	Y	X	Y	Y	H	H
				CTTCTACGT CCATTTGAT CGGAG							
PenSNP00062	04632DD19186CD	JX650039	A/T	AAATGGGT CAGCTGAA ATTTCGCA	CTCTTCTTAC TCTGTTTTTCT TCTTTT	Y	Y	X		Y	H
				AAATGGGT CAGCTGAA ATTTCGCT							
PenSNP00063	05160DD08243CD	JX650040	C/G	TCGATCGTTG AAATGATAAT TGATACAAG	GATCCATA GACTTCTTTT AAGGATTCTAA	Y	X	Y	Y	H	H
				CGATCGTTG AAATGATAA TTGATACAAC							

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00064	06332DD03627CD	JX650041	A/C	ATCAAATGCC ATAGATCCTG CAGATTT ----- CAAATGCCA TAGATCCTG CAGATTG	ACATTCCTAC ACCAACTTCTT CCTACTA	X	X	Y	X	X	H
PenSNP00065	08748DD13630CD	JX650042	C/G	AGCTGTTT AGGAGGT TCATGAATG ----- AGCTGTTCA GGAGTTCA TGAATC	CACCATGTGA ACCAACACT ATTGTCATTT	X	Y	X	X	H	H
PenSNP00066	09773DD14323CD	JX650043	A/G	TCATGCCCA TTCCCCA ----- CATGCCCA TTCCCCG	CCTGGTATGA ACATGGGGA GGTTAT	Y	Y	X	Y	Y	
PenSNP00067	10248DD06150CD	JX650044	C/G	TGTGTCATT GAAATCAA TCCGC ----- GCTTGTGTC ATTGAAAT CAATCCGG	GTTTCATATCT CCCTTTGAGC TTCTTGAA	Y	Y	X	Y	Y	H
PenSNP00068	10624DD11358CD	JX650045	A/G	GTGGCAGT GTGAAACT GCATCA ----- GTGGCAGT GTGAAACT GCATCG	GTTTTCCCT GGGTGCTA AGGTTTAT	Y	X	Y	Y	H	H
PenSNP00069	11267DD06273CD	JX650046	A/C	ACCAAATA CTTATTAGC TCCAGTCGAA ----- CCAAATAC TTATTAGCT CCAGTCGAC	GACTGAAG GATGTTGC GAGAGGC	Y	Y	X	Y	Y	H
PenSNP00070	11564DD17128CD	JX650047	C/T	TGGACTTG GCATTGAA ACAAAAGATC ----- AATTGGAC TTGGCATTGA AACAAAAGATT	ATATGAAA CTCCCCAC AAGAAA	Y	X	Y	X	H	H

Table 7 *Penstemon* SNP marker name, GenBank dbSNP accession ID, polymorphism type, KASPar™ primer sequences (A1, A2 and common allele specific reverse) for all 75 functional SNP assays (Continued)

PenSNP00071	11647DD17264CD	JX650048	C/G	GCACGAGCC AAAATCCT GAGC	ATTGCGATG TGTATCCT GTGTGGGA	X	Y	X	Y	X	H	H
				GCACGAGCC AAAATCCT GAGG								
PenSNP00072	11671DD17144CD	JX650049	C/T	GTGCAGCA ACCCCTATT CATGAC	CCTGTCAA AACATATGAT CTTCATTGGAA	X	Y	X			H	H
				ATGTGCAG CAACCCCT ATTCATGAT								
PenSNP00073	12915DD17470CD	JX650050	A/G	AAGAAAAG GGTGGACAA ATTAACCGT	CAGAACAAC ATCATACTTG ATAAATCTCT	X	X	Y		X	X	H
				GAAAAGGGT GGACAAATT AAACCGC								
PenSNP00074	13828DD14937CD	JX650051	C/T	GTAAGATAT GCTGCCAGA TGG	CTCTGAAGAA GTTTTGTCCT TGATAGCTA	Y	Y	X		Y	Y	H
				GTAAGATAT GCTGCCAG ATGA								
PenSNP00075	14286DD18608CD	JX650052	G/T	GTATTGAG AGCCACT ACCGG	CCACTGAAT TGTTTGAAGA GTTTGGGAA	Y	X	Y		Y	X	H
				CTGTATTGA GAGCCAC TACCGT								

¹These contigs have been deposited at DDBJ/EMBL/GenBank as a Whole Genome Shotgun project under the accessions AKKG00000000 (*P. cyananthus*), AKKH00000000 (*P. dissectus*), AKKI00000000 (*P. davidsonii*), and AKKJ00000000 (*P. fruticosus*). The version described in this paper is the first version for each accession, XXXX01000000.

²The GenBank accession identification for the full sequence for each allele with the specific SNP bp identified.

³KASPar™ primers: A1 and A2 primers are SNP allele specific. All A1 Forward primers had the follow universal primer GAAGGTGACCAAGTTCATGCT added to the 5' end of the allele specific sequence. All A2 Forward primers had the follow universal primer GAAGGTGCGAGTCAACGGATT added to end of the 5' allele specific sequence.

⁴H = heterozygous compared to either homozygous condition for either "X" or "Y".

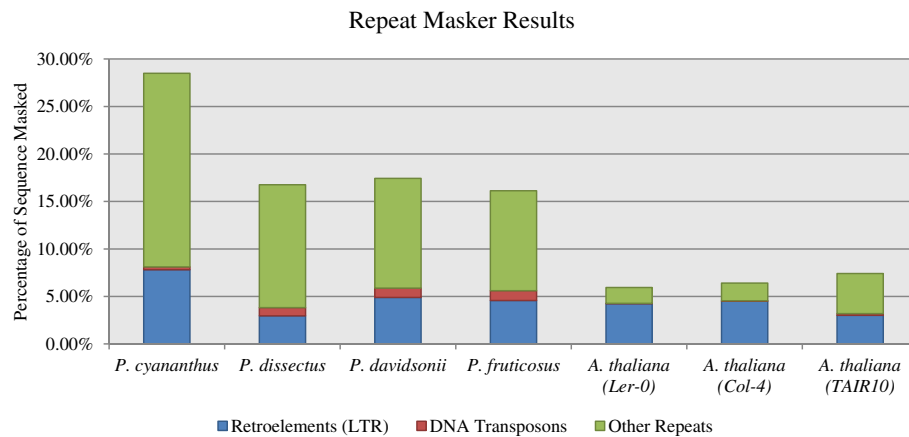


Figure 3 Percentage of retroelements, DNA transposons and other unclassified repeats in *Penstemon cyananthus*, *P. dissectus*, *P. davidsonii*, *P. fruticosus*, and both genome reduced and non-genome reduced *Arabidopsis*¹. ¹ Genome reduced *A. thaliana* sequence from Maughan et al. [35]; *A. thaliana* RILs Ler-0 and Col-4; Non-genome reduced *A. thaliana* sequence downloaded from TAIR (The *Arabidopsis* Information Resource) as whole chromosomes; the diploid (2n = 2x = 16) genome size as reported by Broderick et al. and Schmutz et al. [5,59].

specific repetitive elements. However, without further comparisons we were unable to identify specific repetitive elements associated with the four *Penstemon* species used in this study.

Gene ontology

Using BLASTX we identified an average of 21.5% of the contigs across the four species as putative genes with an average of 13.9% annotated by Blast2GO (Table 5). These putative genes were compared and contrasted in a more detailed study by Dockter [23]. Furthermore, he compared the *Penstemon* sequences

to known genes from the related genera *Antirrhinum* and *Mimulus*, and identified nine putative *Penstemon* genes from *Antirrhinum* and 14 from *Mimulus* with an e-value below $1.0e^{-13}$. Three genes (NADH dehydrogenase from *M. aurantiacus*, ribosomal protein L10 from *M. guttatus*, and ribosomal protein subunit 2 from *M. aurantiacus*, *M. szechuanensis*, and *M. tenellus* var. *tenellus*) were perfect hits (e-value = 0.0).

Conclusions

Penstemon are recognized for their phenotypic variation and their adaptation to multiple environments

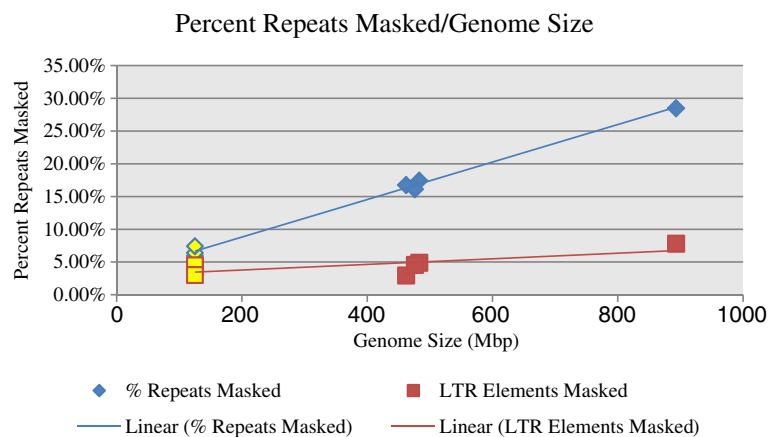


Figure 4 Relationship between genome size and repeat elements in *Penstemon* including the relationship of both LTRs and total repeat elements to genome size for both genome reduced *Penstemon* and genome reduced/non-genome reduced *Arabidopsis*¹ (yellow). ¹ Genome reduced *A. thaliana* sequence from Maughan et al. [35]; *A. thaliana* RILs Ler-0 and Col-4; Non-genome reduced *A. thaliana* sequence downloaded from TAIR (The *Arabidopsis* Information Resource) as whole chromosomes; Genome size as reported by Broderick et al. and Schmutz et al. [5,59].

[6-8,13,14,17,30,31]. Broderick et al. [5] found that this diversity is reflected by a wide range in their genome sizes. Nevertheless, even with this demonstrated plasticity we have identified evidence that there is a high level of sequence conservation across the genus. This apparent sequence conservation is in harmony with the hypothesis that *Penstemon* has rapidly irradiated to its variety of species rather recently in evolutionary time [13,14]. Furthermore, our study identified evidence that the genome size variation in *Penstemon* is rooted in the amount of repetitive elements in each species.

Despite the large differences in *Penstemon*'s genome size, the finding that the genus has a great deal of sequence conservation is invaluable for the development of interspecific markers. The further development and mapping of a number of conserved markers will facilitate the domestication of xeric *Penstemon* cultivars via interspecific hybridization which are largely unexploited largely due to crossing barriers [6-8,10-12,15]. Viehmeyer [16] hypothesized that it might be possible to develop *Penstemon* breeding lines that would facilitate the indirect interspecific hybridization of any two species within the genus. He and others have used traditional breeding techniques to develop a number of interspecific hybrids [7,11,15,17,66]. Clarifying the phylogenetic relationships within the genus should facilitate these objectives [67]. In the largest *Penstemon* phylogenetic study conducted to date, Wolfe et al. [14] sequenced the ITS and two chloroplast genes in 163 species. They concluded that many species are polyphyletic in their origins thus making them difficult to discriminate between one another; thus, requiring additional molecular studies to more accurately define taxonomic relationships.

We tested 51 SSR/INDEL based markers (Table 3), and identified several thousand inter- and intraspecific SNPs (Table 6), all of which have potential as both inter- and intraspecific markers. Of the 51 SSRs/INDELs we selected 12 to test across 93 *Penstemon* taxa. The resulting data was used to more clearly define the phylogenetic relationships of those taxa but our results were incoherent. It is possible that some of these markers may represent more than one locus in the *Penstemon* genome. This situation has been identified by others as a potential weakness in using SSR based markers in interspecific phylogenetic studies [46,47]. A major reason for the vagary in *Penstemon*'s phylogeny is that it appears to have quite recently evolved and rapidly radiated leaving weak species boundaries [13,14]. Furthermore, there are a number of reports of speciation via natural interspecific hybridization found within the genus [14,68-73]. Therefore, like Wolfe et al. [14], we concluded that better marker data sets will be required to reduce present phylogenetic ambiguity.

To gain clearer insights into the relationships of *Penstemon* it will take carefully designed large scale sequencing studies. There are methods which are showing promise to do such studies economically. One example would be to utilize GR-RSC or similar methods which will sample large quantities of homologous sequence of a genome at ever decreasing costs [18,20,74]. Since our SSR/INDEL, sequence, and SNP data have demonstrated broad applicability across *Penstemon* it becomes evident that further studies utilizing this same GR-RSC protocol and downstream analysis on additional species would allow broader comparisons of putative genes, repeat elements, SNPs and SSRs, facilitating a much better understanding of the genus. Furthermore, using this technique on carefully selected parents and their segregating progeny would allow *Penstemon* genetic mapping studies which would greatly enhance the ability to do breeding and domestication studies within the genus. Historically, studies of this nature would have been unthinkable; however, mass homologous loci sequence studies are rapidly becoming feasible [18,20,74]. In the interim it is possible to take the data we report here and further test the 75 SNPs we have reported here along with others not yet developed and for around US\$0.05/data point [18,20] do a much broader study. Studies on homologous SNPs across many *Penstemon* taxa, similar to the *Amaranthus* study of Maughan et al. [20], should assist in developing improved insights into *Penstemon* phylogenetic relationships and produce high quality genetic maps from carefully designed segregating *Penstemon* populations.

Competing interests

The authors declare no competing interests.

Authors' contributions

Rhyan B Dockter, David B Elzinga, Brad Geary, P Jeff Maughan, Leigh A Johnson, Danika Tumbleson, JanaLynn Franke, Keri Dockter, and Mikel R Stevens. RBD performed the GR-RSC technique and either carried out or oversaw the all other steps of the study and participated in all planning and design of all experiments as well as their analysis and did the initial drafting of the manuscript. DBE did or assisted in all bioinformatics performed in this study. BG participated in the design of all aspects of the study as well as advised RBD and was involved in the editing and revising of the manuscript. PJM advised and assisted in the GR-RSC technique as well as advised RBD in relevant issues of the bioinformatics of the study and was involved in the editing and revising of the manuscript. LAJ advised and assisted RBD and MRS in the taxonomy related issues of the study and was involved in the editing and revising of the manuscript. DT, JF, and KD carried out all aspects, including basic analysis, of the marker studies reported. MRS was the senior advisor of RBD and was intricately involved in all aspects of the study and the manuscript. All authors both read and approved the final manuscript.

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