

Forms of *Melanoplus bowditchi* (Orthoptera: Acrididae) collected from different host plants are indistinguishable genetically and in aedeagal morphology

The sagebrush grasshopper, *Melanoplus bowditchi* Scudder (Orthoptera: Acrididae), is a phytophilous species that is widely distributed in the western United States on sagebrush species. The geographical distribution of *M. bowditchi* is very similar to the range of its host plants and its feeding association varies in relation to sagebrush distribution. *Melanoplus bowditchi bowditchi* Scudder and *M. bowditchi canus* Hebard were described based on their feeding association with different sagebrush species, sand sagebrush and silver sagebrush, respectively. Recently, *M. bowditchi* have been observed feeding on other plant species in western Nebraska. We collected adult *M. bowditchi* feeding on four plant species, sand sagebrush, *Artemisia filifolia*, big sagebrush, *A. tridentata*, fringed sagebrush, *A. frigidus*, and winterfat, *Krascheninnikovia lanata*. We compared the specimens collected from the four plant species for their morphological and genetic differences. We observed no consistent differences among the aedeagal parameres or basal rings among the grasshoppers collected from different host plants. Amplified Fragment Length Polymorphism markers were used to test the genetic relationships among the grasshoppers. Analysis of Molecular Variance and distance-based Unweighted Pair Group Method with Arithmetic mean dendrogram failed to reveal significant differences. Although the forms showed behavioral and minor color and size differences, the genetic data show all forms under study interbreed, which indicates they are a single species instead of four species or subspecies. These results indicate that host plant use may influence melanopline phenotype and suggest the need of further genetic analysis of subspecies recognized based on morphology, distribution, and ecology.

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**Forms of *Melanoplus bowditchi* (Orthoptera: Acrididae) collected from different host plants
are indistinguishable genetically and in aedeagal morphology**

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Key words

20 *Melanoplus bowditchi*, morphological variation, host plant, deme, ecotype, AFLP

21 **Introduction**

22 As a group, grasshoppers are somewhat unusual among herbivorous insects in that most
 23 are polyphagous, feeding selectively on plants from a number of unrelated plant families (Otte &
 24 Joern 1977). Because of their polyphagy, most grasshopper populations are not expected to
 25 experience disruptive selection associated with host choice. There are, however, some
 26 grasshopper species with restricted host ranges and a small number that are truly host specific
 27 (Otte & Joern 1977, Sword & Dopman 1999). Host specific grasshoppers also show differences
 28 in development rates, lifespan, and size relating to host use (Traxler & Joern 1999). In addition,
 29 host plant-associated genetic differences have also been observed in the study of *Hesperotettix*
 30 *viridis* (Thomas) and *Schistocera lineata* Scudder (Sword *et al.* 2005).

31 The sagebrush grasshopper, *Melanoplus bowditchi* Scudder, was described by Scudder in
 32 1878 (Scudder 1897). This grasshopper is a phytophilous species that is widely distributed in the
 33 grasslands of the western United States. Although it occurs in mixed-grass, shortgrass, desert
 34 shrub and bunchgrass prairies, it feeds almost exclusively on sagebrush species (Mulkern *et al.*
 35 1969) and its distribution is dependent on sagebrush plants. Six host plants are identified for *M.*
 36 *bowditchi* in Pfadt (1994), with the primary hosts being silver sagebrush, *Artemisia cana*, and
 37 sand sagebrush, *A. filifolia*. The other four species of sagebrush, along with silver sagebrush, are
 38 found in mixed-grass prairie and are reportedly consumed in minute quantities by *M. bowditchi*
 39 (Pfadt 1994). The species is potentially damaging, especially for silver sagebrush (Pfadt 1994).
 40 While silver sagebrush, is broadly distributed through western North America, sand sagebrush,
 41 *Artemisia filifolia* Torrey, is usually associated with deep sand deposits and serves as the host
 42 plant for *M. bowditchi* in areas where silver sagebrush is limited (Harvey 1981).

The subspecies, *Melanoplus bowditchi bowditchi* Scudder was proposed after the description of *Melanoplus bowditchi canus* Hebard (Hebard 1925). The original series of *M. bowditchi bowditchi* was found feeding on *A. filifolia*, while *M. bowditchi canus* was collected from big sagebrush, *Artemisia tridentata* (Hebard 1925). *Melanoplus bowditchi canus* is usually dark gray in color and is common in the northern Great Plains. Its preferred food plant is silver sagebrush, although it has also been observed feeding on other sagebrush species. It is normally found on taller plants until after oviposition, when it becomes abundant on shorter plants. It is seldom found on the ground (Anderson & Wright 1952). In comparison to *M. bowditchi canus*, *M. bowditchi bowditchi* has a larger body size, brighter yellow and brown colors, and very clear tegmina (Hebard 1925).

Hebard (1925) suggested that the gray patterned coloration of *M. bowditchi canus* was a result of a close relationship to the *Melanoplus cinereus* group rather than to other forms of the *Melanoplus flavidus* group (Hebard 1925). However, geographical differences in host plant use and morphology might also be the result of environmentally triggered variation among populations. For example, for a specialized flea beetle, areas with abundant hosts and frequent oviposition show a high level of host acceptance resulting in less use of low-ranking hosts (Tahvanainen & Root 1972). Where preferred plants are uncommon or their availability is obscured by related members of the plant community, thresholds for host acceptance are expected to fall, making the use of other plants more likely (Stanton & Cook 1983, Wiklund 1975).

Recent observations have revealed adult *M. bowditchi* feeding on other *Artemisia* species and winterfat, *Krascheninnikovia lanata*. These forms seem to be distinct both size and color pattern (Fig. 1) and exhibit behavioral differences. Specimens of *M. bowditchi* collected from fringed sagebrush, *Artemisia frigida* are exceptionally pallid (Fig. 1) ranging from a pale tan to pale gray and superficially resembling *Melanoplus angustipennis* (Dodge). In addition,

specimens collected from fringed sagebrush are more reluctant to jump than *M. bowditchi bowditchi* and often must be knocked from the small shrubs to be revealed, similar to reports for *M. bowditchi canus*. Fringed sagebrush is common in dry, well-drained soils or in disturbed areas. In mixed-grass prairie it is found with western wheatgrass *Pascopyrum smithii*, blue grama *Bouteloua gracilis*, and winterfat.

Specimens of *M. bowditchi* that are collected feeding on winterfat are silvery gray. Behaviorally, specimens of this form are much more wary than those collected from fringed sagebrush and silver sagebrush and jump readily like *M. bowditchi bowditchi*.

The objective of this research was to examine the genetic and aedeagal characteristics for the adults of *M. bowditchi* associated with different sagebrush species and to test our hypotheses that the forms associated with host plants are four distinct species.


Materials & Methods

Insects. —A series of adult *M. bowditchi* were collected from four host plants, sand sagebrush, *A. filifolia*, big sagebrush, *A. tridentata*, fringed sagebrush, *A. frigidus*, and winterfat, *Krascheninnikovia lanata* (Table 1). We also collected a distant outgroup, the mottled sand grasshopper, *Spharagemon collare* (Scudder) from bare soil patches in western Nebraska. Specimens were identified based on the available literature of Bruner (1897), Scudder (1897), Pfadt (2002) and Brust *et al.* (2008).

Aedeagal analysis.—For aedeagal studies, abdomens from three or four grasshoppers collected from each plant species were examined (Table 1). In each case the terminal part of the abdomen was separated, intestinal contents removed, and the remaining structure soaked in a solution of 5 % NaOH for 8-10 h, transferred to 70% ethanol for 10 min., and the aedeagi removed. Aedeagi

were cleaned under a dissecting microscope to remove connective tissue. They were preserved in 70% ethanol until examination. Photographs of aedeagi were taken through a dissecting microscope. A visual comparison was made of the structure of the terminal end of the aedeagus, and of the paramere structures.

Genetic analysis - Extraction and Quantification of DNA.—The locations, date of collection, number of specimens and plants from which specimens were collected for genetic analysis are presented in Table 2. Hind femora of specimens from each host plant were preserved in 95% ethanol and stored at -80 °C prior to genetic studies. A total of 35 grasshoppers were examined for genetic differences (Table 2). DNA was isolated from the hind femur of each form specimen using acetyletrimethylammonium bromide (CTAB) extraction protocol modified from Doyle & Doyle (1987). Each hind femur was placed in an autoclaved 1.5 ml microcentrifuge tube and rinsed for 10 min in Nanopure® water. The entire hind femur was homogenized in 250 µl CTAB buffer (100 mM Tris-HCl, 1.4 M NaCl, 0.02 M EDTA, 2% CTAB, and 0.2% β-mercaptoethanol) using sterile white quartz sand and plastic pestles. Another 250 µl of CTAB was added to the tubes to make a volume of 500 µl. RNase A (15 µl of 0.05 g ml⁻¹) was added to each tube, and incubated for 2 h at 65° C. Proteinase K (15 µl of 0.02 g ml⁻¹) was added, and incubated for 1h at 37° C. Samples were centrifuged for 5 min at 20° C and 12,000 rpm. The supernatant from each tube was transferred to new autoclaved tubes and the tissue discarded. Chloroform: isoamyl alcohol (24:1) (500 µl) was added to the supernatant, and tubes were centrifuged at room temperature for 20 min at 12,000 rpm. The upper aqueous layer was transferred to new 1.5 ml Eppendorf tubes and the chloroform: isoamyl alcohol step was repeated to isolate the refined top aqueous phase. Chilled isopropanol (400 µl, -20° C) was added to the tubes to precipitate the DNA, and samples were stored over night at 4° C.

Samples were centrifuged at 12,000 rpm for 30 min at 4° C, to ~~form a pellet of DNA at the~~
~~bottom of the tube~~. The supernatant was discarded, and the DNA was washed with 400 µl of
chilled absolute ethanol followed by centrifugation for 5 min. The supernatant was decanted and
the wash was repeated using 70% ethanol. Tubes were centrifuged again for 5 min, then the
ethanol was removed and the samples allowed to air dry. The pelleted DNA was suspended in 50
µl autoclaved 1x TE buffer (10 mM Tris-HCL, 0.1 mM EDTA). 

AFLP-PCR methods for genetic analysis.—The Amplified Fragment Length Polymorphism
(AFLP) technique, modified from Vos *et al.* (1995) was used for DNA analysis. AFLP consisted
of digestion using *MseI* and *EcoRI* restriction enzymes, ligation of specific nucleotide adapters, a
preselective amplification using universal primers, and a selective amplification using specific
primer pairs.

Template preparation.—Restriction digestion was performed using 1.25 µl NEB Buffer 4 (New
England Biolabs, Foster City, CA), 0.125 µl bovine serum albumin (New England Biolabs),
0.0625 µl *EcoRI*, 0.0625 µl *MseI* (New England Biolabs), 3.94 µl Nanopure® water and 7 µl of
~20ng/µl DNA template for a total volume of 12.5 µl. The restriction digestion was incubated on
a GeneAmp 2720 thermal cycler (Applied Biosystems, Foster City, CA, USA) at 37 °C for 2.5 h.
A ligation mixture (5 µl) consisting of 0.5 µl *EcoRI* and *MseI* prepared adapters, (Integrated DNA
Technologies, Coralville, IA, USA), 0.5 µl T4 DNA ligase, 0.15 µl 10x T4 DNA ligase buffer
(New England Biolabs), and 3.35 µl Nanopure® water was dispensed into the tubes containing
the digestion product and incubated at 25 °C for 8 h. The ligation product was then diluted using
135 µl of 1x TE buffer. A Nanodrop® spectrophotometer (Thermo Fisher Scientific, Walltham,
MA, USA) was used to determine the quantity and quality of DNA in ng/µl from each tube.

134 *Preamplification.* A preamplification mix consisting of 10 µl Preamplification Primer Mix II (LI-
 135 COR Biosciences, Lincoln NE, USA), 0.25 µl Amplitaq 360 DNA polymerase, 0.75 µl 25 mM
 136 MgCl₂, and 1.25 µl 10x PCR buffer II (Applied Biosystems, Foster City, CA) was mixed with
 137 1.25 µl of ligation product and run on a PCR program of 20 cycles (30 s at 94 °C, 1 min at 56 °C,
 138 and 1 min at 72 °C), then stored at 4 °C. Nanopure® water was used to dilute the product to a
 139 ratio of 1:20. Nucleotide sequences of adapters, preamplification primers and selective primers
 140 tested are shown in Table 3. A combination of different primer sets was tested and the best
 141 working primer sets for grasshopper DNA were chosen (Table 4).

142 *Selective amplification.*—The selective PCR mix was prepared consisting of 1.2 µl 10x PCR
 143 buffer II, 0.72 µl 25 mM MgCl₂, 0.24 µl (10 mM) deoxynucleotide triphosphate mix, 0.07 µl
 144 Amplitaq 360 DNA polymerase (Applied Biosystems), 0.5 µl of *Mse*I primer (5.0 µM), 0.3 µl
 145 *Eco*RI (1.0 µM) IRD-700 labeled primer (Integrated DNA Technologies, Coralville, IA), 6.97 µl
 146 nanopure® water, and 1.5 µl of the preamplification template DNA. This step was performed in
 147 the dark due to light sensitivity of the labeled primers. Selective amplification was performed on
 148 a GeneAmp 2720 thermal cycler (Applied Biosystems) with one pre-PCR cycle (30 s at 94 °C, 30
 149 s at 65 °C, 1 min at 72 °C), 12 cycles of 30 s at 94 °C, 30 s at 65 °C → 56 °C, 60 s at 72 °C, and 23
 150 cycles of 30 s at 94 °C, 30 s at 65 °C → 56 °C and 60 s at 72 °C. Blue stop solution (LI-COR
 151 Biosciences, Lincoln, NE) (2.5 µl) was used to end the reaction. The product was then denatured
 152 for 3 min at 94 °C and stored at -20 °C.

153 *Data scoring and analysis.*—The amplified DNA was electrophoresed in KB^{ph} 6.5%
 154 polyacrylamide gel on a GeneReader 4200 DNA analyzer (LI-COR Biosciences) which detects

bands through infrared inflorescence. An IRDye-700 labeled 50-700 bp size standard was used to estimate fragment size. The correlation of % coefficient of variation and the total number of markers was estimated using Bootsie (Foster unpublished).

Gels were scored on Saga Generation 2 version 3.3.0 software (LI-COR Biosciences). Data were converted to a binary matrix for analysis, with 1 = presence of a band and 0 = absence of band. Data were analyzed using Arlequin version 3.5 (Excoffier & Lischer, 2010) and Popgene version 1.32 (Yeh *et al.* 1999). Phylogenetic relationships were examined using distance-based methods for the different host associated *M. bowditchi*. An Unweighted Pair Group Method with Arithmetic mean (UPGMA) dendrogram was constructed using presence/absence characters with the software package PAUP* version 4.0beta (Swofford 2001). Bootstrapping was performed with 1,000 replicates.

Results

Aedeagus examination.—All examined grasshoppers collected from different host plants had similar aedeagi. The structure and angle of parameres were similar among specimens (Figs. 2 and 3). The mean (± 1 S.E.) paramere lengths (0.81 ± 0.07 mm) were shortest in grasshoppers collected from fringed sagebrush and longest in grasshoppers collected from winterfat (0.98 ± 0.02 mm); however, differences were not significant (ANOVA, $P = 0.085$). Aedeagal lengths were also not significantly different, although specimens collected from sand sagebrush had a mean length of 0.81 ± 0.02 mm compared to those from winterfat, which had a mean length of 0.63 ± 0.02 mm (ANOVA, $P = 0.054$).

Genetic variation study.—The *M. bowditchi* from different host plants were initially screened for a total of 10 primer pairs of which six primer pairs (Table 4) were used for analysis. A total of

469 markers were scored using the six primer pairs and 63% of the loci were polymorphic. Using Bootsie examination (J. Foster, unpublished), approximately 96% of the variation in the *M. bowditchi* populations was captured with the chosen markers.

~~A dendrogram was constructed using a distance-based Unweighted Pair Group Method with Arithmetic mean (UPGMA). The UPGMA analysis did not reveal significant genetic structure differences among the *M. bowditchi* collected from different host plants and there were few nodes with bootstrap values greater than 70% (Fig. 4). The molecular sequences of grasshoppers collected from winterfat were spread across the dendrogram, whereas (Fig. 4).~~

The *M. bowditchi* populations were arranged in two groups based on host plant associations and descriptions of recognized subspecies (Hebard 1925, Pfadt 1994). Grasshoppers collected from winterfat were paired with those collected from fringed sagebrush (the *Melanoplus bowditchii canus* group) and those collected from fringed sagebrush were paired with those collected from sand sagebrush (the *Melanoplus bowditchi bowditchi* group). The AMOVA showed the majority of molecular variation (86.8%) occurred within populations (Table 5). Only 7.9% of the genetic variation occurred among populations within groups while the remaining 5.3% was due to the variation among groups (Table 6). Nei's (1987) gene diversity (G_{ST}) is described as the coefficient of gene differentiation, while fixation index (F_{ST}) is the measure of differentiation in sub-populations and is only applicable when there are only 2 alleles at a locus. Nei's genetic diversity (G_{ST}) is analogous to Wright's genetic divergence (F_{ST}). G_{ST} measures the degree of differentiation in multiple populations. The genetic divergence (F_{ST}) and gene diversity (G_{ST}) were low (0.1320 and 0.0879 respectively) while the N_m values (5.1905) were high (Table 6) indicating extensive gene flow among populations.

Discussion

A number of phytophagous insect species contain locally adapted host specific populations, although they utilize a number of host plants across their range (Futuyma & Peterson 1985, Thompson 1994). This phenomenon is only occasionally reported for grasshoppers (Sword & Dopman 1999). The sagebrush grasshopper *M. bowditchi* feeds on several species of sagebrush, although *A. cana* and *A. filifolia* serve as the main host plants (Mulkern *et al.* 1969, Pfadt 1994). The *M. bowditchii* subspecies differ somewhat in distribution with *M. bowditchi bowditchi* found in the southern grass plains and *M. bowditchi canus* found in the northern sagebrush plains, while the ranges broadly overlap in Wyoming and southwestern South Dakota (Hebard 1929). The geographical distribution of the host plants is very similar to the range of the subspecies of *M. bowditchi* (Pfadt 1994), and the feeding preference of this grasshopper has been shown to vary based on local plant availability. Examination of the crop content of *M. bowditchi* collected from North Dakota showed feeding on silver sagebrush, while the populations from western Nebraska ate sand sagebrush (Pfadt 1994). Even though there are some differences in distribution, color, size, and host use between the two described subspecies of *M. bowditchi*, it is important to question the rationale of naming a subspecies solely on these characteristics. Further, it is important to test if *M. bowditchi* feeding on other host plants represent cryptic species or additional subspecies.

In Nebraska and South Dakota, we identified four populations of *M. bowditchi* feeding on different plant species where other host plants were not present. Specimens exhibited behavioral differences and slight color and size differences consistent with previous descriptions of subspecies (Figure 1). Grasshoppers collected from sand sagebrush and winterfat were both very active, either jumping and flying to another host plant when approached within two meters or actively moving into the basal branches. In contrast, specimens collected from fringed sagebrush

generally did not fly far and had to be disturbed before they jumped or flew. Individuals found feeding on big sagebrush were more sedentary than most other *Melanoplus* forms (M. Brust, pers. obs). In our examination of male genitalia, we did not find consistent differences. We recognize that we had a small sample size and that a series of more individuals from each host would be better. However, among our samples, the aedeagi appeared similar and there were no significant differences in overall length or paramere angle or lengths (Figs. 2 and 3) suggesting that they are physically able to interbreed. Even if morphological differences had been noted, breeding tests would be required as variation in aedeagi occurs among individuals of a species and support for the hypothesis of genitalic incompatibility among species is controversial (Eberhard et al. 1998, Masto 2012). Thus, genetic testing of gene flow can aid in determining the presence of populations and incipient species.

We found no genetic differentiation or distinct lineages for *M. bowditchi* in relation to different host plants (Fig. 4) despite collecting grasshoppers from populations located more than 230 km apart (Ardmore, SD and Scottsbluff, NE). We found G_{ST} values between 0.06 and 0.2. A G_{ST} value of 1 would indicate nearly complete isolation of subpopulations while 0 indicates no isolation. A G_{ST} value greater than 0.5 indicates some genetic isolation among subpopulations (Nei 1987). Thus, the low G_{ST} in this study reflects the relative measure of variation among subpopulations with reference to total variation (Table 6). In this study, we were unable to identify any clusters in the dendrogram (Fig. 4) that could separate the populations of *M. bowditchi* into distinctive groups.

These results are similar to the conclusions of Brust *et al.* (2010) who found no genetic differences among *M. foedus foedus* (Scudder), *M. foedus fluviatilis* Bruner, and *Melanoplus packardii* Scudder. Also, Chapco & Litzenberger (2002), found no genetic differences between *M. foedus* and *M. packardii* nor between *Melanoplus angustipennis* (Dodge) and *Melanoplus*

femurrubrum (De Geer). The Analysis of Molecular Variance indicates that most of the variation (86.8%) was within populations with a small portion (5.3%) observed among groups, suggesting frequent interbreeding. Similarly, the variation among *M. bowditchi* from different host plants was low, supporting consistent gene flow. The F_{ST} value of 0.1320 supports the conclusion that genetic exchange occurring among the four subpopulations was sufficient to prevent either genetic differentiation or structuring into genetically differentiated subpopulations of *M. bowditchi*.

There is support in the literature for host plants resulting in distinct phenotypes. For example, Miller (1987) and Futuyma (1990) documented host-specific phenotypes in papilionid butterflies and *Ophraella* leaf beetles. The grasshopper specimens in this study were collected as adults feeding on specific plants, and they differed in color and behavior; however, they do not appear to have consistent genetic differences.

The use of the trinomial for *M. bowditchi* appears invalid; however, we found color morphs that differed in behavior associated with a different hosts and geography. These differences in phenotype appear to be influenced by the environment. These differences are potentially related to the diet, but other environmental factors may play a role. It is further unknown how variable this species is west of the Rocky Mountains. Thus, further detailed investigations for *M. bowditchi* with morphological and behavioral differences associated with host-plant use should be conducted and the genetic variation among forms should be investigated on a larger scale.

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270 **Author Contributions**

271 Mathew Brust collected the grasshopper specimens and identified host plants. Muhammad Irfan
 272 Ullah dissected grasshoppers and prepared genitalia for examination. Muhammad Irfan Ullah,
 273 Fatima Mustafa, and Kate Kneeland prepared specimens for genetic analysis and conducted
 274 analyses under the supervision of John E. Foster. W. Wyatt Hoback and Shripat Kamble provided
 275 guidance on experimental design and analysis. All authors discussed the results and commented
 276 on the manuscript.

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
352 Table 1. State, county location, host plant and date of collection for *Melanoplus bowditchii*
353 specimens used in aedeagus analysis.

State	County	Location	Host Plant	Date	Quantity
Nebraska	Dawes	8 km S of Chadron	Fringed sagebrush <i>Artemisia frigida</i>	July 24, 2010	3
South Dakota	Fall River	24 km N of Ardmore	Big sagebrush <i>Artemisia tridentata</i>	August 21, 2010	3
Nebraska	Morill	14.4 km SW of Alliance	Sand sagebrush <i>Artemisia filifolia</i>	July 24, 2010	3
Nebraska	Scotts Bluff	12 km N of Minatare	Winterfat <i>Krascheninnikovia lanata</i>	July 17, 2010	4

354 Table 1. Collection information for specimens of *M. bowditchi* from different host plants and
355 outgroup, *Spharagemon collare* used in genetic analysis. Specimens of *S. collare* were collected
356 with sweep nets from bare soil.

Species	State	County	Location	Host Plant	Date
<i>Melanoplus bowditchi</i>	Nebraska	Dawes	8 km S of Chadron	Fringed sagebrush <i>Artemisia frigida</i>	July 24, 2010 August 7, 2010
<i>Melanoplus bowditchi</i>	South Dakota	Fall River	24 km N of Ardmore	Big sagebrush <i>Artemisia tridentata</i>	August 21, 2010
<i>Melanoplus bowditchi</i>	Nebraska	Morrill	17.7 km SW of Alliance, 6.4 km E of Broadwater	Sand sagebrush <i>Artemisia filifolia</i>	July 9, 2010
<i>Melanoplus bowditchi</i>	Nebraska	Scotts Bluff	12 km N of Minatare	Winterfat <i>Krascheninnikovia lanata</i>	July 18, 2010
<i>Spharagemon collare</i> (outgroup)	Nebraska	Dawes	4.8 km S of Chadron	None	August 22, 2010

357 Table 3. Nucleotide sequences of adapters, preamplification primers and selective primers used in
358 this study. Sequences were described by Vos *et al.* (1995).



Oligonucleotide	Purpose	Sequence
<i>Eco</i> RI-1 (forward)	Adapter	5'-CTCGTAGAC
<i>Eco</i> RI-2 (reverse)	Adapter	5'-AATTGGTAC
<i>Mse</i> I-1 (forward)	Adapter	5'-GACGATGAG
<i>Mse</i> I-2 (reverse)	Adapter	5'-TACTCAGGA
E (N+0)	Preamplification Primer	5'-GACTGCGTA
M (N+1)	Preamplification Primer	5'-GATGAGTCC
M-CAA	Selective Primer	5'-GATGAGTCC
M-CTC	Selective Primer	5'-GATGAGTCC
M-CAG	Selective Primer	5'-GATGAGTCC
E-AAC	Selective Primer	5'-GACTGCGTA
E-ACT	Selective Primer	5'-GACTGCGTA
E-AGG	Selective Primer	5'-GACTGCGTA
E-ACA	Selective Primer	5'-GACTGCGTA

359 Table 4. Selective Primer combinations used for AFLP analysis and number of marker bands
 360 obtained for each of six types of four-base pair primer sets.

Primer set	<i>Eco</i> RI	<i>Mse</i> I	Number of markers
1	CAAC	ACAA	93
2	CAAC	ACAG	112
3	CAAC	ACTC	54
4	CACA	ACAG	41
5	CACT	ACAG	86
6	CAGG	ACTC	83

361 Table 5. Analysis of Molecular Variance (AMOVA) results and fixation indices. Significance was
362 tested with 1,023 permutations. Group 1 was collected from sand sagebrush and fringed
363 sagebrush and Group 2 was collected from winterfat and big sagebrush.

Source of variation	d.f.	Sum of Squares	Variance Components	Percentage of variation
Among groups	1	200.253	4.69095 Va	5.30
Among populations within groups	2	257.326	6.99008 Vb	7.90
Within populations	28	2150.327	76.79740 Vc	86.80
Total	31	2607.906	88.47843	

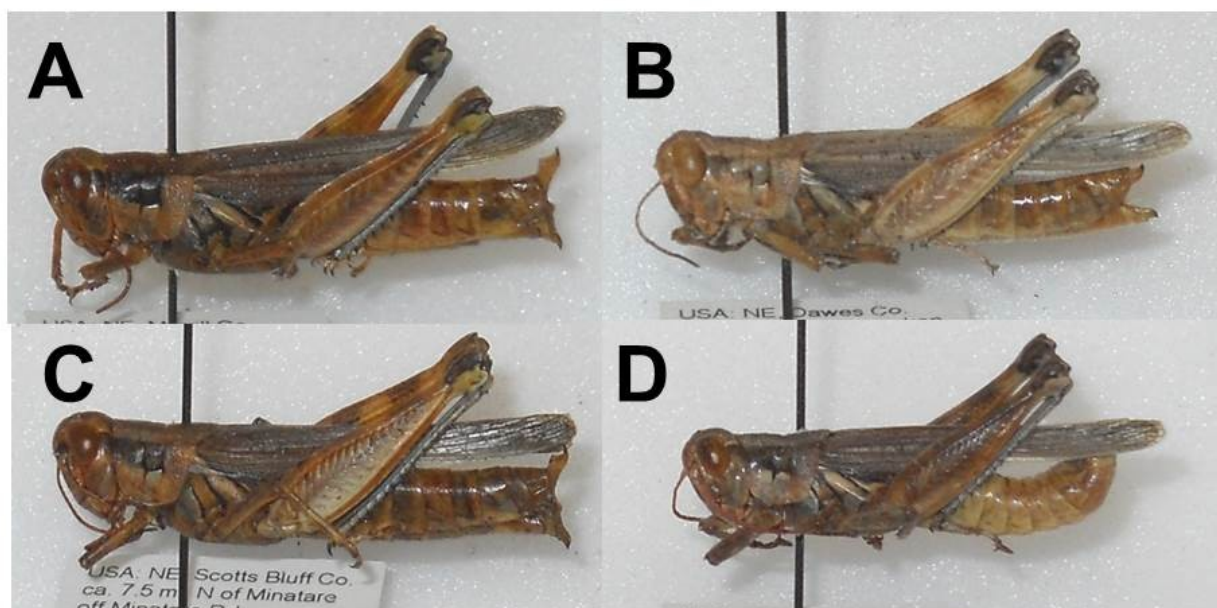
364 Fixation Indices

365 F_{ST} : 0.13202

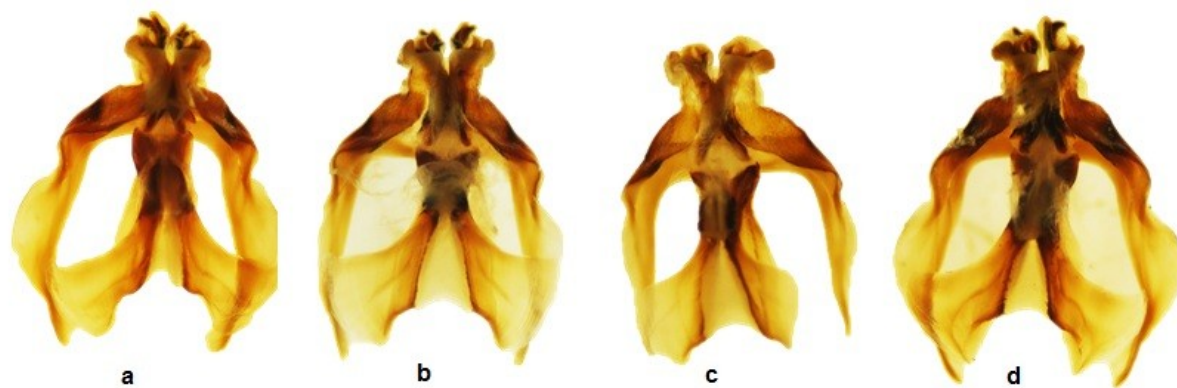
366 Table 6. Analysis of Nei's genetic diversity in subdivided populations. Low G_{ST} values suggest
 367 diversity among populations, and very high N_m values (>1.0) indicate significant gene flow
 368 between grasshopper populations. Group 1 was collected from sand sagebrush and fringed
 369 sagebrush and Group 2 was collected from winterfat and big sagebrush.

	Ht	Hs	G_{ST}	N_m
Group 1	0.2843	0.2266	0.2030	1.9630
Group 2	0.2862	0.2665	0.0690	6.7499
All populations	0.3127	0.2853	0.0879	5.1905

370 Ht = Total diversity
 371 Hs= Diversity within populations
 372 G_{ST} =Diversity among populations
 373 N_m = Estimate of gene flow based on G_{ST}



374 Figure 1. Lateral view of *Melanoplus bowditchii* grasshoppers collected feeding on A) sand
375 sagebrush, B) fringed sagebrush, C) winterfat, and D) big sagebrush.



376 Figure 2. Dorsal view of the aedeagus of *Melanoplus bowditchi* collected from (a) sand
377 sagebrush, (b) fringed sagebrush, (c) winterfat, and (d) big sagebrush.

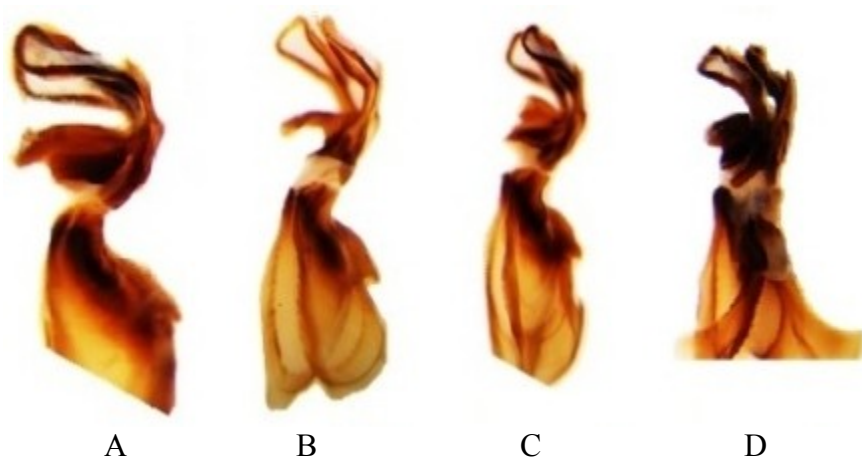
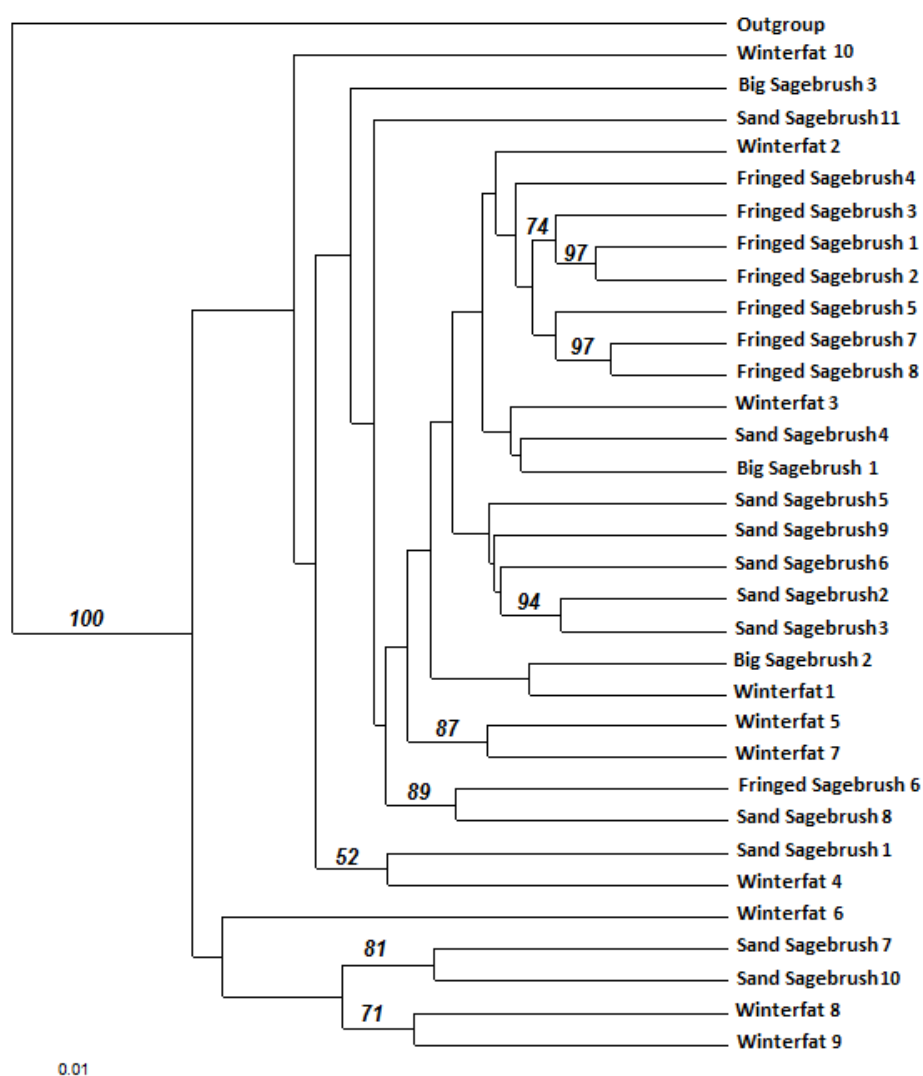


Figure 3. (A) Lateral view of the aedeagus of *Melanoplus bowditchi* collected from (a) sand sagebrush, (b) fringed sagebrush, (c) winterfat, and (d) big sagebrush..



381 Fig. 4. Distance-based Unweighted Pair Group Method with Arithmetic mean (UPGMA)
382 dendrogram of *M. bowditchi* grasshoppers using 1,000 bootstrap replicates. The dendrogram
383 shows the relationship among individuals. Numbers indicate bootstrap support >50% for
384 populations collected from different host plants.