Eutrema salsugineum (Cruciferae) new to Mexico: a surprising generic record for the flora of Middle America

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Abstract
The paper reports Eutrema salsugineum as a novelty to the flora of Mexico and Middle America in general. The finding stands ca. 1600 km apart from the closest known locality in the Rocky Mountains of Colorado, USA. The species is considered native to NW Mexico and its late discovery in the region is presumably explained by its tiny habit, early flowering time, and subephemeral life cycle. The phylogenetic position of this Mexican population in a haplotype network based on the chloroplast DNA fragment psbA-trnH confirms this hypothesis and also suggests, in contrast to the previously held viewpoint, multiple colonizations of North American continent from Asia.

Keywords
Brassicaceae, floristic finding, geographic disjunction, halophyte, model species, native distribution range, Thellungiella

Introduction

Eutrema salsugineum (Pall.) Al-Shehbaz & Warwick, long known as Thellungiella salsuginea (Pall.) O.E. Schulz, is, along with E. edwardsii R. Br., the most widely distributed representative of the genus Eutrema R. Br., occurring in temperate and, to a lesser extent, the Arctic zone of Eurasia and North America. Being an obligatory halophytic annual...
related to *Arabidopsis thaliana* (L.) Heynh., it has become a new model for studying abiotic stress (salt, cold, and draught) tolerance in plants (Bressan et al. 2001; Amtmann 2009). As a result, *Eutrema salsugineum* became one of the first representatives of Brassicaceae to have its genome sequenced (Wu et al. 2012). Nevertheless, fundamental (e. g., geographic) information on the species is incomplete (Koch and German 2013). Due to the peculiar ecological requirements of the species, *E. salsugineum* has a highly disjunct distribution in both continents (Korobkov 1975; Ovchinnikova 1994; Rollins 1993; Al-Shehbaz 2010). Despite being described in the 18th century (Pallas 1773) and being noted for its geographic disposition (Schulz 1924), the exact distribution of this species remained unknown, until recently. One of the reasons for the uncertainty was the misidentification of multiple specimens from Russia and Kazakhstan incorrectly referred to as “*Thellungiella salsuginea*”. Of these, all specimens from the western-most part of the range belong to another species, *E. botschantzevii* (D.A. German) Al-Shehbaz & Warwick [*Thellungiella botschantzevii* D.A. German] (German 2002, 2006, 2008). This finding now means that *E. salsugineum* is completely absent from Europe being bounded at the west by the lower reaches of Amu Darya river, Aral Sea, and the Turgai Valley. Our knowledge on the distribution of the species in America had also undergone improvements, and nowadays *E. salsugineum*’s range is known to extend discontinuously from boreal and Arctic Canada (Alberta, Manitoba, Northwest Territories, Saskatchewan, Yukon) down through the Rocky Mountains within British Columbia, and south to Montana and Colorado. Previous reports for other states such as Idaho (Rydberg 1917) and Ontario (Scoggan 1957) were not confirmed by subsequent studies (Al-Shehbaz 2010; Brouillet et al. 2010). Here we report of *Eutrema salsugineum* specimens collected by the second author in 2010 from arid regions in north-eastern Mexico, revealing another highly disjunct and we believe indigenous population. Our findings are supported with additional genetic analyses from which we draw further biogeographic conclusions.

**Materials and methods**

The specimen documenting the occurrence of *E. salsugineum* in Middle America is deposited in HEID. For estimation of the newly revealed disjunction in the distribution area of the species, available information regarding the closest occurrence has been used (Weber 1966). Results were interpreted in light of the most recent phylogeographic study of *E. salsugineum* by Wang et al. (2015). In this study various plastid genes (*ndhF1, ndhF2, psbA-trnH, rpoC1, rbcL, trnD-trnT, trnL, trnL-trnF, trnV*) have been sequenced and DNA polymorphisms have been detected in one marker only. In order to find the inter-species affinity of the Mexican population, its position in the haplotype network based on the respective variable chloroplast DNA fragment, *psbA-trnH*, was determined. For this purpose, representatives of the two geographically closest populations (from Montana and Colorado) not studied molecularly before and one accession from Canada, as internal control, were also sequenced.

Total DNA was extracted from 100 mg of herbarium tissue using the Invisorb Spin Plant Mini Kit (Stratec Biomedical AG, Birkenfeld, Germany). PCR ampli-
fication was performed in a volume of 25 μL, using 10 μM of each primer, respectively, a total of 2.0 mM MgCl₂ and 0.5 U of Mango-Taq polymerase (Bioline, Luckenwalde, Germany). The primers used for amplification were psbA-for: 5′-GTG ATG CAT GAA CGT AAT GCT C-3′, and trnH-rev: 5′-CGC GCA TGG TGG ATT CAC AAT CC-3′). All primers were extended by the M13 sequence for subsequent sequencing using M13 universal sequencing primers. The amplifications were run on a PTC 200 Peltier Thermal Cycler (MJ Research, Waltham, Massachusetts, USA) under the following conditions: 3 min initial denaturation at 95°C; 30 cycles of amplification with 30s at 95°C, 30s at 50°C, and 1 min at 72°C; and 5 min of final elongation at 72°C. PCR success was checked with electrophoresis in a 1% agarose gel in TAE-buffer. PCR product clean-up was executed using the Wizard SV Gel and PCR Clean-Up System (Promega, Madison, USA). Custom Sanger-sequencing was performed with GATC-Biotech (Konstanz, Germany). The electropherograms were checked and trimmed to the borders of the analysed markers using the program SeqMan DNA-Star Lasergene software package (DNASTAR, Madison, Wisconsin, USA).

DNA sequence variation was compared with cpDNA haplotypes detected by Wang et al. (2015). Haplotype network analysis was conducted with SplitsTree4 vers. 4.14.4 (Huson and Bryant 2006) with gaps treated as additional binary character.

Apart from the Mexican one, the newly sequenced specimens are: 1) USA, Colorado, Park Co., 11 August 1965. W.A. Weber 12925 (GH); 2) USA, Montana, Beaverhead Co., 20 June 1920. E.B. Payson & L.B. Payson 1730 (GH); 3) Canada, Saskatchewan, Jameson, 15 miles south of Regina, 15 June 1983. G.F. Ledingham 7937 (MO).

**Results and discussion**

*Eutrema salsugineum* (Pall.) Al-Shehbaz & Warwick, Harvard Pap. Bot. 10(2): 134. 2005. Described from: [NE Kazakhstan], …circa lacus et lacunas sale praesertim amaro abundantes ad Irtin inter fortalitia Shelesenka et Jamyschewa. Lectotype (designated by German 2011: 52): [Kazakhstan, Pavlodar province, Irtysch valley], “*Sisymbr. salsuginosum*. [29 May 1771, *P. S. Pallas s.n.*], Herb. Pallas. Herb. Fischer” (LE!) ≡ *Sisymbrium salsugineum* Pall., Reise Russ. Reichs 2: 466, 740, tab. V. 1773. ≡ *Arabidopsis salsuginea* (Pall.) N. Busch, Fl. Sibir. Orient. Extr. 1: 136. 1913. ≡ *Thellungiella salsuginea* (Pall.) O.E. Schulz in Engler, Pflanzenreich 86 (4, 105): 252. 1924.

As taxonomy/nomenclature is not the focus of the present study, only most frequently used synonyms are given here. For detailed synonymy, Schulz (1924: 252) and Tropicos (tropicos.org) can be consulted.

New locality: “Mexico: Est. Nuevo León, Los Enebros, Cerro el Potosi, Sierra Madre Oriental, saline-sodic soils, limestone. 24°52′35″N, 100°23′38″W, 1882 m a.s.l., 14 March 2010. Marcus Koch s.n.” (HEID 501412: http://gartenbank.cos.uni-heidelberg.de/img/HEID501412).

This is the first record of *Eutrema salsugineum* (and the genus) from Middle America (specifically Mexico) which shifts the southern limit of the distribution of the species
ca. 1600 km to south-southeast from the closest, isolated fragment of the distribution area of the species, confined to Park County, Colorado (Weber 1966). This introduces a hitherto unrecognized disjunction within the New World part of *Eutrema salsugineum*’s range. In Sierra Madre Oriental the plants are found at elevations of ca. 1880 m a.s.l. in a semi-open habitat among halophytic, oak/ *Juniperus* (*J. flaccida*)- and grass-dominated vegetation on alkaline soil which fits well the ecological requirements of the species (Fig. 1). The habitat at the Mexican location of *E. salsugineum* is apparently natural and shows no obvious signs of anthropogenic modification. Having in mind the naturally disjunct biogeography of this species we consider it as a natural element of the flora of Sierra Madre Oriental which was overlooked by previous studies due to

Figure 1. Habitat of Mexican accession of *Eutrema salsugineum* (Los Enebros, Cerro el Potosi, Sierra Madre Oriental). **A, B** General view **C–E** Closer look of *E. salsugineum* and co-occurring Brassicaceae species with herbarium numbers of respective specimens: **C** *Descurainia pinnata* (Walter) Britton s. l. (HEID 501409 to 501411) **D** *Eutrema salsugineum* (HEID 501412) **E** *Lepidium alyssoides* A. Gray (HEID 501414, 501415) **F** *Nerisyrenia linearisfolia* (S. Watson) Greene (HEID 501413). All images were taken on March 14th 2010 by M.A. Koch.
its tiny habit, trivial appearance, early flowering time, and short, almost ephemeral life cycle. Results of the haplotype analysis (below) apparently confirm this conclusion.

DNA sequence data revealed a new plastid DNA haplotype of Mexican *E. salsugineum* indicating its genetic distinctiveness. The haplotype code is indicated in Table 1 summarizing also haplotypes detected earlier (Wang et al. 2015). The geographic distribution of the various accessions and their plastid haplotypes are given in Fig. 2. SplitTree analysis indicates that the Mexican plastid haplotype is most closely related to haplotype H5 (numbering following Wang et al. 2015), which was originally found only once in Tuva (South Central Siberia) (Fig. 2; Wang et al. 2015). However, the same H5 haplotype was found here in Montana and Colorado, largely representing the mountain regions of the western United States. As expected, the Canadian sample from Saskatchewan analyzed herein carries haplotype H2, which is widely distributed in NE China and the amphi-Beringian region (Fig. 2c).

In their recent phylogeographic study Wang et al. (2015) discovered extremely low level of genetic diversity throughout the whole distribution area of *E. salsugineum*, especially in NE Asia/America, apparently reflecting a very recent (ultimately Late-Pleistocene/very Early Holocene) range expansion. These authors suggested wind to be the main agent mediating the fast and long-distance dispersal of *E. salsugineum* and a single colonization of the New World by the species (Wang et al. 2015). However, a present-day fragmentary sublongitudinal distribution of *E. salsugineum* in the Rocky Mountains along with our finding that relevant populations carry a haplotype (H5) which is identical to a unique haplotype found in South Siberia, suggests that the North American continent has been colonized independently two times. This conclusion is supported by the fact that the newly described accession occurring in Mexico carries a haplotype which most likely derived directly from haplotype H5. This overlapping distribution pattern of an amphi-Beringian group carrying H1/H2 and a disjunct group (South Siberia versus western North America) carrying H5 plus the Mexican type can also be best explained by a first and older immigration of *E. salsugineum*

**Table 1.** Variable sites of the polymorphic cpDNA fragment *psbA-trnH* in *Eutrema salsugineum* and closely related species (modified from Wang et al. 2015). GenBank code of the Mexican haplotype is KY073435.

| Species             | Haplotype | psbA-trnH  |
|---------------------|-----------|------------|
|                     |           | 43 108 129 156 162 208 277 304 |
| *E. salsugineum*    |           |            |
| H1                  | A C       | TGAATTT A TTTCTAT A - C |
| H2                  | A C       | TGAATTT A TTTCTAT A - A |
| H3                  | A C       | TGAATTT A ATAGAAA A - A |
| H4                  | A C       | TGAATTT A TTTCTAT C A A |
| H5                  | A C       | TAAATC G ATAGAAA A A A |
| H6                  | A T       | TGAATTT A TTTCTAT A - A |
| Mexican             | A C       | TAAATC A TTTCTAT A A A |
| *E. halophilum*     | H7        | A C       | TGAATTT A TTTCTAT A A A |
| *E. botschantzevii* | H8        | G C       | TGAATTT A TTTCTAT A A A |
Figure 2. Distribution and relationships of *E. salsugineum* haplotypes. **A** distribution of accessions analyzed herein and in an earlier study (modified from Wang et al. 2015) **B** SplitsTree graph showing genetic relationships among the nine haplotypes (Median-joining network: Bandelt et al. 1999) **C** detailed distribution of haplotypes H1 and H2 in North-West China (modified from Wang et al. 2015).

From Asia predating the last glaciation. During the Last Glaciation Maximum the northern populations in America apparently went extinct and/or migrated southwards. If this scenario is correct, the last latitudinal shift of vegetation belts could enable the species to reach the Middle Americas and subsequently survive there in appropriate habitats at higher elevation. The wide amphi-Beringian distribution of H1 and H2 might reflect postglacial expansion as demonstrated by the phylogeography inferred by Wang et al. (2015) based on nuclear genes. This multiple immigration pattern of various temperate-cold and often draught adapted Brassicaceae taxa between East Asia...
and the North America has been documented e.g. also for *Arabidopsis* and *Arabis* L. (Schmickl et al. 2008, 2010; Koch et al. 2010; Koch and Grosser 2017), highlighting the general significance of this spatio-temporal pattern.

**Conclusion**

As shown herein, Asian-North American halophyte *Eutrema salsugineum* has a far more southern distribution in the New World than previously considered almost reaching the Northern Tropic. The character of the discovered habitat suggests native rather than human-mediated occurrence of the species in Mexico, a conclusion supported by molecular footprints. A *psbA-trnH*-based haplotype network demonstrates more complicated infraspecific structure and biogeographic history of *E. salsugineum* than previously thought and suggests its multiple invasions to the New World.

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**References**

Al-Shehbaz IA (2010) *Eutrema* R. Brown. In: Flora of North America Editorial Committee (Ed.) Flora of North America North of Mexico, vol. 7. Oxford University Press, New York & Oxford, 555–556.

Amtmann A (2009) Learning from evolution: *Thellungiella* generates new knowledge on essential and critical components of abiotic stress tolerance in plants. Molecular Plant 2: 3–12. https://doi.org/10.1093/mp/ssn094

Bressan RA, Zhang C, Zhang H, Hasegawa PM, Bohnert HJ, Zhu JK (2001) Learning from the *Arabidopsis* experience: the next gene search paradigm. Plant Physiology 127: 1354–1360. https://doi.org/10.1109/10.1093/mp/ssn094

Bandelt HJ, Forster P, Röhl A (1999) Median-joining networks for inferring intraspecific phylogenies. Molecular Biology & Evolution 16(1): 37–48. https://doi.org/10.1093/oxfordjournals.molbev.a026036

Brouillet L, Coursol F, Meades SJ, Favreau M, Anions M, Bélisle P, Desmet P (2010+) *Eutrema salsugineum* (Pallas) Al-Shehbaz & Warwick. In: VASCAN, the Database of Vascular Plants of Canada. http://data.canadensys.net/vascan/taxon/4131 [accessed 4. 11. 2016]
German DA (2002) About the genus *Thellungiella* O. E. Schulz (Cruciferae). Turczaninowia 5, 2: 32–41. http://ssbg.asu.ru/turcz/p32-41.pdf

German DA (2006) Additions to Cruciferae of flora of Kazakhstan. Botanicheskii Zhurnal (Moscow & St. Petersburg) 91(8): 1198–1211.

German DA (2008) Genus *Thellungiella* (Cruciferae) in Europe. Botanicheskii Zhurnal (Moscow & St. Petersburg) 93(8): 1273–1280.

German DA (2011) Typification of selected names of Cruciferae taxa from Siberia and some neighboring regions. Turczaninowia 14(1): 45–54. http://ssbg.asu.ru/turcz/turcz_14_1_45-54.pdf

Huson DH, Bryant D (2006) Application of phylogenetic networks in evolutionary studies. Molecular Biology & Evolution 23(2): 254–267. https://doi.org/10.1093/molbev/msj030

Koch MA, German DA (2013) Taxonomy and systematics are key to biological information: *Arabidopsis, Eutrema* (*Thellungiella*), *Noccaea* and *Schrenkiella* (Brassicaceae) as examples. Frontiers in Plant Science 4: 267. https://doi.org/10.3389/fpls.2013.00267

Koch MA, Grosser J (in revision) East Asian *Arabis* species (Brassicaceae) exemplify past hybridization and subsequent emergence of three main evolutionary lineages in East Asia, America and the amphiberingian region.

Koch MA, Karl R, Kiefer C, Al-Shehbaz IA (2010) Colonizing the American continent: Systematics of the genus *Arabis* in North America (Brassicaceae). American Journal of Botany 97(6): 1040–1057. https://doi.org/10.3732/ajb.0900366

Korobkov AA (1975) *Thellungiella* O. E. Schulz. In: Tolmatschev AI (Ed.) Flora arctica URSS, vol. 7. Nauka, Leningrad, 52–53.

Ovchinnikova SV (1994) *Thellungiella* O. E. Schulz. In: Malyschev LI, Peschkova GA (Eds) Flora Sibiriae, vol. 7. Nauka, Novosibirsk, 59–60.

Pallas PS (1773) Reise durch verschiedene Provinzen des Rußischen Reichs, T. 2. Kayserliche Academie der Wißenschaften, St. Petersburg, 744 pp.

Rollins RC (1993) The Cruciferae of Continental North America. Stanford University Press, Stanford, 996 pp.

Ryderberg A (1917) Flora of the Rocky Mountains and Adjacent Plains: Colorado, Utah, Wyoming, Idaho, Montana, Saskatchewan, Alberta, and Neighboring Parts of Nebraska, South Dakota, North Dakota, and British Columbia. The author, New York, 1100 pp.

Schmickl R, Jørgensen MH, Brysting AK, Koch MA (2008) Phylogeographic implications for North American arctic *Arabidopsis lyrata*. Plant Ecology & Diversity 1(2): 245–254. https://doi.org/10.1080/17550870802349138

Schmickl R, Jørgensen MH, Brysting AK, Koch MA (2010) The evolutionary history of the *Arabidopsis lyrata* complex: A hybrid in the amphi-Beringian area closes a large distribution gap and builds up a genetic barrier. BMC Evolutionary Biology 10: e98. https://doi.org/10.1186/1471-2148-10-98

Schulz OE (1924) Cruciferae-Sisymbrieae. In: Engler A (Ed.) Das Pflanzenreich. 86(IV, 105), 1–388.

Scoggan HJ (1957) Flora of Manitoba. Department of Northern Affairs and National Resources, Ottawa, 619 pp.

Tropicos.org. Missouri Botanical Garden. http://www.tropicos.org [accessed 20. 6. 2016]
Wang X-J, Shi D-C, Wang X-Y, Wang J, Sun Y-S, Liu J-Q (2015) Evolutionary Migration of the Disjunct Salt Cress *Eutrema salsugineum* (= *Thellungiella salsuginea*, Brassicaceae) between Asia and North America. PLoS One 10(5): e0124010. https://doi.org/10.1371/journal.pone.0124010

Weber WA (1966) Additions to the flora of Colorado – IV. University of Colorado Studies. Series in Biology 23: 37[1–24]. http://scholar.colorado.edu/sbio/37/

Wu H-J, Zhang ZG, Wang J-Y, Oh D-H, Dassanayake M, Liu B, Huang Q, Sun H-X, Xia R, Wu Y, Wang Y-N, Yang Z, Liu Y, Zhang W, Zhang H, Chu J, Yan C, Fang S, Zhang J, Wang Y, Zhang F, Wang G, Yeo S, Cheeseman JM, Yang B, Li B, Min J, Yang L, Wang J, Chu c, Chen S-Y, Bohnert HJ, Zhu J-K, Wang X-J, Xie Q (2012) Insights into salt tolerance from the genome of *Thellungiella salsuginea*. Proceedings of the National Academy of Sciences of the United States of America 109: 12219–12224. https://doi.org/10.1073/pnas.1209954109
