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Phylogenetic relationships of the bumblebees *Bombus* moderatus, B. albocinctus, B. burjaeticus, B. florilegus and B. cryptarum based on mitochondrial DNA markers: a complex of closely related taxa with circumpolar distribution

(Hymenoptera: Apidae: Bombus)

With 6 figures and 4 tables

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Zusammenfassung

Königinnen von Bombus moderatus aus Alaska/USA und Alberta/Canada und von B. burjaeticus and B. patagiatus aus dem Russischen Transbaikal wurden an verschiedenen Orten im Frühjahr gefangen, um künstliche Kolonien zu züchten. Zusätzlich wurden Männchen von B. florilegus in Hokkaido/Japan gesammelt. Teilsequenzen (Länge 1005 bp) mitochondrialer Cytochrome Oxidase Untereinheit I (COI) wurde sequenziert. Zum Vergleich wurden auch Museumsproben von B. albocinctus und B. burjaeticus sequenziert. Die Divergenz der Sequenzen innerhalb der Taxa beträgt 1 bis 2 Basen-Substitutionen und die Tamura-Nei Genetische Distanz 0.001-0.002. Die Divergenz der Sequenzen zwischen B. moderatus, B. albocinctus und B. burjaeticus beträgt nur 1-5 Basen-Substitutionen und die Tamura-Nei Genetische Distanz 0.001-0.005, während die Divergenz der Sequenzen zwischen B. lucorum, B. magnus, B. patagiatus und B. cryptarum 22-44 Basen Substitutionen beträgt und die Tamura-Nei Genetische Distanz 0.027-0.042. Zusätzlich zu den Clustern für B. lucorum, B. magnus und B. patagiatus zeigt die Topologie des Phylogramms (MrBayes Maximum Likelihood Tree) ein umfangreiches Cluster, in dem B. albocinctus, B. burjaeticus, B. moderatus und B. florilegus vereinigt sind. Da die COI Sequenzen keine Lücken aufweisen, können die einzelnen Nukleotide wie homologe Positionen verwendet werden. Jedes Taxon besitzt 8-20 eigene Substitutionen, die als diagnostische Positionen zur Charakterisierung des Taxons verwendet werden können. Die Analyse der diagnostischen Positionen der Taxa bestätigt die Topologie des Maximum Likelihood Phylogramms. Um die Lücke zwischen den östlichsten bekannten Vorkommen von B. cryptarum im Kaukasus und im Elburz und den Vorkommen von B. burjaeticus/B.albocinctus im Russischen Transbaikal und in Russisch Fernost zu überbrücken, wurden 12 weitere Museumsproben aus Zentralasiatischen Gebirgen und dem Himalaja sequenziert. Durch Analyse der diagnostischen Positionen der teilweise 100 Jahre alten DNA kann gezeigt werden, dass keine dieser Proben zu B. lucorum gehören kann, alle bilden ein Cluster mit den Taxa des cryptarum Komplexes.

Summary

Spring queens of *Bombus moderatus* from Alaska/USA and Alberta/Canada, and of *B. burjaeticus* and *B. patagiatus* from the Russian Transbaikal region were collected at different localities. In addition, males of *B. florilegus* were collected from Hokkaido/Japan. Partial sequences (length 1005 bp) of mitochondrial cytochrome oxidase subunit I (COI) were sequenced from specimens from each locality and species. For comparison, museum specimens of *B. albocinctus* and *B. burjaeticus* were also sequenced. The intraspecific sequence divergence was only 1–2 base substitutions and about 0.001–0.002 in Tamura-Nei genetic dis-

tance. The interspecific sequence divergence between *B. moderatus, B. albocinctus* and *B. burjaeticus* was only 1–5 base substitutions and about 0.001–0.005 in Tamura-Nei genetic distance, whereas the sequence divergence between *B. lucorum, B. magnus, B. patagiatus* and *B. cryptarum* was about 24–44 base substitutions and approximately 0.027–0.042 in Tamura-Nei genetic distance. The MrBayes maximum likelihood tree generated a tree topology with three separate clusters for *B. lucorum, B. magnus*, and one large cluster which united *B. albocinctus, B. burjaeticus, B. moderatus* and *B. florilegus*. Because there are no gaps in the alignments of COI sequences, single nucleotide sites were used as positional homologies. Each taxon was characterised by about 8–20 substitutions which were unique ("private") and could be used as diagnostic characters to define and identify these taxa. An analysis of the number of diagnostic characters confirmed the clustering of the maximum likelihood tree. To bridge the gap of about 5000 km between the most eastern known localities for *B. cryptarum* in the Caucasus and Elburz Mountains and *B. burjaeticus/B. albocinctus* in the Russian Transbaikal region and the Russian Far East, 12 more museum specimens from the Central Asiatic Mountains and the Himalayas were sequenced. By analysing the diagnostic positions in the sequences of this almost 100-year-old museum DNA, it was shown that none were connected with *B. lucorum*: they all clustered within the *cryptarum*-complex taxa.

Introduction

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The taxon Bombus cryptarum (FABRICIUS 1775), initially described from the type locality Hafniae (Copenhagen), re-described as B. lucorum var. pseudocryptarum SKORIKOV (1913) from Poland and Russia, and as B. lucorum var. lucocryptarum BALL (1914) from Belgium, was established as a species by RASMONT (1981, 1983). It took another 25 years until this species, with a European distribution, was accepted as separate from B. lucorum by morphology, male labial gland secretions and DNA sequences (BERTSCH 1997; BERTSCH et al. 2004, 2005; BERTSCH 2009). Bombus cryptarum is distributed throughout Europe. It is more or less abundant in parts of Great Britain, the Netherlands, Benelux, parts of France, in Northern and Middle Germany, Scandinavia, Finland, Poland and Lithuania. The distribution in Belarus and Russia still has to be investigated, but museum specimens and fieldwork in St. Petersburg and Moscow show that the species is also common there. Museum specimens from Tscheljabinsk/Russia make its distribution throughout Northern Russia very probable, at least till the Urals. The available data also show that B. cryptarum is less abundant in the Southern parts of these countries. The abundance and distribution in the Alps of France, Switzerland and Austria still has to be fully investigated, otherwise the alpine distribution continues through the Balkan Mountains and the highlands of North Eastern Anatolia into the Caucasus and the Talesh and Elburz Mountains in Iran.

By investigating isoenzyme markers, SCHOLL et al. (1990) were able to show that the North American *B. moderatus* CRESSON, 1863, was closely related to the European *B. cryptarum*, a view confirmed by DNA sequences (BERTSCH et al. 2010). These sequences also showed that the Russian Far Eastern taxon *B. albocintus* SMITH, 1854, and the North American *B. moderatus* belong to the same taxon, and that this Pan-Pacific *B. albocinctus* is not the Far Eastern representative of *B. lucorum* as discussed by VOGT (1911), BISCHOFF (1930), KRÜGER (1951), TKALCU (1974), LELEJ & KUPIANSKAYA (2000), DAVYDOVA (2001), and DAVYDOVA & PESENKO (2002) but is closely related, if not conspecific, to the European *B. cryptarum*. Between the specimens *B. cryptarum* from the Urals, the Caucasus and the Talesh and Elburz Mountains, and the Russian Far Eastern specimens of *B. albocinctus/burjaeticus*, a gap of about 5000 km has to be closed.

To investigate the relationships between the European *B. cryptarum* and the Russian Far Eastern *B. albocinctus*, we obtained partial sequences from the mitochondrial cytochrome oxidase subunit

I (COI) gene of Bombus sensu stricto taxa from the Russian Transbaikal region, the Russian Far East and museum specimens from the Central Asiatic Mountains (Alai, Kirghizian and Dzungarian Alatau) and the Himalayas, and addressed the following issues: (1) whether the Russian Far East Genbank sequences designated as B. cryptarum were identical with the taxon B. albocinctus; (2) whether there were close relationships between B. albocinctus and the Siberian taxon B. patagiatus NYLANDER, 1948, the Transbaikal region taxon B. burjaeticus KRüger, 1951 and the taxon B. florilegus PANFILOV, 1956 (= *B. terrestris var. japonica* FRIESE 1909) from the Kuril Islands and Hokkaido; and (3) whether it was possible to close the gap between the Transbaikal region taxa and the B. cryptarum specimens from Anatolia, the Caucasus and Elburz Mountains using museum specimens from the Central Asiatic Mountains.

Materials and Methods

Bumblebee samples

Table 1 shows the bumblebee samples used for DNA sequencing with their identification codes and the localities of their origins with geographical coordinates. Figure 1 shows a pacific-centred world map with the distributions of B. cryptarum, B. burjaeticus, B. albocintus, and B. moderatus. Figure 6 shows the distribution of B. florilegus between Hokkaido/Japan and the North Kuril Islands. From the museum collections, 18 specimens were also sequenced and Table 2 gives their identification codes, their original designations and the localities of their origins with geographical coordinates. The localities for these museum specimens with their identification numbers are also included in Figure 1. In order to check the identity

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Code	Locality	Country	Region	Latitude	Longitude	Altitude	
Mag-01	Glenmore Forest	UK	Scotland	57° 09.83' N	03° 41.31' ₩	334 m	Q, $aC \rightarrow M$
Mag-02	Menz	Germany	Brandenburg	53° 06.84' N	12° 68.54' E	85 m	AY530015
Mag-03	Marcinkonys	Lithuania	Alytus County	54° 21.04' N	24° 25.46' E	145 m	Q, aC \rightarrow M
Cry-01	Loch Swannay	UK	Orkney Mainland	59° 08.53' N	03° 12.24' E	51 m	Q, aC \rightarrow M
Cry-02	Hillside	UK	Burray/Orkney Isl.	58° 51.29' N	02° 56.29' E	50 m	Q, $aC \rightarrow M$
Cry-03	Porlock Hill	UK	Devon/England	51° 09.48' N	03° 34.64' W	417 m	Q, aC → M
Cry-04	Roth	Germany	Franken/Bayern	59° 59.87' N	09° 17.03' E	231 m	Q
Cry-05	Menz	Germany	Brandenburg	54° 21.04' N	24° 25.46' E	145 m	Q, aC \rightarrow M
Cry-06	Menz	Germany	Brandenburg	54° 21.04' N	24° 25.46' E	145 m	AY530012
Cry-07	Marcinkonys	Lithuania	Alytus County	54° 21.04' N	24° 25.46' E	145 m	Q, aC \rightarrow M
Cry-08	Vent	Austria	Ötztal/Tirol	46° 52.07' N	10° 54.44' E	2320 m	М
Cry-09	Nassfeld	Austria	Kärnten	46° 34.49' N	13° 06.26' E	1415 m	Q
Cry-10	Sölk Pass	Austria	Steiermark	46° 52.07' N	10° 54.44' E	1780 m	AY181124
Cry-11	Julier Pass	Switzer- land	Albula/Graubünden	46° 28.16' N	09° 43.31' E	2280 m	AY181124
Cry-12	Five Ness/Crail	UK	Five/Scotland	56° 16.41' N	02° 35.15' ₩	10 m	$Q, aC \rightarrow M$
Cry-13	Duncansby Head	UK	Caithness/Scotland	58° 38.34' N	03° 01.46' W	50 m	$Q, aC \rightarrow M$
Cry-14	Duncansby Head	UK	Caithness/Scotland	58° 38.34' N	03° 01.46' W	50 m	AY530011
FLO-01	Iturup/Etoforu	Russia	Kurilsky Distr.	45° 05.45' N	147° 52.43' E		AF279487
Flo-02	Nemuro	Japon	Hokkaido	42° 21.40' N	145° 44.50' E		AF279486
FLO-03	Tomoshiri	Japon	Hokkaido	43° 19.35 'N	145° 39.57' E	25 m	М
Flo-04	Tomoshiri	Japon	Hokkaido	43° 19.35 'N	145° 39.57' E	25 m	М
Alb-01	Kamchatka	Russia	Kamchatka Krai	-20-642 - High-Cold (2006 - 575 - 51		10 1734. 004-014	AF279482
Alb-02	Paramushir	Russia	Sakhalin Obl.	50° 39.48' N	156° 05.45' E		AF279483
Alb-04	Shumshu	Russia	Sakhalin Obl.	50° 44.46' N	156° 22.41' E		AF279484
Alb-04	Magadan	Russia	Magadan Obl.	59° 38.07' N	150° 49.05' E		AF279485
Мод-01	Denali	USA	Alaska	62° 39,91' N	150° 20.42'W	195 m	Q, aC \rightarrow M
Mod-02	Broad Pass	USA	Alaska	63° 19.31' N	149° 09.21'W	725 m	Q, aC \rightarrow M
Mod-03	Paxon	USA	Alaska	63° 00.94' N	145° 31.05'W	950 m	$Q, aC \rightarrow M$
Mod-04	Isabel Pass	USA	Alaska	63° 11.77' N	145° 33.64'W	1095 m	$Q, aC \rightarrow M$
Mod-05	Isabel Pass	USA	Alaska	63° 11.77' N	145° 33.64'W	1095 m	Q
Mod-06	Ya-Ha-Tinda	Canada	Alberta	51° 44.57' N	115° 32.52'W	1615 m	М
Mod-07	Sheep River	Canada	Alberta	50° 39.11' N	114° 21.85'W	1315 m	М
Bur-01	Listvyanka	Russia	Irkutsk Obl.	51° 52.25' N	104° 50.15' E	620 m	$Q, aC \rightarrow M$
Bur-02	Yablonowo	Russia	Chitinsky Distr.	51° 50.58' N	112° 45.20' E	965 m	$Q, aC \rightarrow M$
Bur-03	Ozero Tasey	Russia	Chitinsky Distr.	52° 14.29' N	112° 57.05' E	1005 m	Q
Bur-04	Chita	Russia	Zabaikalsky Krai	52° 00.54' N	113° 28.35' E	750 m	$Q, aC \rightarrow M$
Bur-05	Chita, Kadala Airport	Russia	Zabaikalsky Krai	52° 01.15' N	113° 18.22' E	675 m	Q
Bur-06	Kadachta	Russia	Karymski Distr.	51° 37.22' N	114° 14.56' E	650 m	$Q, aC \rightarrow M$
Bur-07	Kadachta	Russia	Karymski Distr.	51° 37.22' N	114° 14.56' E	650 m	Q
Luc-01	Chita	Russia	Chitinskaja Obl.	52° 00.86' N	113° 28.56' E	730 m	Q, aC \rightarrow M
Luc-02	Yakutsk	Russia	Sakha Republic	62° 01.82' N	129° 44.09' E	115 m	AF279497
Luc-03	Bræstrud	Norway	Buskerud	60° 18.22' N	08° 34.19' E	920 m	AY181120
Pat-01	Yakutsk	Russia	Sakha Republic	62° 01.82' N	129° 44.09' E	115 m	AF279499
Pat-02	Primorsk	Russia	Primorsky Krai				AF279498
Pat-03	Listvyanka	Russia	Irkutsk Obl.	51° 52.25' N	104° 50.15' E	620 m	Q, aC \rightarrow M
Pat-04	Chita	Russia	Zabaikalsky Krai	52° 00.54' N	113° 28.35' E	750 m	$Q, aC \rightarrow M$

Tab. 1: List of 40 *Bombus* samples (MAG = *magnus*, CRY = *cryptarum*, FLO = *florilegus*, ALB = *albocinctus*, MOD = *moderatus*, BUR = *burjaeticus*, PAT = *patagiatus*, and LUC = *lucorum*) used in the present analysis with identification codes, and collection locality information. Q = queen, aC \rightarrow M = artificial colonies with production of males.

of the specimens BUR-01–BUR-07 from the Transbaikal region, tentatively designated as *B. burjaeticus* with the material from Krüger, two paratype specimens of *B. burjaeticus* from Krügers material were sequenced (M-13-1, M-13-2), and as the specimens of *B. albocinctus* from the Russian Far East (ALB-01–ALB-04) were deposited in GenBank by Ito and Tanaka as *B. cryptarum*, three specimens from the type locality Kamchatka for *B. albocinctus* (M-14–M-16) were sequenced for comparison. Voucher specimens have been deposited at the Senckenberg Deutsches Entomologisches Institut, Müncheberg, Germany.

Polymerase chain reaction (PCR) and DNA sequencing of mitochondrial COI

Total DNA was extracted from legs using the QIAamp* DNA Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer's specifications for tissue, and eluted in 150 µl of highly purified water (Ampuwa*, Fresenius Kabi, Bad Homburg, Germany). For sequence analysis, overlapping fragments (in all 1027 bp) of mitochondrial COI were amplified using primers specifically designed for Bombus. BO-1-fwd (5' TAGGATCACCAGATATAGC 3') and BO-K-rev (5' GAGCTCAAACAATAAATCC 3') resulted in the amplification of a 609 bp fragment, whereas BO-5-fwd (5' AATGAAAGAGGTAAAAAAGAAAC 3') and BO-A-rev (5' ATGTTGAGGGAAAAATGTTAT 3') resulted in the amplification of a 510 bp fragment. Polymerase Chain Reaction amplifications were performed as described (BERTSCH et al. 2010). A sample of 10 µl from each reaction was checked on a 1 % agarose gel. Polymerase Chain Reaction products were purified using the AMPure* PCR Purification Kit (Agencourt, Beverly, MA, USA). Both strands were sequenced for all specimens. Sequencing reactions were performed using ABI* BigDye Terminator version 3.1 chemistry (Applied Biosystems, Foster City, CA, USA) according to manufacturer's instructions and they were then analysed on an ABI 3100 sequencer (Applied Biosystems). Sequences were aligned using CLUSTALX. No gaps or poorly aligned regions occurred in the alignment, but missing characters were trimmed from the ends of the alignment to produce an equal sequence length of 1005 bp for all individuals (encoding 335 amino acids). Individual alignments were aligned against the complete COI gene sequence of Bombus ignitus between positions 262 and 1267 (GenBank DQ870926; CHA et al. 2007).

Degraded DNA from museum specimens

The 40–100 year old DNA from the museum specimens was more or less degraded. The integration of these sequences into MrBayes maximum likelihood simulations did not give useful results; depending on the grade of degradation, the specimens clustered in unpredictable ways. Therefore, the original ABI traces of these sequences had to be carefully inspected for degraded positions; whenever double peaks were detected the IUPAC Code for mixed bases (wobbles) was inserted. Where necessary, the Polymerase Chain Reaction was repeated with slightly adjusted amplification conditions, and by this procedure it was possible to get reliable sequences for the control specimens (M13-1, M13-2 and M14-M16), which were suitable for use in MrBayes simulations. About 97 % (863 out of 891) of the positions in the sequences from the fresh material were invariant, therefore, for MrBayes simulations with the rest of the museum specimens, the variability of the degraded DNA was reduced by deleting all the invariant triplets (of the fresh, not the degraded DNA) and only the triplets with parsimony informative positions were included in the analysis.

Code		Locality	Country	Latitude	Longitude	
M-01	lucorum terrestriformis Vogт, 1911:56	Bursa/Bakazak	Turkey	40° 06.09' N	29° 12.56' E	ZSM
M-02	<i>cryptarum armeniensis</i> Rasmont, 1984:154	Ilgaz Pass	Turkey	41° 03.94' N	33° 45.00' E	ZSM
M-03	lucorum (Linnaeus, 1761)	Mt. Bombak	Russia	43° 57.00' N	40° 60.00' E	ZMAS
M-04	lucorum	River Bezymyanka	Russia	??	??	ZMAS
M-05	<i>cryptarum iranicus</i> Krüger, 1951:196	Col de Rovra	Iran	38° 40.00' N	48° 21.00' E	ZSM
M-06	<i>lucorum alaiensis</i> Reinig, 1930: 107	Kisil Beles	Kyrgyzstan	40° 16.98' N	73° 16.19' E	ZMA
M-07	<i>magnus turkestanicus</i> Krüger, 1954:274	Naryn	Kyrgyzstan	41° 26.32' N	76° 01.34' E	ZMA
M-08	<i>magnus borochorensis</i> Krüger, 1954:273	Burchan Canyon	Kazachstan	44° 21.05' N	80° 04.80' E	ZMA
M-09	<i>lucorum jacobsoni</i> Skorikov, 1912:610	Sind Valley Baltal	India	34° 15.26' N	75° 25.00' E	ZMA
M-10	reinigi Tkalcu, 1974:322	Phoksumo Lake	Nepal	29° 10.43' N	82° 56.46' E	ZMF
M-11	<i>lucorum terrestricoloratus</i> Krüger, 1951:196	??	China	<u>}</u> }	?? ??	ZMA
M-12	lucorum mongolicus Vogt, 1909: 42	Irkutsk	Russia	52° 17.21' N	104° 18.45' E	ZMAS
M-13-1	burjaeticus Krüger, 1951:148	Kulskoje	Russia	52° 05.81' N	109° 43.24' E	ZMA
M-13-2	burjaeticus	Nertschinsk	Russia	51° 59.20' N	116° 35.12' E	ZMA
M-14	<i>lucorum albocinctus</i> SMITH, 1854: 397	Petropavlowsk	Russia	53° 01.20' N	158° 39.42' E	ZMAS
M-15	lucorum albocinctus	Kljutschi	Russia	56° 10.05' N	160° 51.21' E	ZMAS
M-16	lucorum albocinctus	Mouth river Opala	Russia	51° 40.87' N	156° 31.62' E	ZMAS
M-17	lucorum albocinctus	Cape Schmidt	Russia	68° 53.50' N	179° 25.00' E	ZMAS

Tab. 2: List of *Bombus* samples from Museum specimens (ZMA = ZOÖLOGICAL MUSEUM AMSTERDAM, ZSM = ZOOLOGISCHE STAATSAMMLUNGEN MÜNCHEN, ZMAS = ZOOLOGICAL MUSEUM RUSSIAN ACADEMY OF SCIENCE ST. PETERSBURG, SEM = SENCKENBERG MUSEUM FRANKFURT/M) used in the present analysis.

Analysis of sequence divergence of mitochondrial COI

The absolute numbers of substitutions were counted based on a pair-wise comparison of the sequences. The analysis of sequence diversity was performed using the maximum composite likelihood method of MEGA 4.0 (TAMURA et al. 2007; KUMAR et al. 2008). Because the distribution of nucleotides in the COI of Hymenoptera is known to be heterogeneous, with a strong A + T bias, we selected the Tamura-Nei model of base substitution (TAMURA & NEI 1993) which corrects this bias in its assumption of sequence evolution. The nucleotide frequencies and the parameters necessary for this model were estimated from the sequence data and Tamura-Nei genetic distances were calculated. The best maximum likelihood model was selected using the JMODELTEST (POSADA 2008). The tree topology was inferred by a maximum likelihood tree based on the general time-reversible model (GTR plus gamma), calculated by Bayesian analysis using MrBayes (HUELSENBECK & RONQUIST 2001). Tree topology was also calculated as a Minimum Evolution tree (ME) with bootstrap sampling using MEGA 4.0 (TAMURA et al. 2007). GENEIOUS PRO 4.5 (Biomatters Ltd.) was used to analyse the alignment and detect diagnostic positions and the GREENBUTTON plugin (InterGrid) was used to do the time consuming MrBayes calculations on a supercomputer cluster. The nucleotide changes along the branches

of cladograms were examined with MACCLADE 4.08 (MADDISON & MADDISON 2002). The COI sequence data of *B. soroeensis* (GenBank AY181159; PEDERSEN 2002) were used as the outgroup.

Results

Nucleotide frequencies and substitution parameters

The aligned data matrix of 36 sequences (*B. moderatus*, *B. albocinctus*, *B. florilegus* and *B. burjaeticus*) of 891 bp length (encoding 297 amino acids) included 863 (97 %) invariants and 28 variable sites. Of these 28 variable positions, 3 were parsimony uninformative (noise) and 25 parsimony informative (signal). However, differences in this pattern were evident in codon positions, where only 1 informative site was in first position and all of the other 24 informative variant sites were in third position. None of these sites was a replacement position; the singleton replacement position 1007 in the GenBank sequence AF279485 was most probably an amplification error or a miscoding lesion. Table 3 shows the pattern of nucleotide substitutions estimated from the data with the maximum composite likelihood model (MEGA). The nucleotide frequencies were 33.1 (A), 42.1 (T), 12.3 (C) and 12.5 (G), which proves the known strong A + T bias typical for sequences of Hymenoptera.

COI sequence divergence between and within species

Table 4 presents the matrix of genetic distances estimated by the Tamura-Nei model and as p-distances within and between the five taxa investigated: *B. cryptarum, B. florilegus, B. albocinctus, B. moderatus* and *B. burjaeticus*. For comparison and discussion the sequences of the European species, *B. lucorum* (LUC-01–LUC-03), *B. magnus* (MAG-01–MAG-03) and the Asiatic species *B. patagiatus* (PAT-01–PAT-03) were included in the analysis. The intraspecific genetic variability was low for all taxa (1–2 nucleotides, Tamura-Nei distance 0.001–0.002). By contrast, the interspecific genetic variability between *B. lucorum, B. magnus, B. patagiatus* and *B. cryptarum* was approximately one order of magnitude larger (25–44 nucleotides, Tamura-Nei distance 0.027–0.042).

The Tamura-Nei genetic distance between *B. burjaeticus* and *B. albocinctus* is only 0.002, both taxa are conspecific, and *B. moderatus*, with a Tamura-Nei genetic distance to *B. albocinctus/ B. burjaeticus* of 0.004, also belongs to the same taxon; the larger genetic distance manifested the geographic separation of this taxon by the Bering Strait. *Bombus florilegus*, with a Tamura-Nei genetic distance between 0.010 and 0.014 to *B. moderatus/B. albocinctus/B. burjaeticus* was clearly distinct but not a separate species. All these taxa had a Tamura-Nei genetic distance between 0.012 and 0.018 to *B. cryptarum*, again with the largest genetic distance for the geographically separated

B. moderatus, therefore it seemed to be useful to treat all these taxa as members of a closely related complex of taxa, the *cryptarum*-complex. *Bombus patagiatus* was well separated from this *cryptarum*-complex by a Tamura-Nei genetic distance of 0.031 to 0.040 and *B. magnus* by a Tamura-Nei genetic distance of 0.030 to 0.037, comparable with *B. lucorum*, which was separated by a Tamura-Nei genetic distance of 0.036 to 0.040. These Tamura-Nei genetic distances are larger by one order of magnitude and revealed that all three taxa are separate species.

Tab. 3: Maximum Composite Likelihood Estimate of the Pattern of Nucleotide Substitution. The transition/transversion rate ratios are $k_1 = 5.25$ (purines) and $k_2 = 8.24$ (pyramidines), the overall transition/transversion bias is R= 1.834.

	А	Т	С	G
А	3	4.74	1.39	7.37
Т	3.73	*	11.42	1.4
С	3.73	39.08	-	1.4
G	19.6	4.74	1.39	-

Tree building by maximum likelihood models

The maximum likelihood tree (Fig. 2) generated using the Bayesian Markov Chain Monte Carlo (MCMC) analysis with 1005 bp full-length sequences was based on the general time-reversible model of base substitution (GTR plus gamma). For comparison, GenBank data of the European taxa *B. lucorum*, *B. cryptarum* and *B. magnus* were included. This tree confirmed the results from the analysis of genetic distances: *B. magnus* and *B. patagiatus* are distinctly separated species, *B. cryptarum*, *B. florilegus* and the cluster γ_3 of *B. albocinctus*/*B. burjaeticus*/*B. moderatus* form a cluster γ of closely related taxa with the geographically separated taxon *B. moderatus* as the most evolved end unit. All clusters had a high posterior probability and were distinctly separate from *B. lucorum*.

Tab. 4: Mean genetic distance within and between taxa. Diagonal and Lower left: TAMURA-NEI model, rates among sites gamma distributed. Upper right: p-distance.

	Cry	Flo	Alb	Mod	Bur	Luc	Pat	Mag
cryptarum	0.001	0.012	0.014	0.018	0.015	0.037	0.029	0.029
florilegus	0.012	0.001	0.010	0.014	0.011	0.034	0.029	0.025
albocinctus	0.014	0.010	0.001	0.005	0.002	0.035	0.037	0.036
moderatus	0.018	0.014	0.005	0.001	0.004	0.037	0.038	0.030
burjaeticus	0.016	0.011	0.002	0.004	0.001	0.036	0.035	0.049
lucorum	0.040	0.036	0.038	0.039	0.039	0.002	0.044	0.033
patagiatus	0.031	0.031	0.036	0.042	0.041	0.038	0.001	0.036
magnus	0.031	0.027	0.030	0.031	0.031	0.031	0.035	0.002

Tree building by diagnostic characters

Because there are no gaps in the alignments of COI sequences, single nucleotide sites can be used as positional homologies (HILLIS 1994). The alignment file (Fig. 3) showed that each taxon was characterised by substitutions which were unique ("private") and could be used as diagnostic characters to define and identify this taxon (FUNK et al. 2001; BERTSCH 2009). In MACCLADE, all of the changes at the nodes and the diagnostic characters at the last branch of the terminal units can be investigated in detail, and a tree with the classical tools for morphological characters can be built (Fig. 4). The conclusions drawn from the genetic distance data and the clustering of the maximum likelihood tree were fully confirmed by analysis of the diagnostic characters. The taxa B. albocinctus, B. burjaeticus and B. moderatus shared four positions, and B. moderatus, as a geographically separated taxon, had two diagnostic positions of its own; the analysis of the diagnostic positions confirmed that these taxa are conspecific. Bombus florilegus is closely related and characterised by two diagnostic positions, which is typical for isolated island populations. The combined B. albocinctus/ B. burjaeticus/B. moderatus taxon and B. florilegus shared five positions and a cluster with B. cryptarum, which was separated by six diagnostic positions. The taxa B. patagiatus, B. magus and B. lucorum had 16, 12 and 17 unambiguous diagnostic characters, respectively, and are separate species.

Museum sequences B. albocinctus and B. burjaeticus

The unpublished sequences from Magadan, Kamchatka and the Northern Kuril Islands, Shumshu and Paramushir, submitted to GenBank in 2000 by Ito & Tanaka, were identified by morphological characteristics as specimens of *B. cryptarum* by Masao Ito. As so far there are no known specimens of typical *B. cryptarum* from these areas, it seemed necessary to prove the identity of these GenBank specimens with typical *B. albocinctus*. Therefore four typical specimens (collare, scutellum and second tergite white-coloured) of *B. albocinctus* from Kamchatka



Fig. 2: Tree topology calculated as Maximum-Likelihood tree using Bayesian MCMC analysis with the general time reversal model of base substitutions with gamma distribution.

(M-14, M-15, M-16) and Chuchotka (M-17) from the collection of the Zoological Museum of the Academy of Sciences St. Petersburg (ZMAS) were sequenced. Two queens from the type series Kulskoje (M-13-1 & M-13-2) of *B. burjaeticus*, from the Vogt collection of the Zoölogical

Museum Amsterdam (ZMA), corresponding in morphology with Krüger's original description (yellow of bands *greenish*, *upper part* of episternum yellow) were sequenced to check their identity with queens collected during fieldwork during June 1994 in Transbaikal region.

The 40 to100-year-old museum specimens had degraded DNA but the genetic connections were established by comparing the diagnostic positions (Fig. 5). As the Tamura-Nei genetic distance between *B. albocinctus* and *B. burjaeticus* was only 0.004 and both taxa are conspecific, the results for the museum specimens of both taxa are identical: they do not belong to *B. lucorum* as none of the 17 diagnostic positions of *B. lucorum* could be found. The sequences for the museum specimen *B. burjaeticus* (M-13-1 and M-13-2) and *B. albocinctus* (M-14 to M-17) shared the positions 328 (T \rightarrow C), 330 (A \rightarrow T), 751 (T \rightarrow C) and 846 (T \rightarrow G) with *B. cryptarum* and they differed from *B. cryptarum* by positions 531 (T \rightarrow C), 855 (T \rightarrow G), 818 (C \rightarrow T), and 1128 (T \rightarrow C), positions characteristic for the Asiatic taxa of this complex.

Degraded DNA of museum specimens

As the alignment of the fresh material showed that 97 % of the positions were invariable, we removed all invariable triplets from the alignment of the museum specimens and generated a maximum likelihood tree (Fig. 5) with the variable triplets using the Bayesian MCMC analysis with the general time-reversible model of base substitution (GTR plus gamma). For comparison, sequences of fresh material of *B. cryptarum*, *B. albocinctus*, *B. burjaeticus* and *B. florilegus* together with sequences of *B. lucorum*, *B. magnus* and *B. patagiatus* were included, as outgroup *B. ignitus* was used (GenBank DQ870926; CHA et al. 2007). This procedure implied the assumption that the museum specimens belong to one of the taxa investigated in this report.

The control specimens M-13-1 and M-13-2 of the type series *B. burjaeticus* and the control specimens M-14 to M-17 of *B. albocinctus* joined the cluster γ_3 with the fresh material of *B. moderatus | B. albocinctus | B. burjaeticus*, which proved that deletion of the invariable sites does not change the overall result. Depending on quality and length (sequences M-02 & M-05 were too short to be included) of the degraded DNA, the specimens M-01, M-03, M-04 from Turkey, the Caucasus and Iran, areas from where *B. cryptarum* has been identified, joined the *B. cryptarum* cluster γ_1 or the *cryptarum*-complex cluster γ . All of the remaining specimens from the Central Asiatic mountains and the Himalayas whether identified as subspecies of *B. lucorum* (M-06, M-09, M-12), as subspecies of *B. magnus* (M-07 and M-08) or as separate species *B. reinigi* (M-10) also joined the cluster γ with the specimens of the *cryptarum*-complex taxa. The posterior probabilities were high; therefore, the conclusion that these specimens are not connected to *B. lucorum* but rather to *B. cryptarum* respectively to the *cryptarum*-complex taxa is reasonable.

Discussion

Bombus patagiatus as a separate species

This species was first described by NYLANDER (1848) and afterwards was neglected for a long time until it was established as a good species by TKALCU (1967). The most obvious characteristic is the typical combination of yellow-white colouration (collare, scuttellum, tergite), which is not a very useful feature in bumblebee taxonomy of critical taxa. TKALCU (1967) described morphological characteristics for this taxon (form of labrum, punctuation of vertex and ocellar field, ocellar distance, proportions of flagellum segments) and the differences between these characters compared to those of *B. lucorum*, but nearly all of these characteristics were so similar that terms such as 'a little bit more' and 'a little bit larger' ore 'sometimes a bit larger' were the best that could be said

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Fig. 3: Alignment of all parsimonious informative triplets (with uninformative sites deleted -), and with a pointer for position number (numbered for total COI) and codon position. Diagnostic (= private) positions marked with colour green = Thymine, violet = Cytosine, red = Adenine and yellow = Guanine.



Fig. 4: Observed diagnostic character changes with position numbers mapped onto the Maximum-Likelihood tree. Black box = unambiguous *diagnostic character* change, grey box = ambiguous *diagnostic character* change, and white box = character change.

[original in French]. RASMONT (1984:141) refers to these characters as morphologically very similar to *B. cryptarum* and comes to the conclusion that 'the two species for the moment cannot be distinguished by their morphology alone' [original in French]. The male labial gland secretions (BERTSCH et al. unpublished results) and the DNA sequences confirmed a close relationship but they also showed also that *B. patagiatus*, with 14 unambiguous diagnostic positions, is a separate species. The sympatric distribution of *B. albocinctus* and *B. patagiatus* in parts of Russia (Yakutia, Primorsk and Sakhalin), where both taxa are abundant also showed their genetic separation.



Fig. 5: Tree topology calculated as Maximum-Likelihood tree using Bayesian MCMC analysis with the general time reversal model of base substitution and gamma distribution for degraded DNA of museum specimens, only parsimony informative triplets included.

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Is Bombus florilegus a separate species?

The taxon Bombus terrestris var. japonica FRIESE, 1909, described from Yesso (= Hokkaido) and re-described as *B. florilegus* PANFILOV, 1956 (*japonicus* = name preoccupied; Dalla Torre 1890) with morphological characters of the sculpture of workers and the male genitalia, is characterised by the completely black abdomen of the females. ITO & SAGAKAMI (1980) listed the morphological differences of females and males of B. florilegus compared to B. albocinctus and B. lucorum, but the variation in most of the characters remains unclear and many of them seem to overlap. Bombus florilegus is a taxon with a very restricted habitat. Figure 6 shows the distribution from Cape Nemuro at Hokkaido through the South Kuril Islands where B. florilegus is the most abundant bumblebee, with a contact zone with B. albocinctus of about 120 km on the small North Kuril Islands Shimushir, Ketoi and Rasshua (ITO & SAGAKAMI 1980; ITO & KURANASHI 2000; LELEJ & KUPIANSKAYA 2000). Compared to the wide distribution of most of the typical Bombus sensu stricto species, this restricted distribution is typical for island populations, as for instance, B. terrestris sassaricus from Sardinia or B. terrestris canariensis from the Canary Islands compared to B. terrestris from the continent. Such isolated island populations are often genetically separated and may have their own diagnostic characters. Bombus florilegus has two unambiguous diagnostic characters that separate this taxon from B. albocinctus, which is comparable to the three unambiguous diagnostic characters separating B. magnus magnus of the British Isles from B. magnus flavoscutellaris of the European continent (BERTSCH 2009) or island populations of B. ignitus from Japan compared to continental populations of *B. ignitus* from Korea (TOKORO et al. 2010). Bombus florilegus is a taxon within the cryptarum-complex, most probably with the status of a subspecies of B. albocinctus. Until further evidence is available it could be treated as B. albocinctus florilegus PANFILOV.

Bombus albocinctus, a Pan-Pacific species with four taxa

This species, described by SMITH (with type locality Kamchatka), was soon treated as the Far Eastern *B. lucorum*, the most obvious separating characters being the white colouration (collare, scutellum and second tergit). DAVYDOVA (2001) and DAVYDOVA & PESENKO (2002) could not find any morphological characters to separate *B. albocinctus* and *B. lucorum* and once more established *B. albocinctus* as the Far Eastern *B. lucorum*, separated only by the white colouration. But the difference in colouration is not always as clear as discussed in the literature; from the 15 queen specimens from Kamchatka (ZMAS, St.Petersburg) many were not clear white (as described by SMITH) but there was a graduation from clear white to light yellow and some were more or less citron yellow, not very different from typical *B. lucorum*. The same is true for 25 queen specimens (ZMAS, St. Petersburg) from Sakhalin, Magadan and Anadyr. Some of these specimens were clear white but some were citron yellow, at least in part, and all intermediate shades were available. But the Tamura-Nei genetic difference of the COI sequences of *B. albocinctus* and *B. lucorum* was in the order of magnitude of other *Bombus* sensu stricto species, therefore *B. albocinctus* cannot be the Far Eastern *B. lucorum*.

Bombus burjaeticus was described by Krüger (from type locality Kulskoje/Burjatia) mainly by characters of colouration. In the VoGT collection (ZMA, Amsterdam) there is a large number of specimens from the type locality Kulskoje (353 queens, collected 19.V.1928 by Klemm, Kulskoje, Uda Valley, *Pinus* forest), most in very good condition. A thorough investigation of the morphological characters still remains to be done, and it might well be that different taxa are included in this Kulskoje material, but one characteristic mentioned in Krüger's original description was quite distinctive: many specimens showed a typical greenish tint of the yellow parts of the colouration, and the dorsal part of the episternum was yellow, both characteristics typical for *B. cryptarum*.

Two queens from the type series conforming to the description of Krüger were sequenced (M-13-1 and M-13-2). The 80year-old DNA was degraded but none of the 17 diagnostic characters of B. lucorum could be found (Fig. 3). Spring queens collected in June 1997 from the surroundings of Chita in Transbaikal region and tentatively designated as B. burjaeticus also delivered COI sequences without any diagnostic characteristics of B. lucorum, but close connections to B. cryptarum were found. These findings were confirmed by the main component of the male labial gland (unpublished results BERTSCH et al.) from artificial colonies of these cf. B. burjaeticus spring queens, which was ethyl dodecanoate (MG = 214, RI = 1507), not ethyl 9-octadecenoate (MG = 254, RI = 1764) as found in B. lucorum (see BERTSCH et al. 2004).

In a short remark RASMONT et al. (1986:677) noted 'the resemblance of the sternite 8 of certain *B. burjaeticus* males with *B. cryptarum*' and they also noted (1986:678) that the females of *B. albocinctus* from Sakhalin



Fig. 6: Distribution of *B. florilegus* (red) from Cape Nemurol Hokkaido through the Southern Kuril Islands and of *B. albocinctus* (blue) from Kamchatka through the Northern Kuril Islands. Arrows = zone of contact.

are not *B. lucorum* but are morphologically closely conform to *B. cryptarum*. Specimens of *B. albocinctus* from Magadan, Kamchatka and the North Kuril Islands deposited in the year 2000 in GenBank by Ito & Tanaka have been identified by morphological characters as *B. cryptarum* (M. Ito in e-mail correspondence). All of these findings and the conclusions corresponded with the results of our sequencing: *B. albocinctus* and *B. burjaeticus* belong to the same taxon, the white colouration of some Kamchatka and Far Eastern specimens is not a specific characteristic and both taxa are genetically very close to the European *B. cryptarum*. And as the North American taxon *B. moderatus* also belongs to *B. albocinctus* (BERTSCH et al. 2010), we have a taxon with a Pan-Pacific distribution. Unless possible connections between the European *B. cryptarum* and these Far Eastern taxa are investigated in detail, the best taxonomic conclusions are to treat *B. burjaeticus* KRÜGER as a junior synonym of *B. albocinctus* SMITH and *B. moderatus* as *B. albocinctus moderatus* CRESSON.

Closing the gap: central Asiatic mountains as a bridge between Europe and the Transbaikal region

Field studies need to be undertaken to close the large gap of about 5000 km between the Elburtz Mountains and the Baikal area. Meanwhile, this can be attempted using the information available in the large museum collections. KRÜGER (1951) was the first to recognise that not all *B. lucorum*like specimens in Asia really belonged to *B. lucorum*. With a trained eye and a lot of experience, as demonstrated when he separated *B. magnus* (first described in detail as *B. latocinctus*, KRÜGER 1939) and *B. lucorum*, he took the chance to fix this insight by designating these specimens extra names. Krüger knew *B. cryptarum*, the type specimen of his *B. lucorum var. lehmanni* from Medingen/Niedersachsen (collected 12/14.IV.1939; Vogt Collection ZMA) was a typical *B. cryptarum*, but he did not treat the taxon as a separate species. Instead he designated all non-*lucorum* specimens from Asia as *B. magnus* (*B. magnus iranicus*, *B. magnus turkestanicus*, *B. magnus borochorensis*, *B. magnus mongolicus*, *B. magnus laevis*, and *B. magnus lantschouensis*). As yet, *B. magnus* has not been detected in Asia, therefore, most of these names should belong to the *cryptarum*complex taxa.

After B. cryptarum was established as separate species, RASMONT et al. (1986) undertook a short survey of how different Asiatic taxa might be connected to *B. cryptarum*. They suggested that some of Krügers Asiatic taxa (borochorensis, turkestanicus, burjaeticus, laevis) resembled B. cryptarum. And there are other specimens with morphological characters which make it probable that they do not belong to B. lucorum. Good examples are B. jacobsoni SKORIKOV and B. lucorum terrestricoloratus KRÜGER; the type materials of these taxa look very much like those of *B. cryptarum*: the collare is a bit greyish instead of yellow with a clear S-band of dark hair at the border of the pronotallobus, separating the collare, and a patch of grey hair at the episternum; these specimens are typical B. cryptarum. All of the museum specimens suspected of belonging to *B. cryptarum* when sequenced, were proven to be distinctly different from B. lucorum. When only the variable triplets were used in maximum likelihood simulation they all clustered within the cryptarum-complex taxa and they shared most of the diagnostic positions of the cryptarum-complex (Fig. 3). The museum specimens from within the known distribution of B. cryptarum (M-01 to M-05) shared the diagnostic positions of B. cryptarum and the specimens from the Central Asiatic Mountains and the Himalayas shared the diagnostic positions characteristic for the Transbaikal region taxa.

A characteristic gap in the distribution of bumblebees was found in the Siberian lowlands, separating two distinct areas of distribution, one in west of the Urals and a second in the Transbaikal region and the Far East (GORODKOV 1984). *Bombus cryptarum* has been found far into the northern regions of Scandinavia, Finland and European Russia (Kola Peninsula), but data are lacking east of the Urals. A distribution of *B. cryptarum* further south in the large belt of steppe vegetation north of the Asiatic Mountains is very improbable. The sequences of the museum specimens M-01 to M-08 formed a bridge from the Elburz Mountains/Iran along the Central Asiatic Mountains Alai/Tadschikistan, Kirghizian Alatau/Kyrgyzstan, to Dzungarian Alatau/Kazakhstan, an area described from the viewpoint of the zoogeography of bumblebees by SKORIKOV (1931), where the ecological conditions may be suitable for *cryptarum*-complex taxa. Additional research in the Russian and Mongolian Altai and in the Sajan Mountains may close the remaining gap in the future. Then will come the time to discuss the European–Asiatic–North American *cryptarum*-complex taxa as most probably one species, a circumpolar *B. cryptarum*.

Museum specimens as a valuable resource for DNA

In the last century it was customary to name new forms according to localities (B. alaiensis, B. borochorensis, B. turkestanicus) or in honour of people (B. jacobsoni, B. reinigi). In a group where differentiation using morphological characters is as delicate as with the many taxa resembling and somehow related to B. lucorum, this approach leads to unsatisfactory results. According to WILLIAMS (1991:84), at least 187 names have been published concerning B. lucorum 'in the broadest sense'. Much more fieldwork is necessary but fieldwork alone will not help to clarify the situation. Collecting fresh material in the Transbaikal region and naming these specimens B. burajeticus because of resemblances in colouration with the type material of Krüger is not sufficient because the characters available are not conclusive. More detailed information is required and it is necessary to connect designated museum specimens with fresh material collected in the field. DNA and especially COI sequences are obvious tools which are suited to solve at least some of these problems. By carefully adjusting the amplification conditions and manually inspecting the ABI traces at the positions with miscoding lesions, it is possible to get reliable sequences from museum specimens, thus, connecting 80-year-old museum specimens and fresh material is possible. However, the standard sequence length (648 bp) used for DNA bar-coding identification by COI might be a bit too short for such purposes, and we think that in the case of museum specimens, with possibly many degraded positions, it would be safer to use sequences of about double that bar-coding length (about 1200 bp) and to carefully compare the diagnostic positions calibrated against many sequences from fresh specimens from a broad spectrum of geographical provenances.

Conclusions

From the taxa investigated in this report, it should now be possible to use their diagnostic characters to relate unambiguously specimens to taxa, and the next step must be to find morphological characters useful for safe identification. Up until now, this task has given unsatisfactory results because the separation of the variability of characters and the delimitation of characters useful for diagnostic work has been impossible without proper designation of the specimens. The number of misidentifications in museum collections is substantial. Consequently, arguments over identification have quite often been circular and a critical re-evaluation of all the work invested into the morphological separation of taxa somehow related to B. lucorum must be done. Just the invention of a B. lucorum 'in the broadest sense' (WILLIAMS 2010) illustrates the problem, and reminds of times not so far back, when, because of difficulties of morphological separation, B. terrestris and B. lucorum from the European continent (both with white tails) were only tentatively separated as species (Hoffer 1883; Schmiedeknecht, 1930; Krüger 1920) or not at all (Faester & FAESTER 1970; WARNKE 1981). But fortunately nobody invented a B. terrestris 'in the broadest sense'. The premature mixing of clearly separable taxa (whatever species definition is used) prohibits scientific progress and insights into biological diversity. Definable unities in nature are the objects of interest; they should be properly named to promote further investigations into their ecology and distribution. Even in museum boxes it is preferable to keep taxa separate. Divide et Impera might also be good strategy to follow in bumblebee taxonomy.

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