Barcoding of Central European Cryptops centipedes reveals large interspecific distances with ghost lineages and new species records from Germany and Austria (Chilopoda, Scolopendromorpha)

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Abstract

In order to evaluate the diversity of Central European Myriapoda species in the course of the German Barcode of Life project, 61 cytochrome c oxidase I sequences of the genus Cryptops Leach, 1815, a centipede genus of the order Scolopendromorpha, were successfully sequenced and analyzed. One sequence of Scolopendra cingulata Latreille, 1829 and one of Theatops erythrocephalus Koch, 1847 were utilized as outgroups. Instead of the expected three species (C. parisii Brolemann, 1920; C. anomalans Newport, 1844; C. hortensis (Donovan, 1810)), analyzed samples included eight to ten species. Of the eight clearly distinguishable morphospecies of Cryptops, five (C. parisii; C. croaticus Verhoeff, 1931; C. anomalans; C. umbricus Verhoeff, 1931; C. hortensis) could be tentatively determined to species level, while a further three remain undetermined (one each from Germany, Austria and Croatia, and Slovenia). Cryptops croaticus is recorded for the first time from Austria. A single specimen (previously suspected as being C. anomalans), was redetermined as C. umbricus Verhoeff, 1931, a first record for Germany. All analyzed Cryptops species are monophyletic and show large genetic distances from one another (p-distances of 13.7–22.2%). Clear barcoding gaps are present in lineages represented by >10 specimens, highlighting the usefulness of the
barcoding method for evaluating species diversity in centipedes. German specimens formally assigned to *C. parisi* are divided into three clades differing by 8.4–11.3% from one another; their intra-lineage genetic distance is much lower at 0–1.1%. The three clades are geographically separate, indicating that they might represent distinct species. Aside from *C. parisi*, intraspecific distances of *Cryptops* spp. in Central Europe are low (<3.3%).

**Keywords**
Barcode, biodiversity, COI, cryptic diversity, introduced species

**Introduction**

The German Barcode of Life project – Myriapoda was started in 2012 with the aim to construct a library of reference sequences from the 200 indigenous Diplopoda and Chilopoda species of Germany (Voigtländer et al. 2011). This project, spearheaded by a study of Bavarian myriapods (Spelda et al. 2011), is still in progress. First results of the “German Myriapod Barcoding Group” were presented by Wesener et al. (2015). With the help of a comprehensive gene database, the taxonomical problems and confusion that exists in many myriapod groups on a species and higher level could be solved in combination with morphological character analyses. Additionally, barcoding could make it possible to determine juvenile and female myriapods; such a determination is often impossible with morphological characters only. Furthermore, in combination with other genetic markers, barcoding might allow analyses of the evolutionary history of species or species groups (e.g. Pilz et al. 2007, Oeyen et al. 2014).

Such a problem of taxonomic confusion applies in particular to the family Cryptopidae of the centipede order Scolopendromorpha. The Cryptopidae show an almost worldwide distribution, as they are present on most continents and many islands (Attens 1930). The family shows their highest diversity in the temperate parts of North and South America, Europe and the Mediterranean region, central and southern Africa, Madagascar, and Australia (Bonato and Zapparoli 2011). Many cryptopid taxa are currently difficult to determine and are in need of revisions. While the phylogeny of the family inside the Scolopendromorpha is still not fully resolved (e.g. Murienne et al. 2010; Vahtera et al. 2013), the monophyly of the diverse and cosmopolitan genus *Cryptops* is currently undisputed (Vahtera et al. 2012).

In Germany and most of Central Europe, the only Scolopendromorpha that occur naturally are two widely distributed species of the genus *Cryptops*: *C. parisi* and *C. hortensis* (Voigtländer et al. 2011). Both species are morphologically distinct and relatively easy to identify, at least in the adult stage. However, in the Austrian Inn-valley, unusual specimens previously assigned to *C. hortensis* have been found (Pichler 1987) which might be different from *C. hortensis*, and in later studies were placed in keys (Lewis 2011) under *C. parisi*.

A third species, *C. anomalans*, is a recent addition to the German fauna (Voigtländer 1988; Fründ 1989; Spelda 2006, Decker and Hannig 2011). Although already mentioned as a possible member of the German fauna by Schubart (1964) this species
was most likely introduced from the Mediterranean realm to northern Europe (Eason 1964; Lindner 2005), as it is mainly confined to parks and gardens. Because the species has few records in Germany (Decker et al. 2014), a special effort was undertaken to collect specimens from the limited number of known German populations.

There are only a handful of barcoding and phylogenetic studies applying molecular data of Scolopendromorpha worldwide (Murienne et al. 2010; Simaiakis et al. 2012; Vahtera et al. 2012, 2013; Joshi and Edgecombe 2013; Oeyen et al. 2014; Siriwut et al. 2015). For Cryptops, there is only a singular molecular study utilizing barcoding genes and it deals with tropical pacific island species (Murienne et al. 2011). Therefore, this study focusing on Central European/German Cryptops is the first of its kind.

Barcoding studies inside the Scolopendromorpha consecutively revealed large interspecific distances (Simaiakis et al. 2012; Joshi and Edgecombe 2013; Oeyen et al. 2014; Siriwut et al. 2015). The only study involving Cryptops (Murienne et al. 2011) revealed exceptionally high intra- and interspecific distances, similar to the observations made in other Scolopendromorpha genera (see above), as well as in a recent study on German geophilomorph centipedes (Wesener et al. 2015).

The aim of this study is to see if barcoding of Cryptops allows (a) a clear separation of the species found in Germany; (b) enables the detection of potential cryptic lineages in the widespread German species; as well as (c) facilitating the correct identification of morphologically distinct specimens from Central Europe.

**Material and methods**

**Specimen collection and preparation**

The focus of the project was Cryptops from Germany, which encompass 85% of the here analysed specimens of the genus (Fig. 1). The remaining 15% (11) successfully sequenced specimens of Cryptops were collected in adjacent countries. Our sample includes six specimens from Austria, two from Italy, and one each from Croatia, Wales, and Slovenia. One of the Italian specimens is of special importance as it came from the type locality of the subspecies Cryptops parisi sebini Verhoeff, 1934. All specimens are stored as vouchers in 95% undenatured ethanol, either at the Museum Koenig, Bonn, Germany (ZFMK), the Senckenberg Museum für Naturkunde, Görlitz (SMNG) or the Bavarian State Collection of Zoology, Munich, ZSM (see Table 1, full specimen information in Suppl. material 1).

The specimens were collected by hand and transferred to vials containing 95% undenatured ethanol within days of collection. The vials contain an individual GBOL number with which the specimens can be connected to the accompanying data. After conservation the specimens were either sent to the GBOL facility at the ZFMK or to the corresponding laboratory at the ZSM. Upon arrival, all specimens were photographed (images are or will be uploaded to BOLD, http://www.boldsystems.org/), and a tissue sample was removed for DNA extraction. For this specific GBOL subproject,
### Table 1.

GBOL numbers, GenBank codes, locality data. GBOL number refers to DNA extraction and BOLD registration; L Nr refers to number of Map (Figure 1). 

**SMNG** = Senckenberg Museum für Naturkunde, Görlitz, Germany; **ZFMK** = Zoological Research Museum A. Koenig, Bonn, Germany; **ZSM** = Zoologische Staatsammlung München, Germany.

| L Nr | GBOL   | GenBank  | Voucher               | Species       | Locality                                                                 |
|------|--------|----------|-----------------------|---------------|---------------------------------------------------------------------------|
| 1    | GBO102755 | KU497147 | ZSM-ART-JSP130822-001 | Scolopendra cingulata | Croatia, Istra, Umag                                                        |
| 2    | GBO102750 | KU497149 | ZSM-ART-JSP130424-007 | Theatops erythrocephalus | Croatia, Istra, Brestova                                                   |
| 1    | ZFMK-TIS-2531556 | KM491707 | ZFMK-MYR 3450 | Cryptops hortensis | Germany, Waren (Müritz), Nationalpark Müritz                              |
| 2    | ZFMK-TIS-2531557 | KM491678 | ZFMK-MYR 3438 | Cryptops hortensis | Germany, Waren (Müritz), Nationalpark Müritz                              |
| 2    | ZFMK-TIS-1470 | KU342047 | ZFMK-MYR 3853 | Cryptops hortensis | Germany, Potsdam, Babelsberg                                              |
| 2    | ZFMK-TIS-2507217 | KU342045 | ZFMK-MYR 3888 | Cryptops hortensis | Germany, Potsdam, Babelsberg                                              |
| 3    | ZFMK-TIS-1543 | KM491700 | ZFMK-MYR 3684 | Cryptops hortensis | Germany, Ilsenburg                                                        |
| 4    | ZFMK-TIS-1528 | KM491595 | ZFMK-MYR 3679 | Cryptops hortensis | Germany, Friedeburg (Saale)                                               |
| 4    | ZFMK-TIS-2519823 | KM491677 | ZFMK-MYR 3824 | Cryptops hortensis | Germany, Friedeburg (Saale)                                               |
| 5    | ZFMK-TIS-1289 | KU342043 | ZFMK-MYR 3551 | Cryptops hortensis | Germany, Hoyerswerda, Dubringer Moor                                       |
| 6    | ZFMK-TIS-15761 | KM491615 | ZFMK-MYR 1057 | Cryptops hortensis | Germany, Bonn - Bad Godesberg, Panoramapark                              |
| 7    | ZFMK-TIS-15555 | KU342044 | ZFMK-MYR 1043 | Cryptops hortensis | Germany, Niederzissen, Bausenpark                                        |
| 8    | GBOL14853 | KU497144 | ZSM-ART-JSP130930-017 | Cryptops hortensis | Germany, Enzberg, Kieselbronn                                              |
| 9    | GBOL02747 | KU497160 | ZSM-ART-JSP110312-009 | Cryptops hortensis | Germany, Zuckerberg SW Stuttgart-Steinhaldenfeld                         |
| 10   | GBOL10885 | KU497162 | ZSM-ART-JSP110312-009b | Cryptops hortensis | Germany, Zuckerberg SW Stuttgart-Steinhaldenfeld                         |
| 11   | GBOL14855 | KU497145 | ZSM-ART-JSP150118-018 | Cryptops hortensis | Germany, Kenzingen, Forlenwald                                             |
| 12   | ZFMK-DNA-112780039 | KM491565 | ZSM-ART-JSP100619-031 | Cryptops hortensis | Germany, Mainau island, 4 km NNE Konstanz                               |
| 13   | GBOL14858 | KU497146 | ZSM-ART-JSP150121-039 | Cryptops hortensis | Germany, Mainau island, 4 km NNE Konstanz                               |
| 14   | ZFMK-DNA-112780041 | KU342046 | ZSM-ART-JSP110208-005 | Cryptops hortensis | Italy, Provincia di Sondrio, Chiavenna, Riserva Naturale Marmite dei Giganti |
| 15   | ZFMK-TIS-19439 | KM491610 | ZFMK-MYR 1948 | Cryptops parisi | Germany, Bochum, Botanical Garden of the Ruhr-University                  |
| 16   | ZFMK-TIS-1619 | ZFMK-TIS-1619 | Cryptops parisi | Germany, Leipzig-Schöenefeld, Partheaue                                    |
| 17   | ZFMK-TIS-15786 | KM491698 | ZFMK-MYR 1082 | Cryptops parisi | Germany, Schwelm-Erlen, nahe Eingang Erlenhöhle,                           |
| L. Nr | GBOL | ZFMK | Voucher | Species | Locality |
|------|------|------|---------|---------|----------|
| 18   | ZFMK-TIS-5767 | KM491624 | ZFMK-MYR 1063 | Cryptops parisi | Germany, Wupperal, NSG Im Holken |
| 19   | ZFMK-TIS-6357 | KM491666 | ZFMK-MYR 5355 | Cryptops parisi | Germany, Weißenberg, Goldener Schla |
| L Nr | GBOL     | GenBank | Voucher               | Species            | Locality                                                                 |
|------|----------|---------|-----------------------|--------------------|---------------------------------------------------------------------------|
| 40   | GBOL14861| KU497141| ZSM-ART-JSP150201-104 | Cryptops parisi    | Germany, W slope of Lercheck, 1 km NW Unterau, 5 km NE Berchtesgaden      |
| 41   | GBOL02742| KU497140| ZSM-ART-JSP130522-015 | Cryptops parisi    | Germany, SW Grafenbachau, 8 km SW Murnau                                 |
| 42   | ZFMK-DNA-112780073| KU342053| ZSM-ART-JSP100510-004 | Cryptops parisi    | Germany, Bad Toelz, Altjoch                                                |
| 43   | ZFMK-TIS-2517130| KU342055| ZFMK-MYR 2470       | Cryptops parisi    | Italy, Lombardia, Brescia, Pisogne, Type locality                       |
| 44   | GBOL12332| KU497142| ZSM-ART-JSP141214-001| Cryptops parisi    | UK, Wales, Aberbargoed                                                    |
| 45   | ZFMK-TIS-1587| KM491706| ZFMK-MYR 4072       | Cryptops anomalans  | Germany, Leipzig, Pleißemühlgraben                                       |
| 46   | ZFMK-TIS-18969| KM491703| ZFMK-MYR 1379       | Cryptops anomalans  | Germany, Bonn, Friedsfeld                                                |
| 47   | ZFMK-TIS-15751| KM491699| ZFMK-MYR 1047       | Cryptops anomalans  | Germany, Bonn - Bad Godesberg, Panoramapark                              |
| 48   | ZFMK-TIS-15752| KM491639| ZFMK-MYR 1048       | Cryptops anomalans  | Germany, Bonn - Bad Godesberg, Panoramapark                              |
| 49   | BCZSMMYR00489| JN266286| ZSM-ART-JSP100619-017| Cryptops umbricus   | Germany, Langenaltheimer Haardt 1 km W Solnhofen, 4 km S Pappenheim     |
| 50   | GBOL02745| KU497151| ZSM-ART-JSP130812-004| Cryptops anomalans  | Germany, Hummelgraben, Stuttgart-Zuffenhausen                            |
| 51   | GBOL14852| KU497158| ZSM-ART-JSP110624-001| Cryptops anomalans  | Germany, SW Stuttgart-Muehlhausen                                         |
| 52   | ZFMK-TIS-1434| KU342048| ZFMK-MYR 1236       | Cryptops croaticus   | Austria, Leithagebirge, Zeiler Berg                                      |
| 53   | ZFMK-TIS-9755| KM491620| ZFMK-MYR-1185      | Cryptops croaticus   | Austria, Leithagebirge I                                                 |
| 54   | ZFMK-TIS-14960| KU497153| ZSM-ART-JSP110425-008| Cryptops sp.         | Croatia, NW Baci and Brestova, 10 km NE Labin                             |
| 55   | ZFMK-TIS-14134| KU342042| ZFMK-MYR 3662       | Cryptops sp.         | Germany, Saxony, Leipzig, Zoo, Gondwanaland                               |
| 56   | GBOL14857| KU497143| ZSM-ART-JSP150118-047| Cryptops sp.         | Slovenia, Osocja 2 km NW Zagon, 5 km NW Postojna                           |
Figure 1. Distribution map of all successfully sequenced Central European specimens of *Cryptops*. Numbers refer to each specimen (see Table 1). Symbols and colours denote species. Blue rectangle = *C. parisi*; red circle = *C. anomalans*; green triangle = *C. hortensis*; brown diamond = *C. croaticus*; orange cross = *C. umbricus*; light blue, orange, and yellow symbols mark undetermined *Cryptops* species.
DNA extraction was attempted for 77 specimens of Cryptops as well as one each of Scolopendra cingulata and Theatops erythrocephalus as outgroups (See Table 1).

Maps were created with ArcGIS 10.

**DNA extraction and sequencing**

At the ZFMK, DNA was extracted from the tissue samples using the BioSprint96 magnetic bead extractor by Qiagen (Germany). After the extraction, samples were outsourced for PCR and sequencing (BGI China). For PCR and sequencing, the degenerated primer pair HCOJJ/LCOJJ (Astrin and Stüben 2008) was used, resulting in a success rate of >75% (38 of 49 extracted specimens).

At the ZSM, a single leg was removed from each specimen and sent in 96 well lysis plates to the Canadian Centre for DNA Barcoding (CCDB, Guelph, Canada) for standardized, high-throughput DNA extraction, PCR amplification and bidirectional Sanger sequencing (http://www.ccdb.ca/resources.php). For PCR and sequencing, a primer cocktail (Hebert et al. 2004, see Table 2) was used, resulting in a success rate of >90% (23 from 25 extracted specimens). All voucher information and the DNA barcode sequences, primer pairs and trace files were uploaded to BOLD (http://www.boldsystems.org).

Sequences were obtained for 61 Cryptops as well as the two outgroup specimens. The three available sequences of Central European Cryptops were added from a previously published dataset (Spelda et al. 2011). Sequence identities were confirmed with BLAST searches (Altschul et al. 1997). All 63 new sequences were deposited in GenBank (see Table 1 for accession numbers). In order to rule-out the accidental amplification of nuclear copies of the mitochondrial COI gene, the whole dataset was translated into amino acids (see Supplemental Material) following the ‘invertebrate’ code in MEGA6 (Tamura et al. 2013); internal stop codons were absent in our dataset. There were a total of 657 positions in the final dataset, gaps were absent.

**Table 2.** List of primers used for amplification and sequencing of the 5’ part of the mitochondrial COI gene.

| Primer name | Sequence | Publication | Used at |
|-------------|----------|-------------|---------|
| LCO1490     | 5’-GGTCAACAAATCATAAAGATATTGG | Folmer et al. 1994 | CCDB for ZSM |
| HCO2198     | 5’-TAAACTTCAGGGGTGACCACAAAAATACTA | Folmer et al. 1994 | CCDB for ZSM |
| LepF1       | 5’-ATTCAACCAATCATAAAGATATTGG | Hebert et. al. 2004 | CCDB for ZSM |
| LepR1       | 5’-TAAACTTCTGGATGTCACAAAAATCA | Hebert et. al. 2004 | CCDB for ZSM |
| C_LepFolF   | cocktail of LepF1 and LCO1490 | www.boldsystems.org/index.php/Public_Primer_PrimerSearch | CCDB for ZSM |
| C_LepFolR   | cocktail of LepR1 and HCO2198 | www.boldsystems.org/index.php/Public_Primer_PrimerSearch | CCDB for ZSM |
| LCO1490-JJ  | 5’-CHACWAAYCATAAAGATATYGG | Astrin and Stüben 2008 | ZFMK |
| HCO2198-JJ  | 5’-AWACTTCTGGRTGVCACAAARAATCA | Astrin and Stüben 2008 | ZFMK |
Phylogenetic analysis

Sequences were aligned by hand in Bioedit (Hall 1999). The final dataset included 66 nucleotide sequences with 657 positions (63 newly sequenced). Phylogenetic analyses were conducted in MEGA6 (Tamura et al. 2013). A Modeltest, as implemented in MEGA6 (Tamura et al. 2013), was performed to find the best fitting maximum likelihood substitution model. Models with the lowest BIC scores (Bayesian Information Criterion) are considered to describe the best substitution pattern. Included codon positions were 1st+2nd+3rd+Noncoding. Modeltest selected the General Time Reversible model (Nei and Kumar 2000) with gamma distribution and invariant sites as best fitting model (lnL -4725.286624, Invariant 0.505, Gamma 1.65919, R 3.11, Freq A: 0.2844, T: 0.3433, C: 0.2113, G: 0.1606). The tree with the highest log likelihood (-4725.2866) is used here to infer the genetic distances and evolutionary history of the analyzed specimens. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories (+G, parameter = 1.6591)). The rate variation model allowed for some sites to be evolutionarily invariable ((+I), 50.5% sites). The bootstrap consensus tree inferred from 1000 replicates (Felsenstein 1985) is taken to represent the evolutionary history of the analyzed taxa. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site.

Distance analysis

The number of base differences per site between sequences is shown in figures and tables (Fig. 3; Suppl. material 2). The analysis involved 66 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All ambiguous positions were removed for each sequence pair. There were a total of 657 positions in the final dataset. Evolutionary distance analyses were conducted in MEGA6 (Tamura et al. 2013). Two frequency distribution diagrams of all pair-wise intra- and inter-specific distances were produced to further evaluate species divergence in Cryptops. All samples of each species were grouped in the first analysis, while Cryptops parisi was split into the three separate clades C. parisi sensu stricto, C. parisi sebini and C. parisi lineage3 in the second analysis.

Results

Phylogenetic analysis

The monophyly of the genus Cryptops is strongly supported (97%) in our tree (Fig. 2). One undetermined Cryptops sp. collected from the tropical rainforest
Figure 2. Maximum likelihood tree under the GTR+G+I model, 1000 bootstrap replicates. Colours and symbols correspond to Maps (Figs 1, 4). Country of origin given after specimen number: AT = Austria; DE = Germany; GB = Wales; HR = Croatia; IT = Italy; SL = Slovenia. Photograph shows a specimen of Cryptops parisi s.s. from Breckerfeld (photo A. Steiner), western Germany. For full data on all specimens, see Table 1.
Figure 3. Frequency distribution of pairwise intraspecific (blue) and interspecific (red) distances. All lineages of *C. Parisi* treated as one species, *C. Parisi* sensu lato. Basic table see Suppl. material 1.
greenhouse at the Leipzig Zoo in eastern Germany (Fig. 1: 55) is in a basal position juxtaposed to all other Cryptops specimens (Fig. 2). The remaining Central European Cryptops are split into two clades, of which only the C. parisi/C. croaticus clade receives high statistical support (96%). The unsupported clade unites C. umbricus and C. anomalans, three specimens of uncertain identity, and C. hortensis (Fig. 2). C. anomalans is in a basal position regarding this second (unsupported) clade with a single haplotype spread all over Germany, forming a monophylum with C. umbricus from Solnhofen, Germany, representing the first record from this country (Fig. 1: 46). The uncertain Cryptops sp. from Slovenia is a sister group to a weakly supported clade (76% bootstrap support) uniting two unidentified Cryptops sp. specimens with C. hortensis (Fig. 2). The latter two unidentified Cryptops sp. specimens from eastern Austria and Croatia are grouped together, but this grouping is not statistically supported.

The monophyly of the 18 specimens of C. hortensis is strongly supported (100%). Of the shallow clades inside C. hortensis (Fig. 2), only one, a clade uniting five different haplotypes from Italy, eastern and western Germany (Fig. 4), receives some statistical support (78%). Interestingly, a second specimen from Friedeburg, Saxony-Anhalt, the same locality as one of the five haplotypes mentioned above (see Table 1), groups within a separate clade (Fig. 2).

The clade uniting C. parisi sensu lato and C. croaticus receives high statistical support (96%). While both specimens of C. croaticus show the same haplotype, the 32 specimens of C. parisi s. l. are separated into three statistically well-supported (99–100%) clades. The basalmost clade (Fig. 2) includes seven specimens and represents three different haplotypes from the eastern alpine region (Fig. 4: green). The remaining two clades of C. parisi are clearly related (92% support); one represents a western clade (Fig. 4: yellow) and the other is found slightly more to the east (Fig. 4: blue) and also includes the toptotypoid of the subspecies C. parisi sebini.

**Distance analysis**

Cryptops specimens differ from the outgroups Scolopendra and Theatops by 19.8–25.7% (Supplementary Material 2). Interspecific and intraspecific distances of the different nominal Cryptops species show no overlap (Fig. 3). Interspecific distances lie between 13.4–21.1% (Fig. 3), with the lowest observed between C. croaticus and C. parisi s. l. (13.4–14.8%) as well as between C. anomalans and C. umbricus (13.9%). Otherwise, interspecific distances are always >16%, with the highest value of >20% observed between C. anomalans and C. hortensis, as well as between C. parisi sebini and C. hortensis. Intraspecific distances are between 0–11.3%. However, intraspecific distances are low, 0–3.3%, if we treat the three distinct lineages of C. parisi as distinct species (Fig. 5).
Discussion

Distance analysis

Clear intraspecific distances in German or even Central European *Cryptops* are low. The specimens filling the majority of our barcoding gap between 3 and 11.3% are the different lineages of *C. parisi*, which differ by 8.4–11.3% from one another (Fig. 3) and might represent distinguishable taxonomic units (see below). Two specimens directly at the edge between inter- and intraspecific distances (Fig. 5), the two *Cryptops “sp. 2”* specimens from Austria and Croatia (15.9%), require a careful re-study (see below).

The biogeographic and ecological pattern of *C. hortensis* and *C. parisi* in Central Europe

*Cryptops parisi* and *C. hortensis* belong to the South European and Central Asiatic European chorotypes respectively (Zapparoli 2006). In central Europe *C. hortensis* and *C. parisi* s. l. seem to exclude each other either geographically or ecologically. In the lowland areas of north-western Germany and in the Upper Rhine valley it is usually *C. hortensis* that occurs, while in the lower mountain ranges usually *C. parisi* is present. Nevertheless, *C. parisi* mainly avoids higher altitudes. In the eastern part of Germany *C. parisi* dominates.

*Cryptops parisi* is generally classified as a mesophilous woodland species (Spelda 1999, Minelli and Iovane 1987, Voigtländer et al. 1997), but may also occur outside of forests, especially in northern Germany where more anthropogenic influenced places are inhabited.

The two clearly differentiated genetical lineages in *C. parisi* s. s. in Germany (see below) are reflected in distinct ecological differences in the preferred habitats between the western and eastern parts of Germany. In the more Atlantic areas in the West, the species prefers woodland like in its main distribution area. In the more continental influenced East, *C. parisi* inhabits open-dry habitats such as dry meadows, mesoxeric meadows and their successional shrub-stages, as well as dwarf-shrub heaths (Voigtländer 2003a, 2003b, 2005).

A single haplotype in German *Cryptops anomalans*

*C. anomalans* is viewed as a species introduced to Germany and England (Eason 1964; Voigtländer 1988). Specimen records are rare, e.g. the species has only recently been recorded from Germany, where it only occurs in localized areas, usually in parks or gardens (Lindner 2010, Decker and Hannig 2011). Our findings show that a single haplotype (Fig. 2) is present in western, eastern and southern Germany (Fig. 1), while all other *Cryptops* (see Fig. 3), as well as Geophilomorpha species (Wesener et al. 2015)
show different haplotypes across a large geographical area. An identical haplotype from different localities might be interpreted as recent human introductions from a homogeneous source population or a rapid spread of *C. anomalans* in Germany.

**First record of *C. umbricus* in Germany**

Our analyses first showed one outlier *C. anomalans* specimen from Solnhofen, Bavaria (Fig. 1), which strongly differs by 13.9% from the common German haplotype. This was the only specimen of *C. anomalans* in a previous analysis involving German centipedes (Spelda et al. 2011). A morphological check against similar species showed that it was indeed not *C. anomalans* but represents *C. umbricus*, a first record for Germany. This finding shows the usefulness of the barcoding method in detecting previously unrecorded species.

**At least three undetermined *Cryptops* species in Central Europe**

*Cryptops* sp. 1 is only represented in our dataset by a single specimen from Slovenia, which is unfortunately missing the pre-ultimate legs and can therefore not easily be determined morphologically.

*Cryptops* sp. 2 is represented by two specimens that are separated by a wide genetic distance of 15.9%. This distance usually falls right into the lower limit observed between different *Cryptops* species (Fig. 3). The two specimens are from the eastern lowlands of Austria (Burgenland) and Croatia (Brestova). Unfortunately, the Austrian specimen is heavily damaged with missing posterior segments, which prevents any determination. As both specimens of *Cryptops* sp. 2 are related, but potentially not conspecific, they are discussed here together.

These two specimens are similar to *C. hortensis*, but are missing the ventral furrow on the prefemora of the ultimate leg pair. An available name for one of these lines might be *C. rucneri* Matic, 1966. This species was synonymised with *C. hortensis* by Koren (1986), followed by Spelda (1999), but treated as a valid species later (Stoev 2002). The presently discovered genetic diversity brings this name into consideration again. One argument for the identity of one of our lines with *C. rucneri* is the configuration of the prefemur of the ultimate leg pair, where Matic (1966) did not mention a ventral furrow. Although Matic (1966, 1972) did not describe and depict the poison gland in great detail, his figures clearly show that in both *C. hortensis* sensu Matic (1972a) and *C. rucneri*, the calyx of the poison glands lie mainly in the femur and tibia of the forcipule. Matic also records *C. rucneri* from Italy (Matic 1967), Austria: Carinthia (Matic 1972b), and Slovenia (Matic 1979).

Maybe this specimen is the same species to which Pichler (1987) refers to as *Cryptops* cf. *hortensis* from North Tyrol. The shape of the poison gland was not illustrated for *C. cf. hortensis*. The poison gland allows a clear separation from *C. parisii* even in
Figure 4. Distribution map of all successfully sequenced Central European specimens of *Cryptops parisi*. Different colours mark the three different clades. Yellow = *C. parisi* sensu stricto; blue = *C. parisi sebini*; green = *C. parisi* lineage 3 (potentially *C. cf. hortensis* sensu Pichler 1987).
**Figure 5.** Frequency distribution of pairwise intraspecific (blue) and interspecific (red) distances. The three lineages of *C. parisi* treated as different species. Basic table see Suppl. material 1.
very early stages. Without checking the poison gland, juvenile specimens of *C. parisi*,
which lack the characteristics of adult specimens (a central depression on the forcipular
tergite and the pair of occipital sutures), can be easily mistaken for *C. hortensis*. Pichler
(1987) records an unidentate labrum for *C. cf. hortensis*, as does Matic (1966) for *C.
rucneri*. Pichler’s (1987) fig. 18 of the 21st pleurocoxa corresponds to fig. 4 of Matic
(1966) for *C. rucneri*.

Of the two specimens of *Cryptops* sp. 2, the one from Brestova is the most probable
to represent *C. rucneri*. This specimen was collected only 30 kilometres distant from
the type locality of *C. rucneri* and shows the characteristic elongated 20th leg pair,
which is unfortunately missing in the other specimen (as well as in our *Cryptops* sp. 1).
Nevertheless, while having only three sequences of these eastern *C. hortensis*-relatives
and without being able to provide a revision of the *hortensis/rucneri*-complex we prefer
at the moment to keep these specimens under the name *Cryptops* sp.

*Cryptops* sp. 3, previously determined as *C. cf. doriae* Pocock, 1891 is only known
from the Leipzig Zoo in eastern Germany, where it was collected in a large tropical
greenhouse (Decker et al. 2014). It was provisionally identified as *C. doriae*, a member of
the *doriae*-group, which is characterized by having teeth on femur, tibia and tarsus
of the ultimate legs (Lewis 2011). *C. doriae* was already reported from a tropical biome
in England (Lewis 2007) and is so far the only introduced tropical *Cryptops* species
with records in Europe (Stoev et al. 2010). A BLAST search of our specimen against
the sequences of *C. doriae* already deposited on GenBank (11.2015) reveals a large ge-
netic distance between our specimen and the ones from the Pacific, which is the reason
we refer to our specimen as *Cryptops* sp. 3.

**First record of *Cryptops croaticus* in Austria**

*Cryptops croaticus* was originally described from Bakar (formerly Buccari) in Croatia
(Verhoeff 1931) and subsequently recorded from other localities in Croatia, Slovenia
and Bosnia-Herzegovina (Matic 1966, 1979, Kos 1992), Greece (Matic 1976), Bul-
garia (Stoev 1997a, 2002), and Italy (Matic 1960, 1968, Matic and Darabantu 1971,
Minelli 1985, 1992). Currently, *C. croaticus* seems to be absent or not yet found in
Hungary (Dányi 2008). One subspecies (*C. croaticus burzenlandicus*) was described
from Romania (Verhoeff 1931) and was subsequently synonymised with the nominal
subspecies (Matic 1972a), another subspecies, *C. croaticus albanicus*, has been described
from Albania (Verhoeff 1934) and was later synonymized under *C. anomalans* (Stoev
1997b). Several subspecies have been described from Italy, namely *C. croaticus ber-
gomatus* (Verhoeff 1934), *C. croaticus longobardius* and *C. croaticus baldensis* (Manfredi
1948), subsequently cited by Conci (1951) and Boldori (1969). Based on this wide
distribution, the occurrence of *C. croaticus* in Austria is not unexpected. In Austria, it is
currently only known from a southern exposed slope, which is home to numerous relic
species adapted to a warmer climate. *C. croaticus* shares its habitat with the recently
rediscovered population of *Scolopendra cingulata* in Austria (Oeyen et al. 2014), as well
as the thermophilic beetle *Carabus hungaricus* and other thermophilic animals (Böhme et al. 2014). However, the determination of our specimens as *C. croaticus* is only based on the characters given in the original description (Verhoeff 1931) as no better description exists. Numerous important characters, such as the last leg pairs, are unfortunately missing in our specimens. A revision of *C. croaticus* is urgently needed (Matic 1966) as it may be that some of the nominal subspecies represent independent species. One way to clarify this is to collect and sequence topotypic material. Once *C. croaticus* has been properly revised, a re-evaluation of the Austrian specimens should be undertaken.

**The three lineages of *Cryptops parisi* sensu lato**

The three lineages of specimens placed in *C. parisi* by morphological characters differ 8.4–11.3% from one another, while their intra-lineage genetic distance is much lower at 0–1.1%. A large barcoding gap becomes clearly visible in our dataset when we treat the three different lineages of *C. parisi* as separate species (Figs 4, 5). Endosymbionts like *Wolbachia* (Hurst and Jiggins 2005) are an unlikely explanation for the different lineages, as such endosymbionts have never been recorded in the Myriapoda (Witzel et al. 2003).

One lineage clearly represents the *C. parisi* sensu stricto (Fig. 2: yellow). This group shows a western distribution in Germany, with a single specimen from southern Germany (Fig. 4). The type locality of *C. parisi* is, as the species epithet implies, Paris, France. Our only sample from Great Britain (Wales) also falls into this group. Intra-lineage variation is low with 0–1.7%. Inner structure of the lineage is limited due to the small genetic distances inside the group, but one group containing only few haplotypes differing in a single or two basepairs from one another is well-supported. This group contains specimens from western Germany, as well as a single specimen each from southwestern (ZFMK-TIS 2520349) and southeastern Germany (ZFMK-DNA-112780049), but these two were collected in a park and a garden.

A second distinct group (Fig. 2: Blue) contains the topotypic specimen of the subspecies *C. parisi sebini* Verhoeff, 1934. *C. parisi sebini* was recently synonymised under *C. parisi* because no morphological differences could be detected (Lewis 2011). However, the distinctiveness of the subspecies *C. parisi sebini* should be re-evaluated, as our genetic data supports this monophyletic subspecies (100% bootstrap support) with a high genetic distance to *C. parisi* s. s. (8.4–9.4%) in combination with low intra-lineage variation (0–0.6%) despite the large geographical distances between the analyzed specimens from Italy and eastern Germany. This *C. parisi* group 2 shows a distribution to the east of *C. parisi* s. s., with localities in eastern northern Italy and the eastern half as well as the south of Germany (Fig. 4). Another name potentially available for this clade is *C. parisi rhenanus* Verhoeff, 1931, which is characterized by its extremely elongated calyx of the poison gland (Verhoeff 1931). If both names turn out to represent the same species, this taxon would have priority over *C. parisi sebini*, with
which it is compared in the original description (Verhoeff 1934). Unfortunately, Verhoeff (1931) never designated a type for *C. parisi* rhenanus. The specimens represented in the Bavarian State Collection of Zoology originate from a large number of localities.

The specimens of *C. parisi* s. l. belonging to a third group (Fig. 2: green), referred here as *C. parisi* lineage 3, are morphologically and genetically distinct and may also be identical to the specimens of *C. cf. hortensis* in the literature (Pichler 1987, Lewis 2011). Our specimens of *C. parisi* lineage 3 come mainly from alpine habitats in Austria and Germany. In the most recent revision of the species group (Lewis 2011), these specimens were listed in the key under *C. parisi*, but with remarks concerning its unique morphology. Coxal pores are too numerous (~50) for *C. hortensis* and more closely resemble the lower end of *C. parisi*. Other morphological characters prompted Lewis (2011) to place these specimens in his key under *C. parisi*, an affinity confirmed here by our genetic analysis.

However, the large genetic distance of 10–11.3% between *C. parisi* lineage 3 and *C. parisi* s. s. as well as to the lineage containing *C. parisi* sebini, combined with a low intraspecific distance (0–1.1%) are clear indications that these specimens might represent a species of its own.

To find names for our two eastern lines of *C. parisi* one has to go back to C. L. Koch, who described three Cryptops species from around Regensburg, Germany: *C. ochraceus* C. L. Koch, 1844 from the Keilstein (a calcareous mountain east of Regensburg), *C. sylvaticus* C. L. Koch, 1844 from the Naab-valley (north of Regensburg) and *C. pallens* C. L. Koch, 1847 from the moat of Regensburg. More information on these species, such as the precise type localities and more detailed descriptions, are provided in Koch (1863), which has often resulted in these species erroneously being assigned to the date of this second publication.

Attems (1930) indicated that it would be impossible to assign these species to either *C. hortensis* or *C. parisi*, while Matic (1972a) simply synonymized them with *C. hortensis*. Both did not take note of the central depression, often darker than the adjacent parts of the tergite, as a character separating *C. parisi* from *C. hortensis*, at least for adult specimens from southern Germany (own observation, JS). This depression is also described by Attems (1930) as existing in some *C. parisi* specimens, but is not otherwise mentioned in the available keys separating the two species (Attems 1930, Brölemann 1930, Verhoeff 1931, Eason 1964, Matic 1972a, Koren 1982, 1986, Iorio and Geoffroy 2008). Verhoeff (1934) also described this character in *C. parisi* sebini. Koch (1863) clearly states and depicts the depression for his species *C. sylvaticus* and *C. ochraceus*. It seems only to be missing in *C. pallens*, which represents a juvenile specimen. Another argument against a synonymy of these species with *C. hortensis* is the absence of the latter species in our extensive collections from eastern Bavaria. Topotypoids of *C. ochraceus* have already been collected and might clarify this species in the near future.

It should be noted that Matic (1972a) depicts a *C. parisi* with a short poison gland. This specimen surely represents a different species.
Outlook/future studies

Future prospects should include the parallel sequencing of nuclear genes to confirm the relationships drawn from the mitochondrial barcoding fragment. To clarify the taxonomic relationships within *Cryptops parisi*, it would be important to collect further samples to enable an extensive morphological evaluation.

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Supplementary material 1

Full specimen information
Authors: Thomas Wesener, Karin Voigtländer, Peter Decker, Jan Philip Oeyen, Jörg Spelda
Data type: occurrence
Explanation note: Full specimen information.
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Supplementary material 2

Uncorrected P-Distance
Authors: Thomas Wesener, Karin Voigtländer, Peter Decker, Jan Philip Oeyen, Jörg Spelda
Data type: measurements
Explanation note: Uncorrected P-Distance between the 66 analyzed specimens.
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