Description of Some Aquatic Insect Genera in Greater Zab River Branches, North of Iraq

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Aquatic insects samples were collected from 6 sites along the Greater Zab River in the northern Iraq from Duhok and Erbil governorates over 12 months during September 2016 to August 2017, which belong to seven orders (Ephemeroptera, Plecoptera, Trichoptera, Odonatan, Diptera, Coleoptera, and Megaloptera). Clustering mitochondrial DNA cytochrome c oxidase and 16S rDNA genes, morphological keys, and matches in the Barcode of Life Database, we identified 24 species return to 7 orders and 12 families, as indicated in the results. The reported species were: Ephemeroptera 5 members of the family Heptageniidae (Maccaffertium meririvulanum, Raptoheptagenia cruentata, Ecdyonurus dispar, Anepeorus rusticus, Stenonema femoratum), 1 Ephemereillidae (Seratella ignita), 1 Arthropleidae (Arthroplea bipunctata), 6 Baetidae (Baetis alpinus, Baetis braaschi, Baetis noa, Baetis harrisoni, Iswaeon anoka, Heterocloeon amplum), 1 member for each of Diptera, Coleoptera, Megaloptera and Odonatan orders, while Plecoptera 2 members Leuctridae (Leuctra hippopoides, Leuctra inermis) and Tricoptera 4 members 3 Hydropsychidae (Leptonema albovirens, Hydropsyche simulans, Arctopsyche irrorate), 1 Hydroptilidae (Ochrotrichia tenuata). Most of these recorded species and genera were mentioned for the first time and represent new records in Iraq. Presence and distribution of identified species varied between studied sites, as a result of differences in biogeographical and physical conditions.

Keywords: Aquatic insects, Morphology, Mitochondrial DNA cytochrome c oxidase, 16S rDNA gene

1. Introduction
One of the most common approaches for assessing the effects of stresses on the ecological conditions of aquatic ecosystems and water quality monitoring is to studying benthic macroinvertebrate populations [1]. Aquatic macroinvertebrates are critical to ecosystem functioning through regulation of many essential top down and bottom-up ecosystem processes such as energy translocation, nutrient flow, and detrital decomposition, and act as excellent bioindicators of stream health because of their sensitivity to pollution [2] and wide variation of response to pollutants [3-5]. The disappearance or loss of biodiversity in macroinvertebrate communities could be easily assigned to anthropogenic pressure [6] integrating the impacts of multiple chemical and physical stressors, including land-use types [7].

Mayflies, stoneflies and caddisflies (Ephemeroptera, Plecoptera and Trichoptera) are prominent representatives of aquatic macroinvertebrates widely used as indicator species for water quality and environmental assessment. Though, clear morphological identification of EPT species, particularly their immature life stages, is difficult but essential task [8].

Nowadays, DNA barcoding and DNA sequencing in certain gene loci are used as an identification method to provide a valuable way of understanding species diversity in many populations of taxonomically complicated studies and confirming the taxonomy of poorly recognized communities [9, 10].

Within northern Iraq (Iraqi Kurdistan Region), the water resource from Greater Zab River catchment is under population pressure and used for drinking, domestic, agricultural and industrial supplies [11,12]. This study aimed to use the findings from an ecological perspective as bio-indicators for the Greater Zab Tributary.

2. Materials and Method
2.1 Study area:
In the present study, a total of 6 sites for sampling and collecting of aquatic insects were selected along the Greater Zab River within Erbil and Duhok Province as follows (Figure 1).
2.2 Aquatic Insect sampling:
Samples of aquatic insects from sediments, rocks and fallen leaves were collected by using Surber sampler in each sampling sites monthly for 12 months from six different studied sites from September 2016 to August 2017. The samples were washed by river water using sieves with (500 μm mesh). After being transported to the laboratory, benthic macroinvertebrates and insect samples were separated from the substratum materials, washed with water and isolated to the main taxonomic groups. These were first held in the field in 4% formaldehyde solution [13]. Then, samples were placed in 95% ethyl alcohol solution for the molecular identification process [10].

2.3 Morphological examination:
Compound and dissecting microscope were used for identification and counting of aquatic insect depending on the following key references: [14-21]. The results were expressed as individual /m².

2.4 Molecular evaluation:
After morphological recognition, the specimens were processed individually for molecular analyses at Ankara University's Department of Biological Sciences' molecular biology laboratory (Ankara, Turkey).

Tissue samples were obtained from the thorax, abdomen and if the sample is small, whole organisms were used. Cetyl Trimethyl Ammonium Bromide (CTAB) method [22] was followed to extract the nucleic acids and the details were as follows:- specimen was homogenized in 300 μl + 50 μl BuE mixed and incubated at 65°C for 1 hr.; then 250 μl C: IAA was added, mixed well and centrifuged on 13000 rpm for 15 min. the supernatant was taken and added 500 μl isopropanol and incubated for 30 min at -80°C. then centrifuged for 10 min at 13000 rpm. a pellet was washed with pure ethanol and dissolved the dried pellet with 50 μl TE buffer of extraction buffer (100 mM Tris-HCl (pH 8), 100 mM EDTA (pH 8), 100 mM Na-Phosphate buffer (pH 8), 250 mM NaCl, 2 ± % CTAB) by a sterile hand pestle. After incubating at 65°C at water bath, equal volume of chloroform-isooamyl alcohol (24:1) mixture was added, mixed properly and centrifuged at 12,000 rpm for 15 min at 4°C. The clear aqueous phase was separated in a sterile microcentrifuge tube and 0.6 volume of room temperature isopropanol was added, mixed and incubated at −80 °C for 30-45 min to allow precipitation of the DNA. After incubation, centrifugation was carried out at 13,000 rpm for 10 min at 4°C. The resulting pellet was then washed with 70% chilled ethanol twice and dried. Dried pellets were dissolved in TE buffer and stored at −20 °C for further applications. DNA quality and quantity were measured using a Nano VueTM Plus spectrophotometer.

PCR amplification was conducted using two universal primers. the primer pair LC01490 (5′-
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Family: Baetidae

*Baetis alpinus* (Pictet, 1843):
Body length 7-10 mm. Body greenish brown. Pro- and mesonotum with dark spots and strips. Legs darkened near the apex of femur and tibia, femora with a narrow dark brownish strip along margin. Posterior the margin of femora with numerous long and pointed bristles.

*Baetis braschi* (Zimmermann, 1980):
Body length 7-8.5 mm. Thorax yellowish brown to brown, with indistinct markings on pronotum. Meso- and metanotum with brown longitudinal strips and small lateral spots. Legs uniformly light yellowish-brown. Outer margin of femora with short apically rounded bristles, tarsal claws with 8-13 teeth. Gills 3-5 strongly asymmetric, stout bristles along outer margin absent. Paracercus 2/3-3/4 of cerci length.

*Baetis noa* (Yanai & Gattolliat, 2018):
General color brown, body size 4.7-5.8 mm. Head brown with ecru spots on frons, around compound eyes and around antennal bases, labrum light brown. Thorax brown with a few pale marks but with no clear pattern. Legs whitish, with light brown spot on dorsal femora, usually with proximal and distal brownish marks on femora and tibiae. On all terga a pair of median dark dots are occasionally present. Gills milky, margins dark brown. Cerci and median caudal filament ecru without bands or patterns. Median caudal filament at least 1/2 length of cerci. Gill margins with short spines. Distal margin of terga with many short spatulas.

*Baetis harrisi* (plate 4) (Barnard, 1932):
All small minnow mayflies are characterized by a small spindle-shaped body with 6 or 7 pairs of gills on either side of the abdomen, consisting of a single oval plate without basal tufts. The long antennae are one of the most apparent features of this group which are longer than the head. Their color varies from light sand to dark brown depending on the environmental conditions they are exposed to.

*Iswaeon anoka* (plate 5) (Daggy, 1945):
The length of the body varies between around 4.0-8.0 mm. Hind wing pads are absent. Cerci with a central dark band or three lighter narrower bands. Abdominal segment 5 without dark pigment encircling the segment. Pale stripe running longitudinally in the center of abdominal tergites, claws with an extremely reduced second row of denticles.

*Heterocloeon amplum* (Traver, 1932):
Body length 7-9 mm, body not flattened dorsoventrally. Terminal filaments reduced, hind wing pads present. Segment 2 of labial palps with a developed projecting corner, abdominal tergal scale like setae present. Fore coxa without filamentous gill.

Order: Diptera

Family: Simuliiidae

*Simulium venustum* (plate 6) (Say, 1823):
Head capsule usually complete and fully exposed, head showing distinct constrictive separation from the thorax. Prothorax with 1 proleg or a pair of prolegs head capsule with a pair of folding labral fans dorso-laterally abdomen 5-8 swollen, posterior segment terminating in a ring of radiating rows of hooks labral fans present, head with sides parallel anal sclerite with posterior arms, antenna with proximal articles, hypostoma with the apex of median tooth extended anteriorly.

Order: Coleoptera

Family: Gyrinidae

*Orectochilus villosus* (plate 7) (Müller, 1776):
Legs may be small with 3-6 defined segments, legs 5 segment, tarsi 2 claws, abdomen with 2 pairs of stout terminal hook on segment 10, abdominal segments 1-9 bearing lateral gills, body long, greyish color, anterior fronto-clypeal margin truncate in the middle.

Order: Megaloptera

Family: Corydalidae

*Nigrania serricarnis* (plate 8) (Say, 1824):
The larvae are slightly flattened, up to 33mm length. The head antero-laterally possesses 4-5 segmented setaceous antennae and eyes. The mouthparts chewing type. The prothorax is heavily sclerotized with the pronotum relatively large and sub-rectangular. Each thoracic segment bears a pair of well-developed legs consisting of 5 segments. Tarsi are 1-segmented and at the apex bear a pair of hook-shaped tarsal claws. S-shaped body with lateral abdominal filaments without ventral gill tufts.

Order: Plecoptera

Family: Leuctridae

*Leuctra hippopoides* (plate 9) (Kacanski & Zwick, 1970):
Elongate narrow nymphs, without ventral gill tufts on thorax, gills, not conical. The sternum not overlapping, paraglossa and glossa produced forward in the same distance, 2nd tarsal segment shorter than 1st, midline of wing pads parallel, no cervical gills, at most abdomen 1-7 separated by membranous pleurad fold, abdominal segments cylindrical, hind wing pads longer than wide, abdominal terga with a posterior fringe of short or long setae, and its last few segments with 2-4 long setae, more than 12 bristles on each side of the pronotum.

*Leuctra inermis* (plate 10) (Kempny, 1899):
Elongate nymph’s length from 6-9 mm. without ventral gill tufts on the thorax, gills, not conical, the sterna not overlapping, paraglossa and glossa produced forward in the same distance, 2nd tarsal segment shorter than 1st, midline of wing pads parallel, extend hind legs reach far short of ab tip. Abdomen parallel sided, abdominal terga with a posterior fringe of short or long setae, and it is the last few segments with 2-4 long setae, abdominal terga 2-10 each with one pair of long dorsolateral bristles.

Order: Odonata

Suborder: Zygoptera

Family: Lestidae

*Leses inaequalis* (plate 11) (Walsh, 1862):
Nymph slender, head wider than thorax and abdomen, 3 long caudal gills, 1st antennal segment not so elongate, prementum with a deep open median cleft, lateral caudal gills
triangular in cross section, pre-mentum stalked and spoon shaped, a movable hook of each palpal lobe with 2 or 3 setae, distal margin of each palpal lobe divided into 4 processes.

Order: Trichoptera
Family: Hydropsychidae

Leptonema albovires (Walker, 1852):

Larvae construct cylindrical portable cases, anal claw with stout apical hook, metasternum entirely covered by sclerite, abdomen with ventrolateral rows of gills, a tuft of long setae at anal proleg, genae touching ventrally, separating ventral apotome into anterior and posterior parts, posterior ventral apotome no longer than broad, abdominal gills with up to 40 filaments, fore trochanters never forked, tibia and tarsus of each prothoracic leg lacking dense steal fringe, dorsum of head convex and without carina.

Hydropsyche simulans (plate 12) (Ross, 1938):

Larvae construct cylindrical portable cases, anal claw with stout apical hook, metasternum entirely covered by sclerite, abdomen with ventrolateral rows of gills, a tuft of long setae at anal proleg, genae ventrally, separating apotome into anterior and posterior parts or inconspicuous abdominal gills with about 10 filaments near apex of central stalk, fore trochanters usually forked, abdominal sternum viii with pair of sclerite, prosternum with pair of large sclerites in the intersegmental fold, hydro dorsum of the abdomen with numerous plain hairs. Around 16 mm in length.

Arctopsyche irrorata (Banks, 1905):

Larvae length 23-27 mm; head black, pale mid-ventrally, with distinct pattern of pale spots dorsally and mostly confined to fronto-clypeus; thoracic nota with pale median line and mottled laterally. Larvae construct cylindrical portable cases, anal claw with stout apical hook, meta-sternum entirely covered by sclerite, abdomen with ventrolateral rows of gills, tuft of long setae at anal proleg, genae ventrally, separating apotome into anterior and posterior parts or inconspicuous, genae of head capsule completely separated by single ventral apotome, most abdominal segments with long seta, ventral apotome narrowed.

Family: Hydrometridae

Ochrotrichia tenuata (Plate 13) (Blickle & Denning, 1977):

Larvae up to 5.5 mm; abdomen laterally compressed; thoracic legs the same length, foretibia with posteroventral lobe; without anal gills; cylindrical portable cases, consisting of 2 silk valves covered with sand. Meta-sternum covered by sclerite, sometimes pigmented and sometimes with a transverse anteriorly positioned membranous area. Abdomen without branched gills or tuft of setae, anal proleg not projecting from the abdomen. Found in many different types of streams and rivers, on rocks in algal mats and in moss in current.

4. Molecular Analysis

The identification of aquatic insects based on DNA analysis shows high diversity of species and provides much promise as a method for taxonomic study and as a basis for phylogenetic analysis. It is becoming increasingly popular because these insects are excellent indicators of aquatic environmental quality [32,33].

In the present study, the phylogeny of 30 aquatic insect’s larvae in the Greater Zab River, was analyzed by using two molecular markers as cytochrome c oxidase subunit I (Cox1) sequences and 16S RNA Figure 2.

The phylogenetic analysis based on barcode region of the mitochondrial DNA cytochrome c oxidase subunit I sequence ranging 650-660 bp were recovered revealed grouping of five main clades. The first clade, Baetis alpinus and Baetis brauschi are in the same sub-branch with the bootstrap value of 25%, because of differences of environmental factors as [32,34] stated that the environment of the river changes dramatically and hierarchically along its course and the cross-section of a river has a very uneven and diverse habitat on a microhabitat scale.

While, Baetis naa, Serratella ignita, Leuctra hippopodias and Anapos kugleri are single clad (II, III, IV and IV) respectively, because they belonged to different orders and families. In agreement, [35] determined the monophyly of the families Heptageniidae, Baetidae, and Ephemerellidae. Also, [36] endorsed there was no significant Bayesian or parsimony support for the monophyly of the eastern group of Leuctra hippos. Whereas, phylogenetic analysis based on ribosomal RNA nucleotide sequence revealed grouping of four (IV) main clades as shown in Figure 3. The first clad (I) includes Maccaffertium erritudinum and Leuctra inermis at bootstrap 48%. In a study, [37] showed phylogenetic inference of Leuctra inermis and suggested sister taxon relationships between morphologically similar species because of similarity in morphological characters and linked in geographical location of populations. The Clade II includes Orectochilus villus at bootstrap 55%, with clad one. Also, [38] recognized genus-group taxa of (Coleoptera: Carabidae) supported as monophyletic. While previely, [39] showed that Orectochilus sp. was paraphyletic and mitochondrial COI gene as a useful molecular marker for Gyriidae phylogenetic studies. In clad III, Ephemerella cornutus was a single clad. The clad IV at bootstrap 100% includes Hydropsyche simulans and Baetis harrisoni. Recently, [40] confirmed that the results strongly supported the monophyly of the Caenidae, Heptageniidae, Isonychiidae and Vietnamellidae families. On the other hand, [41] found that the family Hydropsychidae was monophyletic based on mitochondrial cytochrome oxidase 1 (658 base pairs).

It was also endorsed by relatively high genetic distances to other species. Several studies of DNA barcoding highlighted the cases where this approach was found unsuitable for species identification [42,43]. However, the results obtained in the studies on mayflies show a close
correspondence between the morphologically defined species and the barcode clusters [23,44-47].

However, in this work, we are unable to identify all small larvae precisely at the species level but obtaining sequences from particular specimens. Thus, these results may be regarded to variations in sampling sites as habitat characteristics are very important aspects when considering species distribution, as agreement with Heino and de Mendoza [48].

Overall, our findings suggest that DNA barcoding provides an excellent foundation for macroinvertebrate species identification (including insects) and offers new insight on local biodiversity assessment.

Plate 1. Ecdyonurus dispar.
Plate 2. Stenonema femoratum.
Plate 3. Seratella ignita.
Plate 4. Baetis harrisoni.
Plate 5. Iswaeon anoka.
Plate 6. Simulium venustum.
Plate 7. Orectochilus villosus.
Plate 8. Nigrania serricarnis.
Plate 9. Leuctra hippocoides.
Plate 10. Leuctra inermis.
Plate 11. Lestes inaequalis.
Plate 12. Hydropsyche simulans.
Plate 13. *Ochrotrichia tenuata* (A) whole body; (B) cylindrical portable cases.

Table 1. Show Coordination of studied locations.

| Site | Location | Latitude   | Longitude   | Elevation |
|------|----------|------------|-------------|-----------|
| 1    | Ava sheen| 37°01′05″ N | 43°50′09″ E | 550       |
| 2    | Sheladzae| 37°01′25″ N | 43°45′36″ E | 570       |
| 3    | Ble      | 36°52′15″ N | 44°03′06″ E | 477       |
| 4    | Rezan    | 36°50′33″ N | 44°08′26″ E | 446       |
| 5    | Khalan   | 36°41′26″ N | 44°20′08″ E | 406       |
| 6    | Qandel   | 36°39′23″ N | 44°13′40″ E | 375       |

Figure 2: Molecular Phylogenetic analysis of COI by Maximum Likelihood method

The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model (1). The tree with the highest log likelihood (−4214.58) is shown. The percentage of trees in which the associated taxa clustered together is shown next to the branches. 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. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 8 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All positions containing gaps and missing data were eliminated. There was a total of 625 positions in the final dataset. Evolutionary analyses were conducted in MEGA7 (2).
The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model (1). The tree with the highest log likelihood (−3087.57) is shown. The percentage of trees in which the associated taxa clustered together is shown next to the branches. 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. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The analysis involved 10 nucleotide sequences. Codon positions included were 1st + 2nd + 3rd + Noncoding. All positions containing gaps and missing data were eliminated. There was a total of 367 positions in the final dataset. Evolutionary analyses were conducted in MEGA7 (2).

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