

**Bangor University** 

# MASTERS BY RESEARCH

Assessing the Utility of Aqueous eDNA for Invertebrate Biodiversity Assessment in Reens and Ditches.

Harris, Sophie

Award date: 2023

Awarding institution: Bangor University

Link to publication

#### General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

You may not further distribute the material or use it for any profit-making activity or commercial gain
You may freely distribute the URL identifying the publication in the public portal ?

# Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.



# Assessing the Utility of Aqueous eDNA for Invertebrate Biodiversity Assessment in Reens and Ditches.

# Sophia Harris

2021-2023

A thesis submitted to Bangor University in candidature for the degree Master of Science by Research



School of Natural Sciences Bangor University, Deiniol Road, Bangor, LL57 2UW

# Contents

Declaration						
Ac	Acknowledgements					
AŁ	Abstract4					
1.	Introduction	5				
	1.1 Freshwater Ecosystems	5				
	1.2 Freshwater Monitoring Systems	5				
	1.3 Morphological Macroinvertebrate Monitoring Methods	8				
	1.4 eDNA Biomonitoring in Freshwater Systems	10				
	1.5 Study Site	12				
	1.6 Study Aims	14				
2.	Materials and Methods	14				
	2.1 Sample Collection	14				
	2.2 DNA Extraction and Sequencing	16				
	2.3 Bioinformatics	17				
	2.4 Statistical Analysis	18				
	2.5 Methods of previous surveys within St. Brides SSSI	19				
	2.6 Ecological Quality Indices Assessment	19				
3.	Results	20				
	3.1 Pipeline Output	20				
	3.2 Invertebrate Diversity	20				
	3.3 Comparison of Previous Surveys	25				
	3.4 Ecological Quality Indices	26				
4.	Discussion	27				
	4.1 Water Quality Effect on Biodiversity	28				
	4.2 Abiotic Factors Influencing Biodiversity	29				
	4.2.1 Water Filtration Quantity	29				
	4.2.3 Temperature	30				
	4.2.4 pH	31				
	4.2.10 Seasonality	37				
	4.3 eDNA as a Tool for Assessing Water Quality	38				
5.	Conclusion	39				
6.	References	40				
7.	Appendices	50				
	Appendix II. Complete Species List at 99% filtration identity level.	51				
	7.1 Appendix i. Biological Monitoring Working Party (BMWP) Average Score per Taxon (ASPT) Scoring System	67				

# Declaration

I hereby declare that this thesis is the results of my own investigations, except where otherwise stated. All other sources are acknowledged by bibliographic references. This work has not previously been accepted in substance for any degree and is not being concurrently submitted in candidature for any degree unless, as agreed by the University, for approved dual awards

\_\_\_\_\_

-----Yr wyf drwy hyn yn datgan mai canlyniad fy ymchwil fy hun yw'r thesis hwn, ac eithrio lle nodir ynwahanol. Caiff ffynonellau eraill eu cydnabod gan droednodiadau yn rhoi cyfeiriadau eglur. Nid ywsylwedd y gwaith hwn wedi cael ei dderbyn o'r blaen ar gyfer unrhyw radd, ac nid yw'n cael eigyflwyno ar yr un pryd mewn ymgeisiaeth am unrhyw radd oni bai ei fod, fel y cytunwyd gan yBrifysgol, am gymwysterau deuol cymeradwy.

# Acknowledgements

First and foremost, I would like to thank National Resources Wales for their generous contribution to this project. Due to their interest and funding, I could conduct my Master's project from sample collection to writing. I sincerely hope the work presented in this paper can aid their future management plans to conserve freshwater invertebrates in St. Brides SSSI. Furthermore, a special thanks to Dr Andrew Lucas from NRW for obtaining the funding and arranging to access private land within St. Brides. Alongside this, I am grateful to the landowners for allowing me to conduct my research.

My gratitude also goes to Professor Simon Creer from Bangor University. Simon took me on as a Master's student and allowed me to conduct my research in an area of great interest to me. Furthermore, Simon's contribution aided my development as a scientist and ensured that my personal developmental goals were reached to help me achieve my career plan.

Thirdly, a massive thank you to Dr William Perry for his unwavering support throughout the project. His support and wisdom have been invaluable, from sample design to collection to my write-up. I genuinely could not have completed this project without him, and I am forever grateful.

Another debt of gratitude is owed to my second supervisor Dr Amy Ellison. Working with her in the laboratory has been an incredible opportunity, and I can genuinely say I have learnt my lab skills from the best. Her patience is greatly appreciated; she has cemented my passion for the lab for future research projects.

A special mention goes to those in the Molecular Ecology and Evolution group at Bangor (MEEB). The support from my peers gave me hope and determination, and I will cherish the friendships and memories I made throughout my time at Bangor. I will take the wisdom shared with me wherever I go and remember their encouraging words when things do not go to plan in the future.

Finally, a massive 'thank you' to my friends and family. Their support has been extremely important to me regarding completing my project, and they always knew the right thing to say when I felt overwhelmed and celebrated with me when I made even the smallest of achievements. I look forward to celebrating the completion of this thesis with everyone.

# Abstract

Observing aquatic invertebrate diversity can provide ecological insights into changing environments. Research into moving waterbodies has primarily focused on rivers, with little exploration into biodiversity within ditches. Ditches often drain into larger 'reens', which are artificial structures designed to prevent water-logging in winter and provide livestock with water in summer There has been little investigation into ditches and reens as important habitats for invertebrates, partly due to difficulty surveying them. Often, macroinvertebrate surveys use morphological monitoring methods to understand the area's biodiversity; however, these methods are often time-consuming, require expert knowledge, and are highly invasive. In this study, an alternative method using environmental DNA (eDNA) was applied to understand the alpha and beta diversity of ditches and reens in St. Brides SSSI, South Wales. Here, the data presented demonstrated that eDNA analysis produces far greater taxonomic information than morphological analysis and that abiotic factors, such as waterbody type and temperature, significantly impact alpha diversity, while the amount of water filtered and salinity influenced beta diversity. These results indicate the importance of utilizing consistent methods during water sample collection. In addition, ditches and reens showed differences in invertebrate diversity despite the waterbodies being connected. We anticipate that the findings from this study can aid ditch and reen management plans to ensure that invertebrate biodiversity is maintained. Furthermore, this study highlights the importance of using invertebrates as indicator species for water quality assessment and displays the benefits of using environmental DNA monitoring in combination with morphological monitoring.

# 1. Introduction

## **1.1 Freshwater Ecosystems**

Freshwater ecosystems, rich in species diversity and endemism, are essential to sustaining human existence, with human settlements forming preferentially near freshwater (Revenga *et al.*, 2005). However, human interaction causes anthropogenic pressures such as pollution, exploitation, habitat degradation and the introduction of invasive species (Kuntke *et al.*, 2020). At both regional and local levels, freshwater environments are often in poor condition for biodiversity compared to terrestrial habitats (Clarke, 2015). Changes in freshwater ecosystems may not be visible, and declines in biodiversity can occur for long periods without detection (Linke *et al.*, 2018).

One major cause of freshwater species loss has been the simplification and channelization of rivers and their floodplains. In some areas of the United Kingdom, to counteract species loss, drainage schemes in floodplains and low-lying areas have created networks of channels designed to carry water and maintain lower water levels, providing flood control and prevention of erosion or improved navigation (Brookes, 1985, 1986; Keller, 1976). Artificial drainage networks are characterized by larger and smaller channels, referred to here as ditches. Many of the ditches now surround areas of agriculture or urban zones and are managed to maintain specific water levels through regular vegetation management. Despite these anthropogenic pressures, ditches provide an essential freshwater environment for wildlife and have been found to support a wide variety of invertebrate species, including species of conservation interest.

## **1.2 Freshwater Monitoring Systems**

To develop a standardized monitoring system that could provide insight into water quality, the Institute of Freshwater Ecology (IFE) developed a technique for evaluating the biological quality of rivers in the UK (Wright, 1994). The River Invertebrate Prediction and Classification

System (RIVPACS) focuses on benthic macroinvertebrates for biological assessments as macroinvertebrate taxonomy is well known, and a wide variety of species are found in freshwater habitats (Hellawell, 2012; Wright, 1994). RIVPACS, a statistical model software, incorporates the General Quality Assessment classification of river statuses based on Ecological Quality Indices (EQIs) based on two indices (Paisley, Trigg and Walley, 2014). The first index is The Biological Monitoring Working Party (BMWP), which has been used by regulatory authorities in the UK since the 1980s as the basis of their river invertebrate status classification system (Paisley, Trigg and Walley, 2014). The BMWP provides allocated scores to families based on their sensitivity to pollution (Table 1). The more sensitive taxa are to pollution, usually due to Biological Oxygen Demand (BDO), the higher the BMWP score (Paisley, Trigg and Walley, 2014; Aquilina, 2013). The second index incorporated into the EQIs is the average BMWP score per taxon, also known as the Average Score Per Taxon (ASPT). The ASPT was created due to bias experienced when conducting score systems for invertebrate monitoring, as the sample size primarily affects the number of taxa in a sample. To counteract this bias, the ASPT was developed, whereby the BMWP score is divided by the number of contributing taxa (Appendix 1), thus providing an average score (Hawkes, 1998).

BMWP score	ASPT score	Quality interpretation
>150	>6	Very good
101-150	>5	Good
51-100	>4	Moderate
16-50	<4	Poor
0-15		Very Poor

**Table 1.** The Biological Monitoring Working Party (BMWP) and Average Score Per Taxon (ASPT) scoring system.

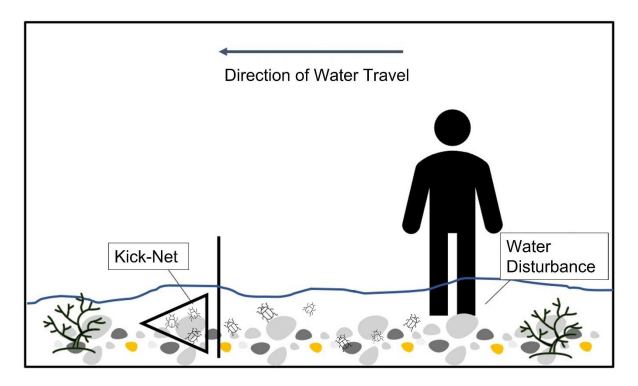
In December 2000, the EU Water Framework Directive (WFD) was created as the primary legislative tool to improve polluted inland water surfaces, transitional, coastal and ground waters to prevent a decline in water quality using monitored macroinvertebrates as biological quality elements (European Commission, 2021; Lathouri *et al.*, 2021). The WFD has monitored the status of aquatic ecosystems by characterizing biological communities and physiochemical and hydromorphological conditions using the BMWP and ASPT (Pawlowski *et al.*, 2018). The WFD sets ecological water surface standards for 27 countries globally and is divided by the range of EQI scores into five classes: very good, good, moderate, poor and very poor (European Commission, 2021).

Monitoring invertebrate populations can provide an early warning for ecological change by observing the presence and abundance of species and allowing for the development of mitigations to slow species decline (Schmeller, 2008; Beever, 2006). Biological monitoring has several limitations (practical and theoretical) that need to be considered to allow them to be applied successfully (Beever, 2006). Despite this, those conducting surveys (e.g. government agencies, local councils, water companies, and researchers) often find themselves in a difficult position where data needs to be gathered quickly, at a low cost, and perhaps without clear objectives (Witmer, 2005). Invertebrate studies are often complicated as species of interest are often poorly understood, rare or strongly influenced by human activities (Witmer, 2005).

The first step in an invertebrate survey is sample collection. Many aquatic invertebrate surveys sample relatively small areas or volumes and only collect a small proportion of species in the survey area (Halse *et al.*, 2002). Aquatic macroinvertebrate surveys are often conducted using equipment such as the Surber sampler and kick-nets (Poikane *et al.*, 2016; Brua, Culp and Benoy, 2011; Sharma, Arambam and Sharma, 2009).

#### **1.3 Morphological Macroinvertebrate Monitoring Methods**

Kick-net (Figure 1) and U-net sampling are recognized internationally for small, regional, and national bioassessments (Brua, Culp and Benoy, 2011). Both methods involve a net being held downstream while the substrate upstream is disturbed, causing invertebrates to pass into the net. While the methods have been proven effective at obtaining macroinvertebrates, they are highly invasive due to the disturbance of the water substrate and the removal of specimens (Brua, Culp and Benoy, 2011). However, these sampling methods are low-cost, easy to transport, and valuable for surveying various habitats (Carter and Resh, 2001). Conversely, the netting size significantly affects biodiversity assessment, as smaller organisms are often missed by passing through the net or becoming trapped within silt, clogging the net, and larger organisms actively climbing out of the net. In addition, collected invertebrates are often difficult for biologists to identify, and the sample size varies depending on the season due to fluctuations in water levels (Stein, Springer and Kohlmann, 2008; Carter and Resh, 2001). High currents caused by increased water levels can also increase the likelihood of organisms missing the net (Stein, Springer and Kohlmann, 2008).



**Figure 1.** Kick-net sampling is a standard method used for sampling macroinvertebrates. The researcher stands upstream, disturbing the water so that invertebrates are swept downstream into the net.

Surber samplers are nets of a given mesh size fastened around a heavy metal square frame, allowing a known isolated area of the riverbed to be sampled (Surber, 1937). They are used because they are practical and highly suited to varying habitats (Ghani *et al.*, 2016). However, they have a small substrate area and fail to collect macroinvertebrates drifting in the water column above the sample (Ghani *et al.*, 2016). The small size of the sampler also makes it difficult to place it correctly on rough substrates (Al-Shami *et al.*, 2013). Larger specimens can crawl out of the samplers, and others can detect the physical disturbance in the water and avoid the nets altogether (Ghani *et al.*, 2016). Despite this, the quantitative collection from the Surber sampler is reliable in estimating the abundance and diversity of benthic invertebrates (Ghani *et al.*, 2016).

#### 1.4 eDNA Biomonitoring in Freshwater Systems

Environmental DNA (eDNA) metabarcoding is becoming an increasingly valuable and viable tool for ecologists (Taberlet *et al.*, 2018). eDNA analysis is a non-invasive molecular species detection method, and eDNA is defined in this paper as DNA deposited from organisms into the environment, which can be collected in samples such as water soil and sediment, without the presence of the organism (Ruppert, Kline and Rahman, 2019; Taberlet *et al.*, 2018; Turkelboom *et al.*, 2013; Sogin *et al.*, 2006). The eDNA can then be extracted, amplified, sequenced, and categorized, allowing for species identification (Ruppert, Kline and Rahman, 2019; Deiner *et al.*, 2015). The amount of eDNA an individual produces depends on biomass, age and feeding activity of the organism, physiology, life history, and use of habitat space (Hering *et al.*, 2018; Barnes and Turner, 2016; Goldberg, Strickler and Pilliod, 2015).

DNA metabarcoding uses PCR and the deployment of taxonomic-specific oligonucleotide primers, combined with high-throughput sequencing to enable multi-species identification, often with DNA extracted from an environmental sample (Taberlet *et al.*, 2012). However, the definition can also be applied to species identification from bulk samples of entire organisms where they have been isolated prior to analysis (Taberlet *et al.*, 2012; Hajibabaei *et al.*, 2011; Chariton *et al.*, 2010; Creer *et al.*, 2010; Porazinska *et al.*, 2010). Furthermore, metabarcoding has been found to achieve comparable assessment results with morphological studies and offers a powerful alternative, identifying species that would otherwise be unfeasible in morphological studies (Elbrecht *et al.*, 2017).

Environmental DNA metabarcoding combines traditional field-based ecology with in-depth use of molecular methods and advanced computational tools (Ruppert, Kline and Rahman, 2019). While it is still an emerging monitoring method, it can revolutionize modern biodiversity surveys (Ruppert, Kline and Rahman, 2019). eDNA is beneficial for monitoring aquatic taxa due to the DNA shed by an organism into the surrounding environment, which can persist in lotic waters. Understanding the total biodiversity of rivers is fundamental to determining surface water

quality (Fernández *et al.*, 2019; Deiner and Altermatt, 2014). Monitoring using eDNA has been compared to traditional monitoring methods, and it has been concluded that DNA metabarcoding and morphological identification give similar correlations with water quality instream conditions when linked to the watershed size and shifts in forest composition across various water bodies (Emilson et al., 2017; Fernández et al., 2019). However, inconsistencies remain in the taxonomic composition produced by the two approaches, especially regarding macroinvertebrate and microbial communities (Keck *et al.*, 2022).

eDNA pipelines can facilitate high throughput processing of samples compared to low throughput traditional methods, allowing for greater replication and more geographically and temporally broad surveys (Seymour *et al.*, 2020). eDNA can also better detect ecological signals than morphological methods, presenting higher taxonomic richness due to the improvement of taxon assignment in some groups (e.g. midges, mayflies, caddisflies and black flies) (Fernández *et al.*, 2019). Multi-gene eDNA studies have also been compared against traditional methods revealing interactive networks linked to ecological assessment criteria (Seymour *et al.*, 2020). Moreover, eDNA analysis can potentially include a broader range of taxa and indicator groups that would otherwise fail to be included in traditional taxonomic identifications, thus focusing on higher levels of invertebrate biodiversity than morphological identification methods (Seymour *et al.*, 2020).

The biomonitoring capabilities of eDNA analysis has meant that it is a powerful tool for conservation, and many studies have focused on detecting invasive species in natural systems. This was first demonstrated by targeting American bullfrogs (*Rana catesbeiana*) in French wetlands (Ficetola *et al.*, 2008). Research has also detected invasive species in transit, such as non-native organisms in the ballast waters of transoceanic ships (Egan *et al.*, 2013; Mahon *et al.*, 2013; Li *et al.*, 2011). Furthermore, similar studies have successfully identified benthic invertebrates and their resting stages within ballast tank sediments (Briski *et al.*, 2011;

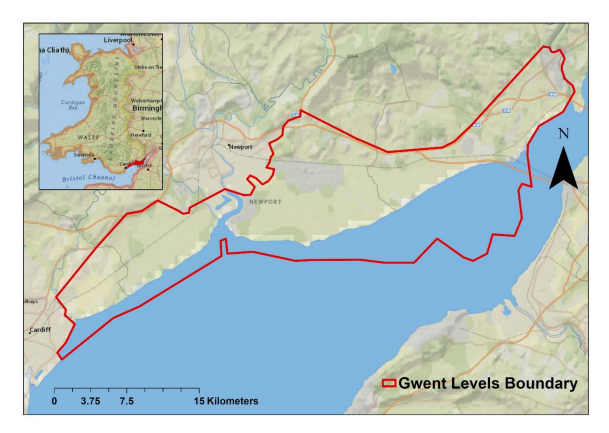
Harvey, Hoy and Rodriguez, 2009; Darling and Tepolt, 2008). Conversely, eDNA has been used to locate rare indigenous species; for example, in the UK, the great crested newt (*Triturus cristatus*) has been a focus of many eDNA surveys (Rees *et al.*, 2014). As eDNA analysis is less invasive, it is more applicable than traditional approaches in certain situations and becoming an increasingly used tool (Stein *et al.*, 2014; Mahon *et al.*, 2013).

# 1.5 Study Site

This study focused on the Gwent Levels (Figure 2), a lowland area between Cardiff and Chepstow, UK, where a network of drainage ditches has been maintained since the roman ages (Living Levels, 2021; Countryside Council Wales, 1991). The Levels are an example of one of the most extensive areas of reclaimed wet pasture in the UK and are rich in biodiversity, including in aquatic invertebrates (CCW, 1991). The Levels have provided habitat for nationally rare or notable species such as the water beetles *Halipus mucronatus* and *Hydrophilus piceus*. The area is also crucial in Wales for its snail and dragonfly populations, including *Physa heterostropha* and *Brachytron pratense* (CCW, 1991). St. Brides SSSI is one in a series of SSSIs located in the Gwent Levels consisting of 5700 hectares (Countryside Council Wales, 2008). The entire landscape is artificial and formed from a 2,000-year history of land reclamation from the sea (Living Levels, 2021). The land is below sea level; however, extensive sea defences prevent submersion (Living Levels, 2021).

Ditches are artificial bodies of water primarily draining excess water and groundwater seepage from agricultural lands in winter while providing livestock with drinking water in warmer months (Verdonschot, 2012). Field ditches are relatively small in width and depth and have varying water volumes and vegetation cover around the edges of agricultural fields. Field ditches feed into more extensive reens, usually between 2-8 m wide and up to 1 m deep, which are generally well maintained. Both ditch types are found in temperate and boreal zones in the Northern Hemisphere in almost all low-lying or wetland areas (Herzon and Helenius, 2008).

With little water movement, field ditches and reens have intensive organic and inorganic matter exchange with the surrounding terrestrial matrix (Herzon and Helenius, 2008). As a result, they must be regularly managed, mowing aquatic vegetation to avoid accumulated sediment (Twisk, Noordervliet and ter Keurs, 2000; Beltman, 1984). Without such maintenance, complete territorialization would occur (Verdonschot, Keizer-vlek and Verdonschot, 2011). The ditches are drained by gravity, pumps, and other water controls (i.e., sluices) to control water levels. Water levels are kept high in summer for fencing and providing livestock water. In winter, however, structures that maintain high water levels are removed to allow floodwater to drain into the estuary when the tide goes out. Many reens and ditches on the Gwent Levels are periodically cleared as part of management regimes to prevent silting (CCW, 2008).



**Figure 2.** The Gwent Levels comprise eight SSSIs on the south coast of Wales, stretching from the east of Cardiff to Caldicot. Sources: National Geographic, Esri, Garmin, HERE, UNEP-WCMC, USGS, NASA, ESA, METI, NRCAN, GEBCO, NOAA, increment P Corp.

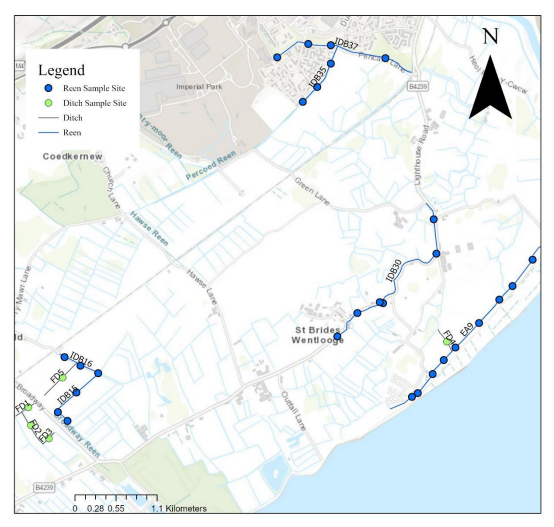
## 1.6 Study Aims

While eDNA is used more frequently to explore aquatic invertebrate biodiversity, it is often not utilized in monitoring schemes and water quality surveys. Here, we explore how alpha ( $\alpha$ ) and beta ( $\beta$ ) diversity differ between ditches and reens in the Gwent Levels while providing a comprehensive insight into the community composition of St. Brides SSSI using eDNA. We investigate how a range of physiochemical parameters and land use impacts diversity, measured via eDNA analysis. In addition, we identify critical methodological considerations and the relative inputs from terrestrial and aquatic systems when sampling from ditches and reens. Finally, we compare how eDNA biomonitoring in the Gwent Levels compares to traditional morphological monitoring techniques through species analysis of the previous surveys.

# 2. Materials and Methods

#### 2.1 Sample Collection

Water samples consisting of 26 reen samples and five ditch samples were selected across St. Brides SSSI, South Wales (Figure 3) from December 1<sup>st</sup> – 6<sup>th,</sup> 2021. Only five ditch sites were selected due to high organic content clogging the filters after filtering so it was thought little DNA could be extracted from the samples. The amount of water filtered at the ditch per replicate varied significantly, with FD2 and FD3 filtering an average of 10ml, compared to FD5, which filtered an average of 867ml. Each EA9.1 replicate also filtered an average of 10ml, compared to IDB16.1 and IDB16.2, which filtered the full 1000ml. However, most reens filtered ~ 200ml per replicate. The sites were selected to capture both urban and rural environments. Three ecological replicates were taken at each site, and three negative controls were taken in the field on the last day. Each water sample was collected using sterilized 1-litre bottles. The water was pumped through a Sterivex filter using a Geopump<sup>TM</sup> Peristaltic Pumps until the filter for each replicate became clogged before adding 1ml of Qiagen lysis buffer ATL. The Sterivex filter was then sealed with a Leur Lock Syringe Cap and stored at 4°C until extraction. Water depth was measured at each site along with the following water physiochemical measurements: depth (cm), sample amount (ml), waterbody type, pH, RDO concentration (mg/L), salinity (PSU), turbidity (NTU), temperature (°C). Physiochemical measurements were taken last to prevent contamination using an Aqua Troll 500, while depth was measured with a measuring stick.



**Figure 3.** A total of 33 sites were selected in the St. Brides SSSI on various reens and ditches covering both urban and rural sites to create sample diversity. Sources: Esri, UK, Esri, HERE, Garmin, GeoTechnologies, Inc., USGS, METI/NASA.

#### 2.2 DNA Extraction and Sequencing

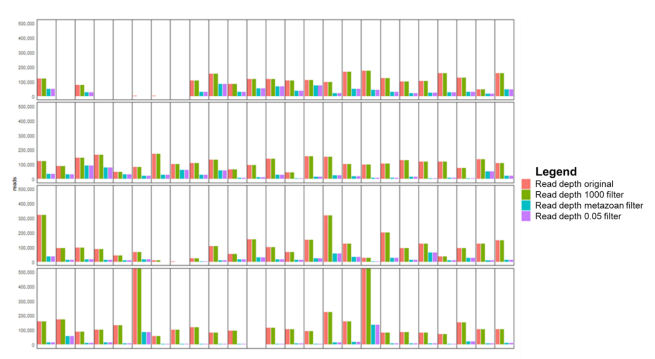
DNA extractions and PCR (Polymerase Chain Reaction) reactions took place in the Molecular Ecology and Evolution laboratories at Bangor University. The extractions were conducted using the Spens et al. (2016) Capsule Methodology protocol and the Qiagen DNeasy Blood and Tissue Kit. Three negative controls from the field were extracted, and a negative PCR control was added to each PCR plate, along with a positive control. COI primers m1COIintF (5'-GGWACWGGWTGAACWGTWTAYCCYCC-3') and jgHCO2198 (5'-TAIACYTCIGGRTGICCRAARAAYCA-3') with universal tails were used (Leray et al., 2013). Each sample library was amplified using the primer set in triplicates and a Qiagen Mastermix kit using an identical library build protocol as in Brennan et al. (2019). Specifically, a 2-step PCR approach was used with a 25 µL total volume consisting of 12.5 µL Qiagen multiplex PCR Mastermix, 0.5 µL of forward primer, 0.5 µL of reverse primer, 10.5 µL PCR grade water and 1 µL template DNA. A Thermal Cycler was used for the following COI PCR Protocol; 95°C 15 mins, then 35 cycles of 94°C 30s; 54°C 90s; 72°C 1min; 72°C 10 min. The triplicates were pooled, and the products from the first round of PCR were purified using 9 µL of Agencourt AMPure XP beads. The second round PCR protocol consisted of 12.5 µL of the Qiagen multiple PCR Master Mix, 1 µL of Round 2 primer index i5-i7, 6.5 µL of PCR water and 5 µL of purified Round 1 PCR product. The PCR machine was run for 10min at 95°C, ten cycles of 30s at 98°C, 30s at 65°C and 30s at 72°C. AMPure XP purification was performed using 12 µL of the Agencourt AMPure XP to 20 µL of the Round 2 PCR product. The PCR round 2 products were quantified using a Qubit dsDNA Broad Range kit, and the equimolar concentrations for the library were pooled before being cleaned using Agencourt AMPure XP beads. Each amplicon library was normalised to 4ng/l by diluting with PCR water and then 1 µL of each normalised amplicon was added to the final pool. The final library was loaded at 12 pM with 10% Phi-X spike-in, and sequenced using an Illumina MiSeq® Reagent Kit v2 (500 cycle) in accordance with the manufacturer's instructions at the Centre for Environmental Technology (CEB), Bangor, Wales.

#### 2.3 Bioinformatics

After obtaining the raw demultiplexed read data, the DADA2 (Callahan et al., 2016) pipeline was run using the statistical software R (R Core Team, 2022), with the default parameters unless specified. By default in the DADA2 pipeline, forward reads with higher than 2 "expected errors" were discarded after trimming and reverse reads with higher than 5 "expected errors" were discarded. Filtering included trimming reads at the first instance of a quality score less than or equal to 2. Basic Local Alignment Search Tool (BLAST) was used for taxonomic assignment in conjunction with the MIDORI COI reference database (Leray et al., 2018; Altschul et al., 1990). Any Amplicon Sequence Variants (ASVs) which did not reach at least 70% percentage identity and query cover were removed from the analysis. Samples with a read depth lower than 1,000 were removed based on the rarefaction curves, and ASVs with a read depth of less than 0.05% of the total reads in a sample were also removed. The proportion of reads per sample were calculated by dividing ASV read depth by total sample read depth. For invertebrate analyses, the following phyla were included: Arthropoda, Gastrotricha, Platyhelminthes, Annelida, Rotifera, Mollusca, Cnidaria, Nematoda, Tardigrada, Porifera, Placozoa, Onychophora, Nemertea, Echinodermata and Bryozoa to obtain a comprehensive overview of invertebrate biodiversity. The greatest impact on read depth was the removal of non-metazoans (Figure 4).

For comparisons with traditional sampling methods and to provide National Resources Wales with a species list, the accuracy of the taxonomic assignment was more important. ASVs were searched against the NCBI Nucleotide database using BLAST with a 1e<sup>-10</sup> e-value threshold. Taxonomic assignments were performed using the "Assign-Taxonomy-with-BLAST" python script (Sevigny, 2018) using default parameters except for a 70% length cut-off to remove potential PCR amplification errors and sample cross contaminations (Macher *et al.*, 2023; Jacot *et al.*, 2021; Corse *et al.*, 2017; Ransome *et al.*, 2017; Leray and Knowlton, 2015) and a 99% species-level assignment threshold to increase identification accuracy when producing

a species list. Invertebrate species were then manually checked to ensure they were UK species.



**Figure 4.** The raw ASVs underwent a series of filtration steps in R studio using the DADA2 pipeline presented on the x-axis, while the number of reads can be viewed on the y-axis. The first step was removing all sample sites with less than 1000 reads. Secondly, only metazoan ASVs remained before those with a depth of less than 0.05 were removed.

# 2.4 Statistical Analysis

Analyses were conducted in R (2022) with scripts provided by Dr William Perry at Cardiff University. All plots were created using the package ggplot2 (Hadley, 2016), apart from the Venn diagram, which was created using the package VennDiagram (Hanbo, 2022). A PERMANOVA (Permutational Analysis of Variance) was used to explore  $\beta$ -diversity, using the adonis2 function in the package vegan (Oksanen *et al.*, 2022), the Bray-Curtis method and 999 permutations. The relationship between the number of ASVs and environmental factors was assessed using a linear model. Furthermore,  $\alpha$  diversity was calculated by observing the number of ASVs. The full model contained the following fixed effects: waterbody type, depth (cm), pH (pH), dissolved oxygen concentration (mg/L), salinity (PSU), turbidity (NTU), temperature (°C) and land use and the amount of water filtered. After a step function using automatic backward elimination, the final model contained water body type, pH, and salinity. Finally, a sparse partial least squares (sPLS) analysis was conducted between 12,278,512 reads and all of the physiochemical variables and then plotted on a heatmap using the package mixOmics (Rohart *et al.*, 2017).

#### 2.5 Methods of previous surveys within St. Brides SSSI

In 2011 Boyce conducted an aquatic macroinvertebrate survey on the Gwent Levels to develop a monitoring strategy designed to aid NRW's commitments under the Common Standards Monitoring programme. The survey was conducted in June and concentrated on collecting aquatic beetles, bugs and snails using Kick-net sampling before the organisms were sorted on a polyethene sheet (Boyce, 2012). The specimens were later organised into the taxonomic groups: Coleoptera, Hemiptera and Mollusca before being identified at the species level (Boyce, 2012). The reens: EA9, IDB15, IDB16, IDB30, IDB35, IDB37 in both the 2011 survey and this eDNA survey.

Graham and Hammond's (2022) survey, which focused on terrestrial and aquatic macroinvertebrates, was conducted in August 2020 using nets and 24-hour bottle traps. The taxa were identified at the species level, either in the field or the laboratory. While this study focused on the Gwent levels, one site, Fair Orchard Farm, was based in St. Brides SSSI.

## 2.6 Ecological Quality Indices Assessment

Water quality assessment was conducted once the sequences were assigned to the family level using the Biological Monitoring Working Party score (BMWP) and the Average Score Per Taxa (ASPT). The BMWP scores were calculated using the BMWP scoring system (Appendix II), in which the score equals the sum of macroinvertebrate families' tolerance scores in each sample (Mandaville, 2002). If a higher BMWP were calculated, it would reflect a better quality

of water in the ditches and reens (Aquilina, 2013). The ASPT can then be calculated to reduce bias as it is the average tolerance score of all macroinvertebrate families within the sample site. The ASPT score ranged from 0 to 10 and was calculated by dividing the BMWP score by the number of families present (Sor *et al.*, 2021).

# 3.Results

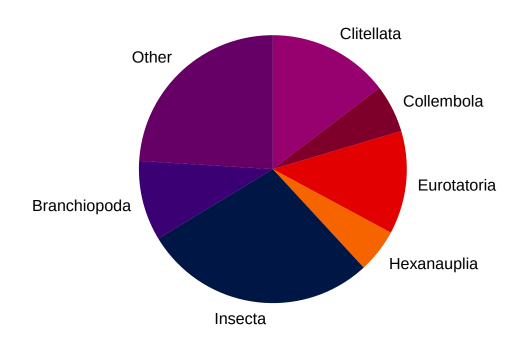
# 3.1 Pipeline Output

A total of 12.2 million paired reads passed through the DADA2 (2016) filtering thresholds. After filtering for metazoan hits only, with a 70% BLAST percentage identity query cover, 2.2 million reads remained, meaning 81% of the reads were of non-metazoan origin. In total, the negative controls had a read depth of <800 and so were removed from the analysis as our baseline was set at a minimum of 1000 reads (Appendix I). To obtain the invertebrate data, the following phyla were included; Arthropoda, Gastrotricha, Platyhelminthes, Annelida, Rotifera, Mollusca, Cnidaria, Nematoda, Tardigrada, Porifera, Placozoa, Onychophora, Nemertea, Echinodermata and Bryozoa. Regarding Class assignment, Insecta (28%) contained the most significant percentage of ASVs, followed by Citellata (15%). In total, 379 species were assigned (Appendix II), including *Dicrotendipes lobiger*, a new species to the Gwent Levels, and Chaetogaster diastrophus, a new species in the UK.

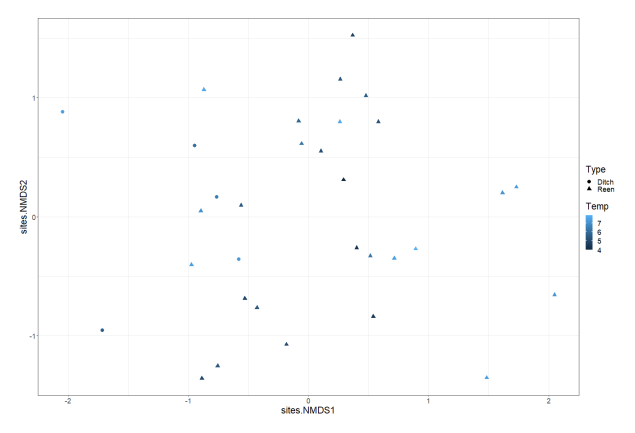
## 3.2 Invertebrate Diversity

The PERMANOVA results (Figure 5) for invertebrates showed that the amount of water filtered ( $F_{1,31} = 1.34$ , Sum sq = 0.50, p = 0.05) and temperature ( $F_{1,31} = 1.40$ , Sum sq = 0.52, p = 0.04) were the only abiotic variables that had a significant effect on beta diversity (Figure 6). The other variables, pH ( $F_{1,31} = 1.21$ , Sum Sq = 0.45, p = 0.1), depth ( $F_{1,31} = 0.97$ , Sum Sq = 0.36, p = 0.55), waterbody (F1, 31 = 1.1, Sum Sq = 0.41, p = 0.25), rugged dissolved oxygen ( $F_{1,31} = 1.26$ , Sum Sq = 0.47, p = 0.1), salinity ( $F_{1,31} = 1.09$ , Sum Sq = 0.40, p = 0.28), turbidity ( $F_{1,31} = 1.26$ , Sum Sq = 0.47, p = 0.1), salinity ( $F_{1,31} = 1.09$ , Sum Sq = 0.40, p = 0.28), turbidity ( $F_{1,31} = 1.26$ ).

= 1.16, Sum Sq = 0.43, p = 0.17) and land use ( $F_{1,31}$  = 0.94, Sum Sq = 0.35, p = 0.61) had no significant effect on the  $\beta$  diversity of invertebrates.

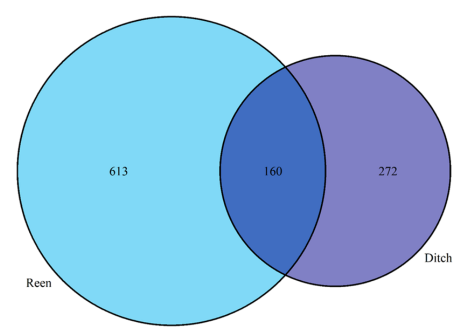


**Figure 5.** A pie chart regarding the division of Class. Insecta was the largest class (28%), followed by Citellata (15%) and Eurotatoria (12%). Branchipodada (10%), Collembola (6%) and Hexanaupalia (5%) all obtained above 5% of assignations compared to the Others (24%), which included; Hydrozoa (4.4%), Malacostraca (4.4%), Polychaeta (4.4%), Arachnida (4%), Gastropoda (3.8%), Bivalvia (1.2%), Chromadorea (1%), Ostracoda (0.9%), Catenulida (0.4%), Diplopoda (0.2%), Rhabditophora (0.2%), Trematoda (0.2%), Cestoda (0.1%), Chilopoda (0.1%), Demospongiae (0.1%), Enoplea (0.1%) and Phylactolaemata (0.1%).



**Figure 6.** The NMDS plots explore the similarities between different variables and invertebrate diversity. Both waterbody type and temperature influenced  $\beta$  diversity. The NMDS plot compares the two different waterbody types, ditches and reens and indicates the impact temperature has on invertebrate diversity, with the colder sites being dark blue.

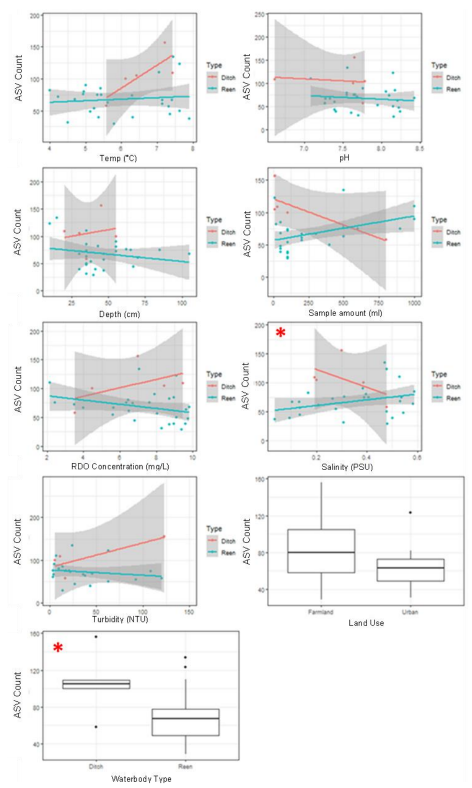
In total, there were 1,045 unique invertebrate ASVs with a greater number of unique invertebrate ASVs in reens, with 613, compared to 272 in ditches (Figure 7) with a 15% species overlap. However, when exploring the  $\alpha$  diversity (Figure 8), there is the implication that the ditches have a higher ASV count when the factor of more reens sampled is removed.



**Figure 7.** A Venn Diagram exploring the relationship of ASVs between ditches and reens. The diagram size is relative to the number of A.S.V.s, with the specific count appearing in the centre. There was only a 15% overlap, which consisted of 160 unique ASVs between ditches and reens.

The average amount of water sampled in ditches was 215 ml compared to 275 ml for reens. Field Ditch 5 (FD5) saw an average of 867 ml filtered, whereas the other ditches combined had an average of 37 ml. Reens IDB15 and IDB16 filtered the highest water quantity; however, that does not appear to have impacted the ASVs per site.

After a step function, the waterbody type, pH, and salinity remained in the final linear model. Alpha diversity was calculated using the number of observed ASVs. Both waterbody type ( $F_{1,28}$  = 8.70, Sum Sq = 6035.0, p = 0.01) and salinity ( $F_{1,28}$  = 4.05, Sum Sq = 2816.3, p = 0.05) had a significant effect on ASV count, but pH did not ( $F_{1,28}$  = 0.43, Sum Sq = 295.4, p = 0.52).



**Figure 8.** The  $\alpha$ -diversity results presenting that water body type (F<sub>1,28</sub> = 8.70, Sum Sq = 6035.0, p = 0.01) and salinity (F<sub>1,28</sub> = 4.05, Sum Sq = 2816.3, p = 0.05) had a significant effect on ASV count. The results indicate that, on average, the ditches had a higher ASV count than reens, although the species present in ditches appear less resilient to salinity as there is a negative correlation.

# 3.3 Comparison of Previous Surveys

Surveys using morphological identification methods have previously been conducted on St. Brides SSSI as the area is a nationally significant assemblage of plants and invertebrate features (Murton, Hunt and Rodgers, 2019; Boyce, 2012). When comparing the eDNA survey results from this study with previous surveys of St. Brides, we see an increase in species detection.

No statistical analysis could be conducted with the previous surveys due to seasonality and location differences. However, when comparing the reens that were used in the eDNA study in the Boyce survey, it was apparent that the eDNA survey detected a far greater number of species than morphological methods (Figure 9). Furthermore, there was minimal overlap between species detected in both surveys.

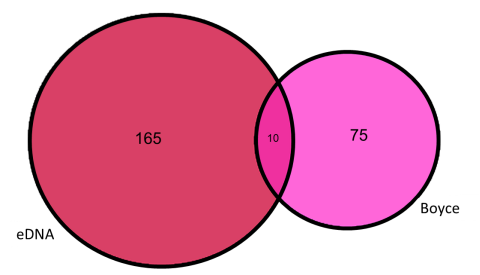


Figure 9. A comparison of two invertebrate sampling methods. Boyce's survey was conducted in 2011 using kick-net sampling. The eDNA survey was conducted in 2021 using molecular techniques. The results implied eDNA was a far more sensitive survey method, although further assessment is required to understand why there is not more of an overlap between species detected.

Despite the combination of methods, Graham and Hammond's survey recorded the least number of species (Figure 10). However, as the survey was conducted in summer and was in the same area, although not the same ditches as this study, no statistical analysis could be made. In total, 28 aquatic macro-invertebrate taxa were identified at Fair Orchard Farm.

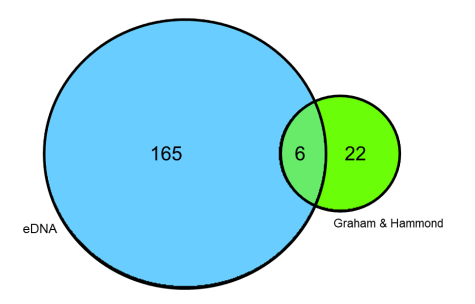


Figure 10. A comparison of Graham and Hammond's morphological identification using kick net and bottle sampling and a survey using eDNA. Graham & Hammond (2022) were able to identify 28 species of invertebrates, with six of those also being detected in the eDNA survey. However, eDNA alone was able to detect 165 species in the reens in St Brides.

Despite all morphological surveys occurring within St. Brides SSSI, different ditches and reens were studied than in this survey. Furthermore, seasonality and site will have influenced the results, so a manual comparison, rather than a statistical analysis, was conducted regarding the morphological survey results and the results from this study.

# **3.4 Ecological Quality Indices**

In total, 31 different families, using a 99% BLAST identity filtering threshold belonging to the Orders: Ephemeroptera, Plecoptera, Trichoptera, Odonata, Hemiptera, Coleoptera, Diptera and Hirudinida were identified and used in the BMWP calculations (Table 2). The field ditches FD2, and FD3 had 'moderate' water quality according to the quality interpretation, while FD1 and IDB35 were both 'good' quality. The rest of the field ditches and reens were all awarded 'very good' quality, with IDB16 having the highest ASPT score, with 8. While FD2 and FD3

had the lowest ASPT scores, with 4.6 each, FD1 had the lowest BMWP score; however, three

ditches surrounding the same field had the lowest BMWP and ASPT scores.

**Table 2.** The results of the water quality analysis according to the WFD guidelines using the Orders sequenced from the eDNA analysis.

Site	BMWP score	ASPT score	Quality interpretation
FD1	20	5	Good
FD2	23	4.6	Moderate
FD3	28	4.6	Moderate
FD4	74	6.1	Very good
FD5	29	7.3	Very good
EA9	101	6.3	Very good
IDB15	76	6.9	Very good
IDB16	72	8	Very good
IDB30	100	6.6	Very good
IDB35	89	5.9	Good
IDB37	84	6	Very good

# 4. Discussion

Following the results of this study, it is apparent that analysing eDNA in reens and ditches can provide valuable insights into invertebrate biodiversity. There were clear differences in the factors affecting diversity, with  $\beta$  diversity significantly impacted by the amount of water sampled and temperature. Furthermore,  $\alpha$  diversity was influenced by salinity and waterbody type. The difference in impact by waterbody type can be seen in the ASVs detected. There was notable turnover in the assigned species to those environments, suggesting different community compositions between the ditches and reens. Furthermore, the species associated with the ASVs could be used to calculate the B.M.W.P. and A.S.P.T., suggesting that the water quality of St. Brides is at least "moderate", with reens having better quality and higher  $\alpha$  diversity.

### 4.1 Water Quality Effect on Biodiversity

Using the invertebrate metabarcoding data to assess water quality has indicated that the field ditches were poorer in quality overall when compared to reens. Three of the five ditches in the same area had the lowest BMWP and ASPT scores ( $\leq$  5). Furthermore, no families with a score of 10 were present in ditches FD1, FD2 or FD3, indicating some pollution. The most common family detected in both ditches and reens was Chrysomelidae, in the Order Coleoptera, which scored 5, followed by Asellidae, which scored three. Two sites detected the *Hydrophilus piceus*, the Great Silver Water Beetle, which is classed as Near Threatened by the IUCN (NBN Atlas, 2021). It has a restricted British distribution, whereas, in Wales, modern records only originate from three 10km squares around the Gwent Levels. Also detected in the Coleoptera family is the declining Nationally Rare species *Choragus sheppardi,* which was detected in one of the field ditches (FD2) in this survey.

Ditches displayed poorer water quality using the Ecological Quality Indices, potentially due to a lack of management. The reens are maintained by NRW, who dredge the water on a sevenyear cycle to prevent overgrowth (Boyce, 2012), compared to ditches, whose management is more sporadic and dependent on landowners, which often results in neglect. Reen and ditch management have been linked to species diversity. Removing sediments and vegetation is required to allow for drainage, which can significantly impact vegetation and invertebrate fauna (Shaw *et al.*, 2015; Milsom *et al.*, 2004; Twisk, Noordervliet and ter Keurs, 2000). In this survey and previous surveys on the Gwent Levels, duckweed species have been allowed to grow undisturbed in ditches indicating eutrophication (Graham and Hammond, 2022; Whitehead, 2022; Boyce, 2012).

Furthermore, dense mats of duckweed suppress the growth of submerged beds of aquatic macrophytes, which are essential niches for aquatic invertebrates (Boyce, 2012). The eDNA survey conducted in this paper did not determine the cause of dominant duckweed; however, studies have shown that high levels of nitrogen and phosphorus from over-fertilization have caused a vegetation shift (Janse and Van Puijenbroek, 1998). In ditches, the plants have moved from mainly submerged aquatic vegetation to eutrophication levels of duckweed (Janse and Van Puijenbroek, 1998). Management, especially of connected ditches, may allow for faster recolonization, while other processes, such as preventing the use of agro-chemicals close to banks, further benefit macroinvertebrates (Shaw *et al.*, 2015; Leng, Musters and de Snoo, 2009; Manhoudt, Visser and de Snoo, 2007).

# 4.2 Abiotic Factors Influencing Biodiversity

#### 4.2.1 Water Filtration Quantity

The amount of water sampled was a significant variable in this study regarding the  $\beta$  diversity, although it was not significant regarding  $\alpha$  diversity in this study. The least amount of water filtered was obtained from the ditches with an average of 203ml, compared to 350ml for reens. Mächler *et al.* (2016) studied water filtration effects on the detection rate for three macroinvertebrate species belonging to the order Mollusca, Ephemeroptera and Amphipoda using volumes ranging from 250 to 2000ml. They concluded that an increase in the volume of extracted DNA screened and primer performance was more important in reducing the false negative detections of some species, although increasing sample volume was also beneficial. Only one species used had a positive relationship between increased sample volume and detection (Mächler *et al.*, 2016).

Similarly, Peixoto *et al.* (2021) investigated the importance of eDNA capture methods' regarding detecting species in aquatic environments in Portugal. The study covered the usual range of applications in eDNA monitoring, considering both targeted detection of ubiquitous species and the overall characterization of amphibian community composition using qPCR,

High Throughput Sequencing and two PCR replication thresholds (stringent and relaxed). They found that the filtration method influenced eDNA recovery and species detection, with filtering methods being more effective than precipitation, implying this was associated with the amount of water filtered (Peixoto *et al.*, 2021). Water filtering quantities are among the first biases experienced when using water eDNA methods (Mächler *et al.*, 2016). A lack of uniform methodology adds to the issue of filtration amounts, while there is little evidence on the optimal amount of extracted eDNA to reduce the likelihood of detecting false negatives (Mächler *et al.*, 2016; Deiner and Altermatt, 2014).

### 4.2.3 Temperature

In this study, temperature (p = 0.05) significantly impacted  $\beta$  diversity, although it did not affect  $\alpha$  diversity. The ditches had an average temperature of 6.55°C, while the reens had an average temperature of 6.03°C. The temperature of lotic environments has been considered important in influencing the life histories of aquatic organisms (Elliott, 1987a, 1987b; Vannote and Sweeney, 1980; Brittain, 1975). Aquatic invertebrate thermal histories cause responses at the organismic, population and community levels of organization, establishing ecological and evolutionary time scales (Ward and Stanford, 1982). Moon (1940) explored the impact of temperature when researching the movements of freshwater invertebrates in lake Windermere and found it could influence the movement of invertebrates. Furthermore, they found that while low temperatures (5°C) limited the amount of movement, they did not completely inhibit invertebrate activity (Moon, 1940). Further research is required to explore temperature impact on seasonal changes and its influence on invertebrate diversity in ditches and reens.

Strickler *et al.* (2015), who studied the effects of UV-B, temperature, and pH on eDNA degradation in aquatic microcosms, found DNA degradation rates were lowest under cold temperatures (5°C), low UV-B levels, and alkaline conditions. Higher degradation rates were associated with higher temperatures, neutral pH, and moderately high UV-B, creating

favourable microbial growth environments. Moreover, they found that temperature positively affected degradation rates in its own right, increasing as the temperature increased (Strickler, Fremier and Goldberg, 2015). However, in a natural system such as that studied here, with a temperature range of ( $4.2 \ ^\circ$ C – 7.9°C), results were robust and showed no evidence of eDNA degradation impacting diversity.

Global climate change may further impact invertebrate biodiversity. Although this was not considered in this study, there is scope for long-term monitoring of the Gwent Levels to understand how seasonal temperature changes will impact diversity. Species react differently to changes in the environment. While some invertebrates may become locally extinct, others may expand their ranges (Davey et al., 2013; Parmesan and Yohe, 2003). Flying insects have seen a dramatic decrease of 75% in biomass over a three-decade period in Germany (Hallmann et al., 2017). Furthermore, there are increasing reports of other Orders experiencing declines, with a potential consequence of changing ecosystem connectivity and function (Hallmann et al., 2019; Powney et al., 2019; van Strien et al., 2019; Dirzo et al., 2014; Schuch, Wesche and Schaefer, 2012; Shortall et al., 2009; Gaston and Fuller, 2007; Conrad et al., 2006). This could be due to streams and rivers being among the most sensitive ecosystems regarding climate change (Durance and Ormerod, 2009). Their temperatures closely track air temperature, particularly in headwaters, with streams warming in response to climate change (Durance and Ormerod, 2007; Caissie, 2006). Over the last few decades, 50 southern English streams have increased by a mean of 2.1-2.9°C in winter and 1.1-1.5°C in summer (Durance and Ormerod, 2009).

## 4.2.4 pH

The pH of water is influenced by the natural conditions of the environment (Baaloudj *et al.*, 2020). In this study, pH was found not to impact significantly  $\alpha$  and  $\beta$ -diversity for invertebrates, with the ditch pH ranging between 6.6 and 7.8 compared to reens, which ranged from 7 to 8.4.

Berezina (2001) found that the highest species diversity for freshwater invertebrates occurred between pH 4.09-8.65, with a decrease in species diversity below 4 and above 9 under experimental conditions. Petrin, Laudon and Malmqvist (2007) investigated macroinvertebrate response to a low pH gradient in rivers across Sweden. They found that Plecoptera richness did not change with varying pH levels in the north or south Sweden. However, while Ephemeroptera richness was susceptible to changes in pH in both regions, Trichoptera decreased with the increasing pH in the north, and in southern Sweden, they increased with increasing pH. The results support the hypothesis that stream invertebrates can tolerate low pH through exaptation or adaptation; however, the degree of which depends on the taxa (Petrin, Laudon and Malmqvist, 2007). Berezina (2001) further supported this, demonstrating that invertebrate survival depends on the tolerance limits of individuals, and their communities are formed depending on relationships between the pH of the aquatic environment and individual adaptability. This further implies that the pH range within this study was not enough to cause a significant impact on  $\alpha$  and  $\beta$ -diversity in freshwater invertebrates.

pH significantly impacted eDNA capture and degradation in a controlled laboratory microcosm experiment (Kagzi *et al.*, 2022). While some research found that eDNA persists for several weeks in water, other studies have demonstrated that even small (<100 bp) eDNA fragments can degrade to undetectable levels within hours or days after an organism's removal (Thomsen *et al.*, 2012; Kagzi *et al.*, 2022; Strickler, Fremier and Goldberg, 2015). Several studies have indicated that more acidic conditions (pH<5) have a significant impact on degradation in both the field and the laboratory (Goldberg, Strickler and Fremier, 2018; Seymour *et al.*, 2018; Strickler, Fremier and Goldberg, 2015). However, evidence has also suggested that pH only significantly influenced the degradation rate when combined with other abiotic variables, such as temperature and UV-B (Lance *et al.*, 2017; Strickler, Fremier and Goldberg, 2015).

### 4.2.5 Salinity

While salinity was not found to impact  $\beta$  diversity (p=0.28) significantly, it was a significant variable for  $\alpha$  diversity (p=0.05). The Water Framework Directive requires monitoring salinity to determine the waterbody's ecological condition (Pickwell *et al.*, 2022; European Commission, 2000). Salinization is a globally important stressor in freshwater ecosystems and is defined as the total concentration of dissolved inorganic ions in water or soil (Cañedo-Argüelles *et al.*, 2013; Williams and Sherwood, 1994). Natural salinization occurs from salts transported by seawater evaporation, catchment weather and sea spray (Williams and Sherwood, 1994). Salinisation can also be impacted by anthropogenic effects through water harvesting, road de-icing, mining activities and changes to vegetation leading to water table movement (Cañedo-Argüelles *et al.*, 2013).

Bray *et al.* (2018) investigated how macroinvertebrate communities from high-salinity sites impacted low-salinity sites along a salinity gradient in experimentally manipulated streams. At the community level, it was expected that salinity and biological interactions between sensitive and tolerant organisms would influence the community composition, reduce diversity and potentially homogenize communities (Bray *et al.*, 2018). This declining pattern with salinity was seen by Ephemeroptera abundance, regarding both the salinity effects and in response to interactions with more tolerant taxa. Salt-sensitivity appeared to increase in Plecoptera and EPT taxa in the presence of tolerant taxa, with little salinity impact in salt-sensitive treatments (Bray *et al.*, 2018). Overall, Bray *et al.* (2018) found that at higher salinities, direct effects of salinity dominated community responses, and this resulted in reduced abundance and altered community composition, with almost a complete loss of Ephemeroptera and a reduction in Trichoptera.

In this study, when exploring the  $\alpha$  diversity, ditches declined in species count regarding salinity levels, thus supporting Bray *et al.*'s (2018) findings. However, reens presented a positive correlation with increasing salinity levels, implying that the taxa present in the reens are more tolerable to salinization stressors. One explanation for the increase in reen diversity

is whether the communities consisted of salt-tolerant taxa. Bray *et al.* (2018) found that interspecific interactions between salt tolerance and salt-sensitive taxa became more critical as the sensitivity to the toxicant increased. This may be due to more salt-tolerant predators; therefore, there are greater predator impacts with increased salinity (Bray *et al.*, 2018; Kefford, 1998).

## 4.2.6 Dissolved Oxygen

Dissolved oxygen (DO) concentration is a key factor in determining macroinvertebrate assemblage composition, despite being non-significant in this survey (Williams, 1996). Macroinvertebrates in ditches face both predictable and unpredictable variations in DO concentration, with low oxygen concentration viewed as an important environmental filter (Verdonschot and Verdonschot, 2014). Several factors could influence the DO concentrations, including salinity, time of day, algal bloom and depth (Natural Resources Wales, 2022). Previous research in laboratory experiments has shown increased mortality, decreased growth rate, and prolonged development time under oxygen stress (Kolar and Rahel, 1993; Moore and Burn, 1968).

Dissolved oxygen levels can be impacted by temperature, and cold water can hold more dissolved oxygen than warm water, leading to frequent higher DO levels in winter (U.S. Geological Survey, 2022). The DO ranges recorded in this study are between 2.1 mg/L and 9.8 mg/L. Hypoxia occurs when DO concentrations fall below 2-3 mg/L (US Environmental Protection Agency, 2015). Therefore, this suggests that DO is not a significant variable for  $\alpha$  and  $\beta$  diversity as the DO levels in most ditches and reens were acceptable for invertebrates. Two sites (IDB16.2 and IDB30.1) had a DO concentration <3mg/L, while the other sites in the same reens had concentrations >3 mg/L, therefore the site did not impact the overall significance of  $\alpha$  and  $\beta$  diversity.

#### 4.2.7 Turbidity

There is little research to suggest turbidity directly impacts freshwater invertebrate communities, which was also found not to have a significant influence within this study. Kefford *et al.* (2007) investigated freshwater invertebrates' response to the gradient of salinity and turbidity in Australia. They found one test species, *Micronecta annae* (Hemiptera: Corixidae), preferred relatively high turbidity (>200 NTU), but only from one of two locations. However, another species, *Austrochiltonia subtenuis* (Amphipoda: Hyalellidae), showed the opposite, responding to low turbidity (<200 NTU). The evidence found was weak, potentially because turbidity levels are not directly harmful to invertebrates, which corroborates what we found in this study (Kefford *et al.*, 2007).

### 4.2.8 Depth

While this study did not find depth to be a significant factor, evidence suggests that water depth is more likely to influence invertebrate community structure throughout the seasons. In winter, water levels in the ditches and reens in St. Brides SSSI are lower than in summer, allowing floodplains to be drained. This could reduce diversity in winter, as aquatic invertebrates may not be able to travel as freely to different reens due to movement barriers, such as sluice gates separating the waterbodies. Although this has not been explored with invertebrates, Katano *et al.* (2003) found this true for fish. Furthermore, research has shown that water depth and shade can have a small impact on invertebrate communities in ditches (Shaw *et al.*, 2015). While in this study, ditches had a positive correlation with depth regarding  $\alpha$  diversity, and reens had a non-significant correlation regarding  $\alpha$  diversity and a slightly negative non-significant correlation relating to  $\beta$  diversity. However, Shaw *et al.* (2015) found increased biodiversity within invertebrate communities in deeper sites. They suggested that shade and water depth were important environmental factors but admitted that their study's significance was minor (Shaw *et al.*, 2015).

#### 4.2.9 Waterbody Type & Land use

There was no significant difference in waterbody type on  $\beta$  diversity; however, there was on  $\alpha$ (p = 0.01) diversity implying no difference in the species composition within the community was observed. However, it suggests a greater variety of species between the ditches and reens. One potential explanation is that more plants are in the ditches due to a lack of maintenance. Clare and Edwards (1983) found that most macroinvertebrates collected from ditch samples were within the water column and on plants rather than in the benthos. Furthermore, Herzon and Helenius (2008) found that terrestrial vegetation on the banks of the ditches was essential for many invertebrate communities, with a specific host and nectar plants providing food and overwintering sites. The ditches in this study had overhanging plants, providing food and shelter and may significantly impact the variety of species that can inhabit that waterbody type. Freshwater ecosystems are directly influenced and or/ indirectly by human activities, so it was essential to understand if there were any anthropogenic impacts in this study (Juvigny-Khenafou et al., 2021). Reens IDB37, IDB35, and half of IDB30 were categorized as urban, while the other reens and ditches were rural. There was no significant difference in biodiversity when comparing invertebrate  $\alpha$  and  $\beta$  diversity between ditches and reens in urban and rural environments, potentially due to minimal urban sites where samples were taken compared to rural. However, other research on lotic ecosystems has found that urbanisation negatively affects the diversity of freshwater macroinvertebrates (Gál et al., 2019). Conversely, Vermonden et al. (2009) found that urban drainage ditches can obtain the same levels of macroinvertebrate biodiversity as those in rural areas. Further research is needed to understand if the linkage between reens means land use has minimal impact on invertebrate biodiversity.

#### 4.2.10 Seasonality

The study presented here was conducted in December, which could impact species detection, especially in groups such as Ephemeroptera, where a single generation overwinters in the nymphal stage (Clifford, 1982). Musters et al. (2019) used a small, well-connected network of drainage ditches to measure the spatiotemporal  $\beta$  diversity of a freshwater macrofaunal metacommunity in the temporal climate zone in the Netherlands from May to November 2011 and 2012. They found no temporal patterns between the years and months (Musters et al., 2019). Conversely, Zizka, Geiger and Leese (2020) used DNA metabarcoding of stream invertebrates in Western Germany to explore spatial-temporal variation, where variation had distinct seasonal effects on their OTU composition at the near-natural river in spring (April/May 2017/2018) and autumn/ winter (September 2016/2017 and November/December 2016/2017). They found variations in taxa response, with taxa that are highly abundant in spring, almost absent, or present only in eggs or overwintering in autumn, and vice versa (Zizka, Geiger and Leese, 2020). Aggregated at the order level, a higher number of reads were assigned to Ephemeroptera in spring than autumn, compared to Plecoptera had a higher number of reads assigned in autumn. However, ecological status assessment remained consistent throughout the seasons and comparable to other assessments where morphological identification was used (Zizka, Geiger and Leese, 2020).

Conversely, Rehinholdt Jenson *et al.* (2021) used eDNA metabarcoding to explore seasonal turnover in the community composition of stream macroinvertebrates in Denmark and found that Plecoptera were most abundant in spring sampling as they are more abundant during the spring season. Šporka *et al.* (2006) explored streams in central Europe to understand the influence of seasonal variation on the bioassessment of streams using macroinvertebrates. They found an increase in Trichoptera and Ephemeroptera abundance when comparing October sampling to spring sampling, a trend not seen in either of the studies by Rehinholdt Jenson *et al.* (2021) or Zizka, Geiger and Leese (2020) (Šporka *et al.*, 2006). They presented

significant differences in communities in spring and autumn, with a considerable increase in taxa detected when sampling in both seasons (Reinholdt Jensen *et al.*, 2021).

### 4.3 eDNA as a Tool for Assessing Water Quality

Using eDNA as an identification tool can potentially reduce processing times, labour intensity, and time spent on sample analysis (Kuntke *et al.*, 2020). Environmental DNA-derived data allows for the inclusion of a much more comprehensive range of taxa and indicator groups that would otherwise be ignored due to the limitations of traditional taxonomic identification (Seymour *et al.*, 2020). Elbrecht *et al.* (2017) compared DNA metabarcoding with a morphology-based protocol and found that eDNA identified more than twice the number of taxa. Traditional morpho-taxonomic methods and eDNA face limitations regarding the quality and completeness of sequence databases or identification keys used for taxonomic assignment (Seymour *et al.*, 2020). With traditional methods such as kick-net sampling, taxonomic assignment is often limited to family or general-level assignments and can often oversimplify or omit groups such as Rotifers, Oligochaeta, or Chironomidae, which are critical environmental indicators (Seymour *et al.*, 2020; Elbrecht *et al.*, 2017; Furse *et al.*, 2009).

When comparing the eDNA survey results from this study with previous surveys of St. Brides, we see an increase in species detection. This is due to one of the benefits of eDNA being that it can detect both aquatic and terrestrial macroinvertebrates. The previous surveys on the Gwent Levels have been conducted using traditional morphological-taxonomic methods and, therefore, due to time or expertise constraints, have either focused on specifically terrestrial or aquatic macroinvertebrates (Graham and Hammond, 2022; Boyce, 2012)

While some of the reens in this study were used in Boyce's survey, they are incomparable due to seasonality. Boyce and Graham, and Hammond conducted the traditional surveys in summer, when invertebrates are more active, compared to the eDNA survey, which was conducted in winter. Temperature both directly and indirectly influences aquatic invertebrate

life histories, with colder temperatures reducing the amount of energy available for growth and reproduction (Hart, 1985). However, despite the eDNA sampling being conducted in winter, the species detection was far greater. Further investigation is required to understand how seasonality effects invertebrate detection in water to understand whether surveys can be conducted throughout the year.

### 5. Conclusion

Monitoring invertebrates is essential for understanding how an aquatic community is functioning as often declines in biodiversity go unnoticed as they are not always visible. As invertebrates are sensitive to ecological changes, they are useful biological indicators; however, many previous invertebrate surveys in the Gwent Levels have used morphological identification methods. This study analysing eDNA has provided a more comprehensive taxonomic database for invertebrates in ditches and reens without the results being impacted by environmental degradation from elements such as pH and temperature. However, some abiotic variables significantly impacted the invertebrate community, such as waterbody type and salinity, indicating an interaction between the two variables and is vital in understanding the  $\alpha$  diversity of communities in these wetland channels, potentially influenced by marine inputs. Regarding  $\beta$  diversity, the amount of water filtered has a significant impact meaning there is further scope to develop more consistent filtering methods. Furthermore, temperature also impacts  $\beta$  diversity, which demonstrates there could be an interaction between the amount of water filtered and temperature; however, this is unlikely.

Further research is required to analyse more eDNA samples from ditches and explore seasonality as a variable. Furthermore, exploration is required to understand if there is any correlation between salinity and waterbody type and the impact it has on  $\alpha$  diversity. Nevertheless, using aqueous eDNA analysis has proven effective in understanding biodiversity in St. Brides SSSI, with the results being applicable to aid future management plans in the protection of invertebrates.

## 6. References

Al-Shami, S.A., Che Salmah, Md Rawi, Abu Hassan, A. and Madziatul, R.M. (2013) Biodiversity of stream insects in the Malaysian Peninsula: spatial patterns and environmental constraints. *Ecological Entomology*. [online]. 38 (3), pp.238–249.

Altschul, S.F., Gish, W., Miller, W., Myers, E.W. and Lipman, D.J. (1990) Basic local alignment search tool. *Journal of Molecular Biology*. [online]. 215 (3), pp.403–410.

Aquilina, R. (2013) *Pre-restoration assessment of the Hogsmill and River Wandle* Wandle River Trust, 21.

Baaloudj, A., Ouarab, S., Kerfour, A., Bouriach, M., Ali Hussein, A., Hammana, C. and N' diaye Djénéba, H. (2020) Use of macro invertebrates to assess the quality of Seybouse River (North-East of Algeria). *Ukrainian Journal of Ecology*. 10 (4), pp.60–66.

Beever, E.A. (2006) MONITORING BIOLOGICAL DIVERSITY: STRATEGIES, TOOLS, LIMITATIONS, AND CHALLENGES. *Northwestern Naturalist*. [online]. 87 (1), Society for Northwestern Vertebrate Biology, pp.66–79.

Beltman, B. (1984) Management of ditches. The effect of cleaning of ditches on the water coenoses. *SIL Proceedings, 1922-2010.* [online]. 22 (3), Taylor & Francis, pp.2022–2028.

Berezina, N.A. (2001) Influence of Ambient pH on Freshwater Invertebrates under Experimental Conditions. *Russian Journal of Ecology*. [online]. 32 (5), pp.343–351.

Boyce, D.C. (2012) *Monitoring invertebrate features on Sites of Special Scientific Interest: aquatic invertebrates on the Gwent Levels: Redwick and Llandevenny SSSI, St Brides SSSI.* Cardiff, Countryside Council for Wales, 87.

Bray, J.P., Reich, J., Nichols, S.J., Kon Kam King, G., Mac Nally, R., Thompson, R., O'Reilly-Nugent, A. and Kefford, B.J. (2018) Biological interactions mediate context and species-specific sensitivities to salinity. *Philosophical Transactions of the Royal Society B: Biological Sciences*. [online]. 374 (1764), Royal Society, p.20180020.

Brennan, G.L. *et al.* (2019) Temperate airborne grass pollen defined by spatio-temporal shifts in community composition. *Nature Ecology & Evolution*. [online]. 3 (5), Nature Publishing Group, pp.750–754.

Briski, E., Cristescu, M.E., Bailey, S.A. and MacIsaac, H.J. (2011) Use of DNA barcoding to detect invertebrate invasive species from diapausing eggs. *Biological Invasions*. [online]. 13 (6), pp.1325–1340.

Brittain, J.E. (1975) The Life Cycle of Baëtis macani Kimmins (Ephemerida) in a Norwegian Mountain Biotope. *Insect Systematics & Evolution*. [online]. 6 (1), Brill, pp.47–51.

Brookes, A. (1986) Response of aquatic vegetation to sedimentation downstream from river channelisation works in England and Wales. *Biological Conservation*. [online]. 38 (4), pp.351–367.

Brookes, A. (1985) traditional engineering methods, physical consequences and alternative practices. *Progress in Physical Geography: Earth and Environment*. [online]. 9 (1), SAGE Publications Ltd, pp.44–73.

Brua, Robert.B., Culp, J.M. and Benoy, G.A. (2011) Comparison of benthic macroinvertebrate communities by two methods: Kick- and U-net sampling | SpringerLink. *Hydrobiologia*. 658, pp.293–302.

Caissie, D. (2006) The thermal regime of rivers: a review. *Freshwater Biology*. [online]. 51 (8), pp.1389–1406.

Callahan, B.J., McMurdie, P.J., Rosen, M.J., Han, A.W., Johnson, A.J.A. and Holmes, S.P. (2016) "DADA2: High-resolution sample inference from Illumina amplicon data." *Nature Methods*. [online]. 13, pp.581–583.

Cañedo-Argüelles, M., Kefford, B.J., Piscart, C., Prat, N., Schäfer, R.B. and Schulz, C.-J. (2013) Salinisation of rivers: An urgent ecological issue. *Environmental Pollution*. [online]. 173, pp.157–167.

Carter, J.L. and Resh, V.H. (2001) After site selection and before data analysis: sampling, sorting, and laboratory procedures used in stream benthic macroinvertebrate monitoring programs by USA state agencies. *Journal of the North American Benthological Society*. [online]. 20 (4), [The University of Chicago Press, Society for Freshwater Science], pp.658–682.

Chariton, A.A., Court, L.N., Hartley, D.M., Colloff, M.J. and Hardy, C.M. (2010) Ecological assessment of estuarine sediments by pyrosequencing eukaryotic ribosomal DNA. *Frontiers in Ecology and the Environment*. [online]. 8 (5), pp.233–238.

Clare, P. and Edwards, R.W. (1983) The macroinvertebrate fauna of the drainage channels of the Gwent Levels, South Wales. *Freshwater Biology*. [online]. 13 (3), pp.205–225.

Clarke, S.J. (2015) Conserving freshwater biodiversity: The value, status and management of high quality ditch systems. *Journal for Nature Conservation*. [online]. 24, pp.93–100.

Clifford, Hugh.F. (1982) Life Cycles of Mayflies (Ephemeroptera), with Special Reference to Voltinism. *Quaestiones Entomologicae*. 18, pp.15–90.

Conrad, K.F., Warren, M.S., Fox, R., Parsons, M.S. and Woiwod, I.P. (2006) Rapid declines of common, widespread British moths provide evidence of an insect biodiversity crisis. *Biological Conservation*. [online]. 132 (3), pp.279–291.

Countryside Council Wales (1991) *Guidelines for selection of SSSIs | JNCC - Adviser to Government on Nature Conservation* [online]. Available from: https://naturalresources.wales/media/640899/SSSI\_0341\_Citation\_EN0014d9a.pdf [Accessed 28 September 2021].

Countryside Council Wales (2008) *Gwent Levels: St.Brides Site of Special Sceintific Interest* Cardiff.

Creer, S. *et al.* (2010) Ultrasequencing of the meiofaunal biosphere: practice, pitfalls and promises. *Molecular Ecology*. [online]. 19 (s1), pp.4–20.

Darling, J. and Tepolt, C. (2008) Highly sensitive detection of invasive shore crab (Carcinus maenas and Carcinus aestuarii) larvae in mixed plankton samples using polymerase chain reaction and restriction fragment length polymorphisms (PCR-RFLP). *Aquatic Invasions*. [online]. 3 (2), pp.141–152.

Davey, C.M., Devictor, V., Jonzén, N., Lindström, Å. and Smith, H.G. (2013) Impact of climate change on communities: revealing species' contribution. *Journal of Animal Ecology*. [online]. 82 (3), pp.551–561.

Deiner, K. and Altermatt, F. (2014) Transport Distance of Invertebrate Environmental DNA in a Natural River. *PLoS ONE*. [online]. 9 (2), p.e88786.

Dirzo, R., Young, H.S., Galetti, M., Ceballos, G., Isaac, N.J.B. and Collen, B. (2014) Defaunation in the Anthropocene. *Science*. [online]. 345 (6195), pp.401–406.

Durance, I. and Ormerod, S.J. (2007) Climate change effects on upland stream macroinvertebrates over a 25-year period. *Global Change Biology*. [online]. 13 (5), Blackwell Publishing Ltd., 9600 Garsington Road, pp.942–957.

Durance, I. and Ormerod, S.J. (2009) Trends in water quality and discharge confound long-term warming effects on river macroinvertebrates. *Freshwater Biology*. [online]. 54 (2), pp.388–405.

Egan, S.P., Barnes, M.A., Hwang, C.-T., Mahon, A.R., Feder, J.L., Ruggiero, S.T., Tanner, C.E. and Lodge, D.M. (2013) Rapid Invasive Species Detection by Combining Environmental DNA with Light Transmission Spectroscopy. *Conservation Letters*. [online]. 6 (6), pp.402–409.

Elbrecht, V. and Leese, F. (2017) PrimerMiner: an r package for development and in silico validation of DNA metabarcoding primers. *Methods in Ecology and Evolution*. [online]. 8 (5), pp.622–626.

Elbrecht, V., Vamos, E.E., Meissner, K., Aroviita, J. and Leese, F. (2017) Assessing strengths and weaknesses of DNA metabarcoding-based macroinvertebrate identification for routine stream monitoring. *Methods in Ecology and Evolution*. [online]. 8 (10), pp.1265–1275.

Elliott, J.M. (1987a) Egg Hatching and Resource Partitioning in Stoneflies: The Six British Leuctra Spp. (Plecoptera: Leuctridae). *Journal of Animal Ecology*. [online]. 56 (2), [Wiley, British Ecological Society], pp.415–426.

Elliott, J.M. (1987b) Temperature-induced changes in the life cycle of Leuctra nigra(Plecoptera: Leuctridae) from a Lake District stream. *Freshwater Biology*. 18 (1), pp.177–184.

European Commission (2000) Directive 2000/60/EC of the European Parliament and of the Council of 23 October 2000 establishing a framework for community action in the field of water policy. *Official Journal of the European Communities*. L327, pp.1–72.

European Commission (2021) Introduction to the EU Water Framework Directive -Environment - European Commission2021 [online]. Available from: https://ec.europa.eu/environment/water/water-framework/info/intro\_en.htm [Accessed 10 January 2022].

Fernández, S., Rodríguez-Martínez, S., Martínez, J.L., Garcia-Vazquez, E. and Ardura, A. (2019) How can eDNA contribute in riverine macroinvertebrate assessment? A metabarcoding approach in the Nalón River (Asturias, Northern Spain). *Environmental DNA*. [online]. 1 (4), pp.385–401.

Ficetola, G.F., Miaud, C., Pompanon, F. and Taberlet, P. (2008) Species detection using environmental DNA from water samples. *Biology Letters*. [online]. 4 (4), Royal Society, pp.423–425.

Furse, M.T., Hering, D., Brabec, K., Buffagni, A., Sandin, L. and Verdonschot, P.F.M. (2009) *The Ecological Status of European Rivers: Evaluation and Intercalibration of Assessment Methods* Springer Science & Business Media.

Gál, B., Szivák, I., Heino, J. and Schmera, D. (2019) The effect of urbanization on freshwater macroinvertebrates – Knowledge gaps and future research directions. *Ecological Indicators*. [online]. 104, pp.357–364.

Gaston, K.J. and Fuller, R.A. (2007) Biodiversity and extinction: losing the common and the widespread. *Progress in Physical Geography: Earth and Environment*. [online]. 31 (2), SAGE Publications Ltd, pp.213–225.

Ghani, W.M.H.W.A., Rawi, C.S.M., Hamid, S.A. and Al-Shami, S.A. (2016) Efficiency of Different Sampling Tools for Aquatic Macroinvertebrate Collections in Malaysian Streams. *Tropical Life Sciences Research*. 27 (1), pp.115–133.

Goldberg, C.S., Strickler, K.M. and Fremier, A.K. (2018) Degradation and dispersion limit environmental DNA detection of rare amphibians in wetlands: Increasing efficacy of sampling designs. *Science of The Total Environment*. [online]. 633, pp.695–703.

Graham, J. and Hammond, M. (2022) *Investigating ditch biodiversity in the Gwent Levels : a survey of vegetation and aquatic macro-invertebrates at 5 sites within the Gwent Levels* 113.

Hadley, W. (2016) *ggplot2: Elegant Graphics for Data Analysis.* [online]. Springer-Verlag New York. Available from: https://ggplot2.tidyverse.org.

Hajibabaei, M., Shokralla, S., Zhou, X., Singer, G.A.C. and Baird, D.J. (2011) Environmental Barcoding: A Next-Generation Sequencing Approach for Biomonitoring Applications Using River Benthos. *PLOS ONE*. [online]. 6 (4), Public Library of Science, p.e17497.

Hallmann, C.A. *et al.* (2017) More than 75 percent decline over 27 years in total flying insect biomass in protected areas. *PLOS ONE*. [online]. 12 (10), Public Library of Science, p.e0185809.

Hallmann, C.A., Zeegers, T., van Klink, R., Vermeulen, R., van Wielink, P., Spijkers, H., van Deijk, J., van Steenis, W. and Jongejans, E. (2019) Declining abundance of beetles, moths and caddisflies in the Netherlands. *Insect Conservation and Diversity*. [online]. 13 (2), pp.127–139.

Halse, S., Cale, D., Jasinska, E. and Shiel, R. (2002) Monitoring change in aquatic invertebrate biodiversity: sample size, faunal elements and analytical methods. *Aquatic Ecology*. [online]. 36 (3), Dordrecht, Netherlands, Springer Nature B.V., pp.395–410.

Hanbo, C. (2022) *VennDiagram: Generate High-Resolution Venn and Euler Plots* [R package version 1.7.3]

Harvey, J.B.J., Hoy, M.S. and Rodriguez, R.J. (2009) Molecular detection of native and invasive marine invertebrate larvae present in ballast and open water environmental samples collected in Puget Sound. *Journal of Experimental Marine Biology and Ecology*. [online]. 369 (2), pp.93–99.

Hawkes, H.A. (1998) Origin and development of the biological monitoring working party score system. *Water Research*. [online]. 32 (3), pp.964–968.

Hellawell, J.M. (2012) *Biological Indicators of Freshwater Pollution and Environmental Management* 3rd edition. Springer Science & Business Media.

Herzon, I. and Helenius, J. (2008) Agricultural drainage ditches, their biological importance and functioning. *Biological Conservation*. [online]. 141 (5), pp.1171–1183.

Janse, J.H. and Van Puijenbroek, P.J.T.M. (1998) 'Effects of eutrophication in drainage ditches' *Environmental Pollution.Nitrogen, the Confer-N-s First International Nitrogen Conference 1998* [online]. 102 (1, Supplement 1), pp.547–552.

Juvigny-Khenafou, N.P.D., Piggott, J.J., Atkinson, D., Zhang, Y., Macaulay, S.J., Wu, N. and Matthaei, C.D. (2021) Impacts of multiple anthropogenic stressors on stream macroinvertebrate community composition and functional diversity. *Ecology and Evolution*. [online]. 11 (1), pp.133–152.

Kagzi, K., Hechler, R.M., Fussmann, G.F. and Cristescu, M.E. (2022) Environmental RNA degrades more rapidly than environmental DNA across a broad range of pH conditions. *Molecular Ecology Resources*. [online]. Available from: https://onlinelibrary.wiley.com/doi/abs/10.1111/1755-0998.13655 [Accessed 31 August 2022].

Katano, O., Hosoya, K., Iguchi, K., Yamaguchi, M., Aonuma, Y. and Kitano, S. (2003) Species diversity and abundance of freshwater fishes in irrigation ditches around rice fields. *Environmental Biology of Fishes*. [online]. 66 (2), pp.107–121.

Keck, F., Hürlemann, S., Locher, N., Stamm, C., Deiner, K. and Altermatt, F. (2022) A triad of kicknet sampling, eDNA metabarcoding, and predictive modeling to assess richness of mayflies, stoneflies and caddisflies in rivers. *Metabarcoding and Metagenomics*. [online]. 6, pp.117–131.

Kefford, B.J. (1998) The Relationship Between Electrical Conductivity and Selected Macroinvertebrate Communities in Four River Systems of South-West Victoria, Australia. *International Journal of Salt Lake Research.* 7, pp.153–170.

Kefford, B.J., Salter, J., Clay, C., Dunlop, J.E. and Nugegoda, D. (2007) Freshwater Invertebrates' Response to Gradients of Salinity and Turbidity: Using Preference as a Rapid Sub-lethal Test. *Australasian Journal of Ecotoxicology*. [online]. 13 (3), Australasian Society for Ecotoxicology, pp.131–142. Keller, E.A. (1976)'Channelization: Environmental, Geomorphic, and Engineering Aspects | 7'Geomorphology and Engineering 1st edition. [online]. Available from: https://www.taylorfrancis.com/chapters/edit/10.4324/9781003028826-7/channelizationenvironmental-geomorphic-engineering-aspects-edward-keller [Accessed 30 September 2022].

Kolar, C.S. and Rahel, F.J. (1993) Interaction of a biotic factor (predator presence) and an abiotic factor (low oxygen) as an influence on benthic invertebrate communities. *Oecologia*. [online]. 95 (2), pp.210–219.

Kuntke, F., de Jonge, N., Hesselsøe, M. and Lund Nielsen, J. (2020) Stream water quality assessment by metabarcoding of invertebrates. *Ecological Indicators*. [online]. 111, p.105982.

Lance, R., Klymus, K., Richter, C., Guan, X., Farrington, H., Carr, M., Thompson, N., Chapman, D. and Baerwaldt, K. (2017) Experimental observations on the decay of environmental DNA from bighead and silver carps. *Management of Biological Invasions*. [online]. 8 (3), pp.343–359.

Lathouri, M., England, J., Dunbar, M.J., Hannah, D.M. and Klaar, M. (2021) A river classification scheme to assess macroinvertebrate sensitivity to water abstraction pressures. *Water and Environment Journal*. [online]. 35 (4), pp.1226–1238.

Leng, X., Musters, C.J.M. and de Snoo, G.R. (2009) Restoration of plant diversity on ditch banks: Seed and site limitation in response to agri-environment schemes. *Biological Conservation*. [online]. 142 (7), pp.1340–1349.

Leray, M., Ho, S.-L., Lin, I.-J. and Machida, R.J. (2018) MIDORI server: a webserver for taxonomic assignment of unknown metazoan mitochondrial-encoded sequences using a curated database. *Bioinformatics*. [online]. 34 (21), pp.3753–3754.

Leray, M., Yang, J.Y., Meyer, C.P., Mills, S.C., Agudelo, N., Ranwez, V., Boehm, J.T. and Machida, R.J. (2013) A new versatile primer set targeting a short fragment of the mitochondrial COI region for metabarcoding metazoan diversity: application for characterizing coral reef fish gut contents. *Frontiers in Zoology*. [online]. 10 (1), p.34.

Li, F., Mahon, A.R., Barnes, M.A., Feder, J., Lodge, D.M., Hwang, C.-T., Schafer, R., Ruggiero, S.T. and Tanner, C.E. (2011) Quantitative and Rapid DNA Detection by Laser Transmission Spectroscopy. *PLOS ONE*. [online]. 6 (12), Public Library of Science, p.e29224.

Linke, S. *et al.* (2018) Freshwater ecoacoustics as a tool for continuous ecosystem monitoring. *Frontiers in Ecology and the Environment*. [online]. 16 (4), pp.231–238.

Living Levels (2021) *Reens, ditches and grips Living Levels*.2021 [online]. Available from: https://www.livinglevels.org.uk/stories/2021/3/12/reens-ditches-and-grips [Accessed 13 October 2022].

Mächler, E., Deiner, K., Spahn, F. and Altermatt, F. (2016) Fishing in the Water: Effect of Sampled Water Volume on Environmental DNA-Based Detection of Macroinvertebrates. *Environmental Science & Technology*. [online]. 50 (1), American Chemical Society, pp.305–312.

Mahon, A.R., Barnes, M.A., Li, F., Egan, S.P., Tanner, C.E., Ruggiero, S.T., Feder, J.L. and Lodge, D.M. (2013) DNA-based species detection capabilities using laser transmission

spectroscopy. *Journal of The Royal Society Interface*. [online]. 10 (78), Royal Society, p.20120637.

Mandaville, S.M. (2002) Benthic Macroinvertebrates in FreshwatersTaxa Tolerance Values, Metrics, and Protocols Soil & Water Conservation Society of Metro Halifax.

Manhoudt, A.G.E., Visser, A.J. and de Snoo, G.R. (2007) Management regimes and farming practices enhancing plant species richness on ditch banks. *Agriculture, Ecosystems & Environment*. [online]. 119 (3), pp.353–358.

Milsom, T.P., Sherwood, A.J., Rose, S.C., Town, S.J. and Runham, S.R. (2004) Dynamics and management of plant communities in ditches bordering arable fenland in eastern England. *Agriculture, Ecosystems & Environment*. [online]. 103 (1), pp.85–99.

Moon, H.P. (1940) An Investigation of the Movements of Fresh-Water Invertebrate Faunas. *Journal of Animal Ecology*. [online]. 9 (1), [Wiley, British Ecological Society], pp.76–83.

Moore, W.G. and Burn, A. (1968) Lethal Oxygen Thresholds for Certain Temporary Pond Invertebrates and Their Applicability to Field Situations. *Ecology*. [online]. 49 (2), pp.349–351.

Murton, K., Hunt, A. and Rodgers, K. (2019) *Conservation Objectives for the reen and field ditch habitat feature and aquatic plant features on the Gwent Levels SSSIs and Newport Wetlands SSSI* Cardiff, Natural Resources Wales, 92.

Musters, C.J.M., Hunting, E.R., Schrama, M., Cieraad, E., Barmentlo, S.H., Ieromina, O., Vijver, M.G. and van Bodegom, P.M. (2019) Spatial and temporal homogenisation of freshwater macrofaunal communities in ditches. *Freshwater Biology*. [online]. 64 (12), pp.2260–2268.

Natural Resources Wales (2022) 'Dissolved Oxygen in Water' *Information Note* Natural Resources Wales.

NBN Atlas (2021) *Hydrophilus piceus : Great Silver Water Beetle | NBN Atlas*2021 [online]. Available from: https://species.nbnatlas.org/species/NBNSYS0000007737 [Accessed 15 August 2022].

Oksanen, J. (2022) vegan: Community Ecology Package [R package version 2.6-2]

Paisley, M.F., Trigg, D.J. and Walley, W.J. (2014) Revision of the Biological Monitoring Working Party (bmwp) Score System: Derivation of Present-Only and Abundance-Related Scores from Field Data. *River Research and Applications*. [online]. 30 (7), pp.887–904.

Parmesan, C. and Yohe, G. (2003) A globally coherent fingerprint of climate change impacts across natural systems. *Nature*. [online]. 421 (6918), pp.37–42.

Pawlowski, J. *et al.* (2018) The future of biotic indices in the ecogenomic era: Integrating (e)DNA metabarcoding in biological assessment of aquatic ecosystems. *Science of The Total Environment.* [online]. 637–638, pp.1295–1310.

Peixoto, S., Chaves, C., Velo-Antón, G., Beja, P. and Egeter, B. (2021) Species detection from aquatic eDNA: Assessing the importance of capture methods. *Environmental DNA*. [online]. 3 (2), pp.435–448.

Petrin, Z., Laudon, H. and Malmqvist, B. (2007) Does freshwater macroinvertebrate diversity along a pH-gradient reflect adaptation to low pH? *Freshwater Biology*. [online]. 52 (11), pp.2172–2183.

Pickwell, A., Constable, D., Chadd, R., Extence, C. and Little, S. (2022) The development of a novel macroinvertebrate indexing tool for the determination of salinity effects in freshwater habitats. *River Research and Applications*. [online]. 38 (3), pp.522–538.

Poikane, S. *et al.* (2016) Benthic macroinvertebrates in lake ecological assessment: A review of methods, intercalibration and practical recommendations. *Science of The Total Environment.* [online]. 543, pp.123–134.

Porazinska, D.L., Giblin-Davis, R.M., Esquivel, A., Powers, Thomas.O., Sung, W. and Thomas, W.K. (2010) Ecometagenetics confirm high tropical rainforest nematode diversity. *Molecular Ecology*. 19 (24), pp.5521–5530.

Powney, G.D., Carvell, C., Edwards, M., Morris, R.K.A., Roy, H.E., Woodcock, B.A. and Isaac, N.J.B. (2019) Widespread losses of pollinating insects in Britain. *Nature Communications*. [online]. 10 (1), p.1018.

R Core Team (2022) *R: A Language and Environment for Statistical Computing.* Vienna, Austria, R Foundation for Statistical Computing.

Rees, H.C., Bishop, K., Middleditch, D.J., Patmore, J.R.M., Maddison, B.C. and Gough, K.C. (2014) The application of eDNA for monitoring of the Great Crested Newt in the UK. *Ecology and Evolution*. [online]. 4 (21), pp.4023–4032.

Reinholdt Jensen, M., Egelyng Sigsgaard, E., Agersnap, S., Jessen Rasmussen, J., Baattrup-Pedersen, A., Wiberg-Larsen, P. and Francis Thomsen, P. (2021) Seasonal turnover in community composition of stream-associated macroinvertebrates inferred from freshwater environmental DNA metabarcoding. *Environmental DNA*. [online]. 3 (4), pp.861– 876.

Revenga, C., Campbell, I., Abell, R., de Villiers, P. and Bryer, M. (2005) Prospects for monitoring freshwater ecosystems towards the 2010 targets. *Philosophical Transactions of the Royal Society B: Biological Sciences*. [online]. 360 (1454), Royal Society, pp.397–413.

Rohart, F., Gautier, B., Singh, A. and Le Cao, K.A. (2017) 'mixOmics: An R package for 'omics feature selection and multiple data integration.' *Public Library of Science.Computational Biology* 13 (11).

Schmeller, D.S. (2008) European species and habitat monitoring: where are we now? *Biodiversity and Conservation.* [online]. 17 (14), pp.3321–3326.

Schuch, S., Wesche, K. and Schaefer, M. (2012) Long-term decline in the abundance of leafhoppers and planthoppers (Auchenorrhyncha) in Central European protected dry grasslands. *Biological Conservation*. [online]. 149 (1), pp.75–83.

Sevigny, J.L. (2018) *Assign-Taxonomy-with-BLAST*Python [online]. Available from: https://github.com/Joseph7e/Assign-Taxonomy-with-BLAST/blob/944a73c8390dcbc20203217ae15097628613dfe8/taxonomy\_assignment\_BLAS T.py [Accessed 30 November 2022].

Seymour, M. *et al.* (2018) Acidity promotes degradation of multi-species environmental DNA in lotic mesocosms. *Communications Biology*. [online]. 1 (1), p.4.

Seymour, M., Edwards, F.K., Cosby, B.J., Kelly, M.G., de Bruyn, M., Carvalho, G.R. and Creer, S. (2020) Executing multi-taxa eDNA ecological assessment via traditional metrics and interactive networks. *Science of The Total Environment*. [online]. 729, p.138801.

Sharma, R.C., Arambam, R. and Sharma, R. (2009) Surveying macro-invertebrate diversity in the Tons river, Doon Valley, India. *The Environmentalist*. [online]. 29 (3), pp.241–254.

Shaw, R.F., Johnson, P.J., Macdonald, D.W. and Feber, R.E. (2015) Enhancing the Biodiversity of Ditches in Intensively Managed UK Farmland. *PLOS ONE*. [online]. 10 (10), Public Library of Science, p.e0138306.

Shortall, C.R., Moore, A., Smith, E., Hall, M.J., Woiwod, I.P. and Harrington, R. (2009) Longterm changes in the abundance of flying insects. *Insect Conservation and Diversity*. [online]. 2 (4), pp.251–260.

Sor, R., Ngor, P.B., Soum, S., Chandra, S., Hogan, Z.S. and Null, S.E. (2021) Water Quality Degradation in the Lower Mekong Basin. *Water*. [online]. 13 (11), Multidisciplinary Digital Publishing Institute, p.1555.

Šporka, F., Vlek, H.E., Bulánková, E. and Krno, I. (2006) Influence of seasonal variation on bioassessment of streams using macroinvertebrates' In: Furse, M.T., Hering, D., Brabec, K., Buffagni, A., Sandin, L. and Verdonschot, P.F.M. (eds.) *The Ecological Status of European Rivers: Evaluation and Intercalibration of Assessment Methods*. Developments in Hydrobiology [online]. Dordrecht, Springer Netherlands, 543–555. Available from: https://doi.org/10.1007/978-1-4020-5493-8\_36 [Accessed 16 August 2022].

Stein, E.D., Martinez, M.C., Stiles, S., Miller, P.E. and Zakharov, E.V. (2014) Is DNA Barcoding Actually Cheaper and Faster than Traditional Morphological Methods: Results from a Survey of Freshwater Bioassessment Efforts in the United States? *PLOS ONE*. [online]. 9 (4), Public Library of Science, p.e95525.

Stein, H., Springer, M. and Kohlmann, B. (2008) Comparison of two sampling methods for biomonitoring using aquatic macroinvertebrates in the Dos Novillos River, Costa Rica. *Ecological Engineering*. 34 (4), pp.267–275.

Strickler, K.M., Fremier, A.K. and Goldberg, C.S. (2015) 'Quantifying effects of UV-B, temperature, and pH on eDNA degradation in aquatic microcosms' *Biological Conservation.Special Issue: Environmental DNA: A powerful new tool for biological conservation* [online]. 183, pp.85–92.

van Strien, A.J., van Swaay, C.A.M., van Strien-van Liempt, W.T.F.H., Poot, M.J.M. and WallisDeVries, M.F. (2019) Over a century of data reveal more than 80% decline in butterflies in the Netherlands. *Biological Conservation*. [online]. 234, pp.116–122.

Surber, E.W. (1937) Rainbow Trout and Bottom Fauna Production in One Mile of Stream. *Transactions of the American Fisheries Society*. [online]. 66 (1), Taylor & Francis, pp.193–202.

Taberlet, P., Coissac, E., Pompanon, F., Brochmann, C. and Willerslev, E. (2012) Towards next-generation biodiversity assessment using DNA metabarcoding. *Molecular Ecology*. [online]. 21 (8), pp.2045–2050.

Thomsen, P.F., Kielgast, J., Iversen, L.L., Møller, P.R., Rasmussen, M. and Willerslev, E. (2012a) Detection of a Diverse Marine Fish Fauna Using Environmental DNA from Seawater Samples. *PLOS ONE*. [online]. 7 (8), Public Library of Science, p.e41732.

Twisk, W., Noordervliet, M.A.W. and ter Keurs, W.J. (2000) Effects of ditch management on caddisfly, dragonfly and amphibian larvae in intensively farmed peat areas. *Aquatic Ecology*. [online]. 34 (4), pp.397–411.

US Environmental Protection Agency (2015) *Hypoxia 101*Overviews and Factsheets24 March 2015 [online]. Available from: https://www.epa.gov/ms-htf/hypoxia-101 [Accessed 11 October 2022].

U.S. Geological Survey (2022) *Dissolved Oxygen and Water*2022 [online]. Available from: https://www.usgs.gov/special-topics/water-science-school/science/dissolved-oxygen-and-water [Accessed 11 October 2022].

Vannote, R.L. and Sweeney, B.W. (1980) Geographic Analysis of Thermal Equilibria: A Conceptual Model for Evaluating the Effect of Natural and Modified Thermal Regimes on Aquatic Insect Communities. *The American Naturalist*. [online]. University of Chicago Press. Available from: https://www.journals.uchicago.edu/doi/10.1086/283591 [Accessed 12 October 2022].

Verdonschot, R. (2012) Drainage ditches, biodiversity hotspots for aquatic invertebrates. Defining and assessing the ecological status of a man-made ecosystem based on macroinvertebrates. Alterra Scientic Contributions. 40.

Verdonschot, R.C.M., Keizer-vlek, H.E. and Verdonschot, P.F.M. (2011) Biodiversity value of agricultural drainage ditches: a comparative analysis of the aquatic invertebrate fauna of ditches and small lakes. *Aquatic Conservation: Marine and Freshwater Ecosystems*. [online]. 21 (7), pp.715–727.

Verdonschot, R.C.M. and Verdonschot, P.F.M. (2014) Shading effects of free-floating plants on drainage-ditch invertebrates. *Limnology*. [online]. 15 (3), pp.225–235.

Ward, J.V. and Stanford, J.A. (1982) Thermal Responses in the Evolutionary Ecology of Aquatic Insects. *Annual Review of Entomology*. [online]. 27 (1), pp.97–117.

Whitehead, P.F. (2022) TRADITIONAL ORCHARD INVERTEBRATE STUDY 2019-2021 REPORT ON THE INVERTEBRATES' *Report on the Invertebrates* 56.

Williams, D.D. (1996) Environmental Constraints in Temporary Fresh Waters and Their Consequences for the Insect Fauna. *Journal of the North American Benthological Society*. [online]. 15 (4), The University of Chicago Press, pp.634–650.

Williams, W.D. and Sherwood, J.E. (1994) Definition and measurement of salinity in salt lakes. *International Journal of Salt Lake Research*. [online]. 3 (1), pp.53–63.

Witmer, G.W. (2005) Wildlife population monitoring: some practical considerations. *Wildlife Research*. [online]. 32 (3), CSIRO PUBLISHING, pp.259–263.

Wright, J.F. (1994) Development of RIVPACS in the UK and the value of the underlying data-base. *Limnetica*. [online]. 10 (1), pp.15–32.

Zizka, V.M.A., Geiger, M.F. and Leese, F. (2020) DNA metabarcoding of stream invertebrates reveals spatio-temporal variation but consistent status class assessments in a natural and urban river. *Ecological Indicators*. [online]. 115, p.106383.

# 7. Appendices

Appendix I.

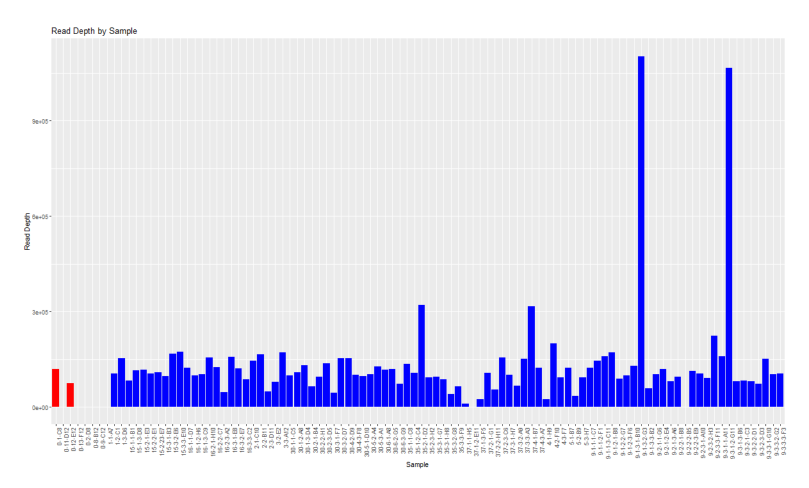


Figure 21. Sample read depths. The samples starting with a 0 code are controls. Those highlighted in red are positive controls, while the negative control read depths were so minimal, they did not appear on the Y axis.

Phylum	Class	Order	Family	Genus	Species
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Allolobophora	Allolobophora chlorotica
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Aporrectodea	Aporrectodea icterica complex sp. L1 DP- 2018
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Aporrectodea	Aporrectodea longa
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Aporrectodea	Aporrectodea rosea
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Bimastos	Bimastos rubidus
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Eiseniella	Eiseniella sp. BIOUG32056-F01
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Eiseniella	Eiseniella tetraedra
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Lumbricus	Lumbricus castaneus
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Lumbricus	Lumbricus festivus
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Lumbricus	Lumbricus terrestris
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Murchieona	Murchieona minuscula
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Octolasion	Octolasion cyaneum
Annelida	Clitellata	Crassiclitellata	Lumbricidae	Satchellius	Satchellius mammalis
Annelida	Clitellata	Enchytraeida	Enchytraeidae	Cernosvitoviella	Cernosvitoviella minor
Annelida	Clitellata	Enchytraeida	Enchytraeidae	Chamaedrilus	Chamaedrilus cognettii
Annelida	Clitellata	Enchytraeida	Enchytraeidae	Cognettia	Cognettia pseudosphagnetorum
Annelida	Clitellata	Enchytraeida	Enchytraeidae	Fridericia	Fridericia striata
Annelida	Clitellata	Enchytraeida	Enchytraeidae	Globulidrilus	Globulidrilus riparius
Annelida	Clitellata	Hirudinida	Erpobdellidae	Erpobdella	Erpobdella testacea
Annelida	Clitellata	Lumbriculida	Lumbriculidae	Lumbriculus	Lumbriculus variegatus
Phylum	Class	Order	Family	Genus	Species

# Appendix II. Complete Species List at 99% filtration identity level.

Annelida	Clitellata	Lumbriculida	Lumbriculidae	Stylodrilus	Stylodrilus heringianus
Annelida	Clitellata	Rhynchobdellida	Glossiphoniidae	Glossiphonia	Glossiphonia complanata
Annelida	Clitellata	Rhynchobdellida	Glossiphoniidae	Helobdella	Helobdella stagnalis
Annelida	Clitellata	Tubificida	Naididae	Aulodrilus	Aulodrilus pluriseta
Annelida	Clitellata	Tubificida	Naididae	Chaetogaster	Chaetogaster cf. diastrophus MK-2019
Annelida	Clitellata	Tubificida	Naididae	Chaetogaster	Chaetogaster diastrophus
Annelida	Clitellata	Tubificida	Naididae	Dero	Dero digitata
Annelida	Clitellata	Tubificida	Naididae	Dero	Dero obtusa
Annelida	Clitellata	Tubificida	Naididae	Ilyodrilus	Ilyodrilus templetoni
Annelida	Clitellata	Tubificida	Naididae	Limnodrilus	Limnodrilus claparedianus
Annelida	Clitellata	Tubificida	Naididae	Limnodrilus	Limnodrilus hoffmeisteri
Annelida	Clitellata	Tubificida	Naididae	Nais	Nais communis
Annelida	Clitellata	Tubificida	Naididae	Nais	Nais communis/variabilis complex sp. A2
Annelida	Clitellata	Tubificida	Naididae	Nais	Nais communis/variabilis complex sp. A3
Annelida	Clitellata	Tubificida	Naididae	Potamothrix	Potamothrix bavaricus
Annelida	Clitellata	Tubificida	Naididae	Potamothrix	Potamothrix hammoniensis
Annelida	Clitellata	Tubificida	Naididae	Potamothrix	Potamothrix heuscheri
Annelida	Clitellata	Tubificida	Naididae	Rhyacodrilus	Rhyacodrilus falciformis
Annelida	Clitellata	Tubificida	Naididae	Spirosperma	Spirosperma ferox
Annelida	Clitellata	Tubificida	Naididae	Stylaria	Stylaria lacustris
Annelida	Clitellata	Tubificida	Naididae	Tubifex	Tubifex tubifex
Annelida	Clitellata	unknown_order	unknown_family	unknown_genus	Oligochaeta sp. 1 RV-2016
Annelida	Polychaeta	unknown_order	Capitellidae	Dasybranchus	Dasybranchus sp. DH1
Arthropoda	Arachnida	Araneae	Anyphaenidae	Anyphaena	Anyphaena accentuata
Phylum	Class	Order	Family	Genus	Species

Arthropoda	Arachnida	Araneae	Clubionidae	Clubiona	Clubiona phragmitis
Arthropoda	Arachnida	Araneae	Linyphiidae	Gnathonarium	Gnathonarium dentatum
Arthropoda	Arachnida	Araneae	Lycosidae	Pardosa	Pardosa amentata
Arthropoda	Arachnida	Araneae	Pisauridae	Pisaura	Pisaura mirabilis
Arthropoda	Arachnida	Araneae	Theridiidae	Anelosimus	Anelosimus vittatus
Arthropoda	Arachnida	Araneae	Theridiidae	Paidiscura	Paidiscura pallens
Arthropoda	Arachnida	Araneae	Theridiidae	Theridion	Theridion varians
Arthropoda	Arachnida	Opiliones	Leiobunidae	Leiobunum	Leiobunum blackwalli
Arthropoda	Arachnida	Opiliones	Phalangiidae	Oligolophus	Oligolophus tridens
Arthropoda	Arachnida	Opiliones	Phalangiidae	Paroligolophus	Paroligolophus agrestis
Arthropoda	Arachnida	Opiliones	Sabaconidae	Sabacon	Sabacon viscayanus
Arthropoda	Arachnida	Sarcoptiformes	Acaridae	Tyrophagus	Tyrophagus curvipenis
Arthropoda	Arachnida	Sarcoptiformes	Acaridae	Tyrophagus	Tyrophagus fanetzhangorum
Arthropoda	Arachnida	Sarcoptiformes	Camisiidae	Platynothrus	Platynothrus peltifer
Arthropoda	Arachnida	Sarcoptiformes	Damaeidae	unknown_genus	Damaeidae sp. AMUEnv005
Arthropoda	Arachnida	Trombidiformes	Eriophyidae	Abacarus	Abacarus hystrix
Arthropoda	Arachnida	Trombidiformes	Eriophyidae	Aculodes	Aculodes mckenziei
Arthropoda	Arachnida	Trombidiformes	Penthaleidae	unknown_genus	Penthaleidae sp. Q091
Arthropoda	Arachnida	Trombidiformes	Pygmephoridae	Elattoma	Elattoma abeskoun
Arthropoda	Arachnida	Trombidiformes	Tydeidae	Tydeus	Tydeus sp. BMOC 17-0901-48
Arthropoda	Arachnida	unknown_order	unknown_family	unknown_genus	Arachnida sp. BOLD:ACM9770
Arthropoda	Branchiopoda	Diplostraca	Daphniidae	Daphnia	Daphnia curvirostris
Arthropoda	Branchiopoda	Diplostraca	Daphniidae	Daphnia	Daphnia longispina
Arthropoda	Branchiopoda	Diplostraca	Daphniidae	Daphnia	Daphnia pulex
Phylum	Class	Order	Family	Genus	Species

Arthropoda	Branchiopoda	Diplostraca	Daphniidae	Simocephalus	Simocephalus vetulus
Arthropoda	Branchiopoda	Diplostraca	Eurycercidae	Eurycercus	Eurycercus lamellatus
Arthropoda	Chilopoda	Geophilomorpha	Himantariidae	Stigmatogaster	Stigmatogaster subterranea
Arthropoda	Collembola	Entomobryomorpha	Entomobryidae	Lepidocyrtus	Lepidocyrtus cyaneus
Arthropoda	Collembola	Entomobryomorpha	Entomobryidae	Lepidocyrtus	Lepidocyrtus lanuginosus
Arthropoda	Collembola	Entomobryomorpha	Isotomidae	Desoria	Desoria trispinata
Arthropoda	Collembola	Entomobryomorpha	Isotomidae	Isotoma	Isotoma viridis
Arthropoda	Collembola	Entomobryomorpha	Isotomidae	Isotomurus	Isotomurus palustris
Arthropoda	Collembola	Entomobryomorpha	Isotomidae	Isotomurus	Isotomurus unifasciatus
Arthropoda	Collembola	Entomobryomorpha	Isotomidae	Parisotoma	Parisotoma aff. notabilis L0
Arthropoda	Collembola	Entomobryomorpha	Tomoceridae	Tomocerus	Tomocerus minor
Arthropoda	Collembola	Neelipleona	Neelidae	Megalothorax	Megalothorax minimus
Arthropoda	Collembola	Poduromorpha	Neanuridae	Neanura	Neanura muscorum
Arthropoda	Collembola	Poduromorpha	Poduridae	Podura	Podura aquatica
Arthropoda	Collembola	Symphypleona	Dicyrtomidae	Dicyrtomina	Dicyrtomina saundersi
Arthropoda	Collembola	Symphypleona	Dicyrtomidae	unknown_genus	Dicyrtomidae sp. BOLD:ACL8646
Arthropoda	Collembola	Symphypleona	Katiannidae	Sminthurinus	Sminthurinus aureus
Arthropoda	Collembola	Symphypleona	Katiannidae	Sminthurinus	Sminthurinus elegans
Arthropoda	Collembola	Symphypleona	Sminthuridae	Sminthurus	Sminthurus viridis
Arthropoda	Collembola	Symphypleona	Sminthurididae	Sminthurides	Sminthurides aquaticus
Arthropoda	Diplopoda	Julida	Julidae	Ophyiulus	Ophyiulus pilosus
Arthropoda	Hexanauplia	Cyclopoida	Cyclopidae	Acanthocyclops	Acanthocyclops vernalis
Arthropoda	Hexanauplia	Cyclopoida	Cyclopidae	Cyclops	Cyclops abyssorum
Arthropoda	Hexanauplia	Cyclopoida	Cyclopidae	Eucyclops	Eucyclops cf. serrulatus BOLD:AAZ6402
Phylum	Class	Order	Family	Genus	Species

Arthropoda	Hexanauplia	Harpacticoida	Canthocamptidae	Canthocamptus	Canthocamptus staphylinus
Arthropoda	Insecta	Coleoptera	Anobiidae	Anobium	Anobium punctatum
Arthropoda	Insecta	Coleoptera	Anthribidae	Choragus	Choragus sheppardi
Arthropoda	Insecta	Coleoptera	Cantharidae	Cantharis	Cantharis rustica
Arthropoda	Insecta	Coleoptera	Cantharidae	Crudosilis	Crudosilis ruficollis
Arthropoda	Insecta	Coleoptera	Carabidae	Bembidion	Bembidion biguttatum
Arthropoda	Insecta	Coleoptera	Carabidae	Carabus	Carabus problematicus
Arthropoda	Insecta	Coleoptera	Carabidae	Leistus	Leistus fulvibarbis
Arthropoda	Insecta	Coleoptera	Carabidae	Leistus	Leistus spinibarbis
Arthropoda	Insecta	Coleoptera	Carabidae	Nebria	Nebria brevicollis
Arthropoda	Insecta	Coleoptera	Chrysomelidae	Donacia	Donacia clavipes
Arthropoda	Insecta	Coleoptera	Chrysomelidae	Donacia	Donacia semicuprea
Arthropoda	Insecta	Coleoptera	Chrysomelidae	Donacia	Donacia simplex
Arthropoda	Insecta	Coleoptera	Chrysomelidae	Luperus	Luperus longicornis
Arthropoda	Insecta	Coleoptera	Chrysomelidae	Plateumaris	Plateumaris sericea
Arthropoda	Insecta	Coleoptera	Coccinellidae	Adalia	Adalia decempunctata
Arthropoda	Insecta	Coleoptera	Coccinellidae	Tytthaspis	Tytthaspis sedecimpunctata
Arthropoda	Insecta	Coleoptera	Curculionidae	Hylastes	Hylastes cunicularius
Arthropoda	Insecta	Coleoptera	Dytiscidae	Agabus	Agabus bipustulatus
Arthropoda	Insecta	Coleoptera	Dytiscidae	Agabus	Agabus guttatus
Arthropoda	Insecta	Coleoptera	Dytiscidae	Agabus	Agabus sturmii
Arthropoda	Insecta	Coleoptera	Dytiscidae	Hydroporus	Hydroporus palustris
Arthropoda	Insecta	Coleoptera	Erirhinidae	Stenopelmus	Stenopelmus rufinasus
Arthropoda	Insecta	Coleoptera	Helophoridae	Helophorus	Helophorus aequalis
Phylum	Class	Order	Family	Genus	Species

Phylum	Class	Order	Family	Genus	Species
Arthropoda	Insecta	Diptera	Chironomidae	Cricotopus	Cricotopus sylvestris
Arthropoda	Insecta	Diptera	Chironomidae	Cricotopus	Cricotopus cf. curtus ATNA376-09
Arthropoda	Insecta	Diptera	Chironomidae	Corynoneura	Corynoneura sp. 4ES
Arthropoda	Insecta	Diptera	Chironomidae	Chironomus	Chironomus muratensis
Arthropoda	Insecta	Diptera	Chironomidae	Chaetocladius	Chaetocladius melaleucus
Arthropoda	Insecta	Diptera	Chironomidae	Chaetocladius	Chaetocladius dissipatus
Arthropoda	Insecta	Diptera	Chironomidae	Camptocladius	Camptocladius stercorarius
Arthropoda	Insecta	Diptera	Chaoboridae	Chaoborus	Chaoborus flavicans
Arthropoda	Insecta	Diptera	Ceratopogonidae	Forcipomyia	Forcipomyia sp. 2ES
Arthropoda	Insecta	Diptera	Ceratopogonidae	Forcipomyia	Forcipomyia aristolochiae
Arthropoda	Insecta	Diptera	Ceratopogonidae	Culicoides	Culicoides impunctatus
Arthropoda	Insecta	Diptera	Ceratopogonidae	Culicoides	Culicoides chiopterus
Arthropoda	Insecta	Diptera	Ceratopogonidae	Brachypogon	Brachypogon nitidulus
Arthropoda	Insecta	Diptera	Calliphoridae	Calliphora	Calliphora vicina
Arthropoda	Insecta	Diptera	Bibionidae	Dilophus	Dilophus febrilis
Arthropoda	Insecta	Diptera	Bibionidae	Bibio	Bibio marci
Arthropoda	Insecta	Diptera	Agromyzidae	Chromatomyia	Chromatomyia milii
Arthropoda	Insecta	Coleoptera	Tenebrionidae	Lagria	Lagria hirta
Arthropoda	Insecta	Coleoptera	Staphylinidae	Ocypus	Ocypus olens
Arthropoda	Insecta	Coleoptera	Staphylinidae	Ocypus	Ocypus aeneocephalus
Arthropoda	Insecta	Coleoptera	Staphylinidae	Lesteva	Lesteva pubescens
Arthropoda	Insecta	Coleoptera	Staphylinidae	Lathrobium	Lathrobium fulvipenne
Arthropoda	Insecta	Coleoptera	Staphylinidae	Lathrobium	Lathrobium brunnipes
Arthropoda	Insecta	Coleoptera	Hydrophilidae	Hydrophilus	Hydrophilus piceus

Phylum	Class	Order	Family	Genus	Species
Arthropoda	Insecta	Diptera	Chironomidae	Trissopelopia	Trissopelopia longimana
Arthropoda	Insecta	Diptera	Chironomidae	Tanytarsus	Tanytarsus sylvaticus
Arthropoda	Insecta	Diptera	Chironomidae	Tanytarsus	Tanytarsus buchonius
Arthropoda	Insecta	Diptera	Chironomidae	Stempellinella	Stempellinella brevis
Arthropoda	Insecta	Diptera	Chironomidae	Smittia	Smittia sp. F190
Arthropoda	Insecta	Diptera	Chironomidae	Rheocricotopus	Rheocricotopus atripes
Arthropoda	Insecta	Diptera	Chironomidae	Pseudorthocladius	Pseudorthocladius pilosipennis
Arthropoda	Insecta	Diptera	Chironomidae	Pseudorthocladius	Pseudorthocladius filiformis
Arthropoda	Insecta	Diptera	Chironomidae	Psectrocladius	Psectrocladius platypus
Arthropoda	Insecta	Diptera	Chironomidae	Parochlus	Parochlus kiefferi
Arthropoda	Insecta	Diptera	Chironomidae	Orthocladius	Orthocladius schnelli
Arthropoda	Insecta	Diptera	Chironomidae	Orthocladius	Orthocladius frigidus
Arthropoda	Insecta	Diptera	Chironomidae	Orthocladius	Orthocladius dentifer
Arthropoda	Insecta	Diptera	Chironomidae	Micropsectra	Micropsectra roseiventris
Arthropoda	Insecta	Diptera	Chironomidae	Micropsectra	Micropsectra pallidula
Arthropoda	Insecta	Diptera	Chironomidae	Limnophyes	Limnophyes sp. 14ES
Arthropoda	Insecta	Diptera	Chironomidae	Limnophyes	Limnophyes pentaplastus
Arthropoda	Insecta	Diptera	Chironomidae	Limnophyes	Limnophyes minimus
Arthropoda	Insecta	Diptera	Chironomidae	Heterotanytarsus	Heterotanytarsus apicalis
Arthropoda	Insecta	Diptera	Chironomidae	Glyptotendipes	Glyptotendipes nr. paripes CH152
Arthropoda	Insecta	Diptera	Chironomidae	Eukiefferiella	Eukiefferiella minor
Arthropoda	Insecta	Diptera	Chironomidae	Eukiefferiella	Eukiefferiella ilkleyensis
Arthropoda	Insecta	Diptera	Chironomidae	Eukiefferiella	Eukiefferiella claripennis
Arthropoda	Insecta	Diptera	Chironomidae	Dicrotendipes	Dicrotendipes pulsus

Arthropoda	Insecta	Diptera	Chironomidae	unknown_genus	Chironomidae sp. DL-2020
Arthropoda	Insecta	Diptera	Chironomidae	unknown_genus	Chironomidae sp. RAK1
Arthropoda	Insecta	Diptera	Culicidae	Coquillettidia	Coquillettidia richiardii
Arthropoda	Insecta	Diptera	Culicidae	Culiseta	Culiseta annulata
Arthropoda	Insecta	Diptera	Drosophilidae	Drosophila	Drosophila suzukii
Arthropoda	Insecta	Diptera	Ephydridae	Hydrellia	Hydrellia maura
Arthropoda	Insecta	Diptera	Ephydridae	Notiphila	Notiphila dorsata
Arthropoda	Insecta	Diptera	Ephydridae	Scatella	Scatella paludum
Arthropoda	Insecta	Diptera	Limoniidae	Dicranomyia	Dicranomyia modesta
Arthropoda	Insecta	Diptera	Lonchopteridae	Lonchoptera	Lonchoptera lutea
Arthropoda	Insecta	Diptera	Mycetophilidae	Mycetophila	Mycetophila lunata
Arthropoda	Insecta	Diptera	Polleniidae	Pollenia	Pollenia labialis
Arthropoda	Insecta	Diptera	Psychodidae	Psychoda	Psychoda phalaenoides
Arthropoda	Insecta	Diptera	Ptychopteridae	Ptychoptera	Ptychoptera contaminata
Arthropoda	Insecta	Diptera	Rhagionidae	Rhagio	Rhagio tringarius
Arthropoda	Insecta	Diptera	Scathophagidae	Scathophaga	Scathophaga sp. BIOUG02375-A02
Arthropoda	Insecta	Diptera	Simuliidae	Simulium	Simulium armoricanum
Arthropoda	Insecta	Diptera	Simuliidae	Simulium	Simulium aureum
Arthropoda	Insecta	Diptera	Simuliidae	Simulium	Simulium velutinum
Arthropoda	Insecta	Diptera	Syrphidae	Eristalinus	Eristalinus sepulchralis
Arthropoda	Insecta	Diptera	Syrphidae	Eristalis	Eristalis pertinax
Arthropoda	Insecta	Diptera	Syrphidae	Helophilus	Helophilus pendulus
Arthropoda	Insecta	Diptera	Tipulidae	Tipula	Tipula oleracea
Arthropoda	Insecta	Diptera	Tipulidae	Tipula	Tipula paludosa
Phylum	Class	Order	Family	Genus	Species

Arthropoda	Insecta	Diptera	unknown_family	unknown_genus	Diptera sp. RAK1
Arthropoda	Insecta	Ephemeroptera	Baetidae	Baetis	Baetis rhodani
Arthropoda	Insecta	Ephemeroptera	Baetidae	Cloeon	Cloeon dipterum
Arthropoda	Insecta	Ephemeroptera	Heptageniidae	Rhithrogena	Rhithrogena semicolorata
Arthropoda	Insecta	Hemiptera	Aphididae	Amphorophora	Amphorophora rubi
Arthropoda	Insecta	Hemiptera	Aphididae	Aulacorthum	Aulacorthum solani
Arthropoda	Insecta	Hemiptera	Aphididae	Ceruraphis	Ceruraphis eriophori
Arthropoda	Insecta	Hemiptera	Aphididae	Elatobium	Elatobium abietinum
Arthropoda	Insecta	Hemiptera	Aphididae	Hyalopterus	Hyalopterus pruni
Arthropoda	Insecta	Hemiptera	Aphididae	Metopolophium	Metopolophium dirhodum
Arthropoda	Insecta	Hemiptera	Aphididae	Periphyllus	Periphyllus hirticornis
Arthropoda	Insecta	Hemiptera	Aphididae	Periphyllus	Periphyllus testudinaceus
Arthropoda	Insecta	Hemiptera	Aphididae	Rhopalosiphum	Rhopalosiphum enigmae
Arthropoda	Insecta	Hemiptera	Aphididae	Rhopalosiphum	Rhopalosiphum padi
Arthropoda	Insecta	Hemiptera	Aphididae	Schizaphis	Schizaphis graminum
Arthropoda	Insecta	Hemiptera	Aphididae	Tuberolachnus	Tuberolachnus salignus
Arthropoda	Insecta	Hemiptera	Aphrophoridae	Neophilaenus	Neophilaenus lineatus
Arthropoda	Insecta	Hemiptera	Cicadellidae	Aphrodes	Aphrodes makarovi
Arthropoda	Insecta	Hemiptera	Cicadellidae	Fagocyba	Fagocyba douglasi
Arthropoda	Insecta	Hemiptera	Cicadellidae	lassus	lassus lanio
Arthropoda	Insecta	Hemiptera	Cicadellidae	Idiocerus	Idiocerus herrichii
Arthropoda	Insecta	Hemiptera	Cicadellidae	Ribautiana	Ribautiana debilis
Arthropoda	Insecta	Hemiptera	Coccidae	Pulvinaria	Pulvinaria idesiae
Arthropoda	Insecta	Hemiptera	Corixidae	Hesperocorixa	Hesperocorixa linnaei
Phylum	Class	Order	Family	Genus	Species

Arthropoda	Insecta	Hemiptera	Corixidae	Hesperocorixa	Hesperocorixa sahlbergi
Arthropoda	Insecta	Hemiptera	Gerridae	Gerris	Gerris lacustris
Arthropoda	Insecta	Hemiptera	Hydrometridae	Hydrometra	Hydrometra stagnorum
Arthropoda	Insecta	Hemiptera	Nepidae	Nepa	Nepa cinerea
Arthropoda	Insecta	Hemiptera	Notonectidae	Notonecta	Notonecta glauca
Arthropoda	Insecta	Hemiptera	Pemphigidae	Thecabius	Thecabius affinis
Arthropoda	Insecta	Hemiptera	Psyllidae	Cacopsylla	Cacopsylla melanoneura
Arthropoda	Insecta	Hemiptera	Psyllidae	Cacopsylla	Cacopsylla sp. SO-2015
Arthropoda	Insecta	Hemiptera	Psyllidae	Psylla	Psylla alni
Arthropoda	Insecta	Hemiptera	Triozidae	Trioza	Trioza urticae
Arthropoda	Insecta	Hemiptera	Veliidae	Velia	Velia caprai
Arthropoda	Insecta	Hymenoptera	Formicidae	Formica	Formica fusca
Arthropoda	Insecta	Hymenoptera	Formicidae	Lasius	Lasius flavus
Arthropoda	Insecta	Hymenoptera	Formicidae	Lasius	Lasius niger
Arthropoda	Insecta	Hymenoptera	Formicidae	Myrmica	Myrmica ruginodis
Arthropoda	Insecta	Hymenoptera	Tenthredinidae	Euura	Euura imperfecta
Arthropoda	Insecta	Hymenoptera	Tenthredinidae	Pristiphora	Pristiphora nigella
Arthropoda	Insecta	Lepidoptera	Crambidae	Cataclysta	Cataclysta lemnata
Arthropoda	Insecta	Lepidoptera	Crambidae	Chrysoteuchia	Chrysoteuchia culmella
Arthropoda	Insecta	Lepidoptera	Crambidae	Donacaula	Donacaula forficella
Arthropoda	Insecta	Lepidoptera	Depressariidae	Depressaria	Depressaria ultimella
Arthropoda	Insecta	Lepidoptera	Elachistidae	Spuleria	Spuleria flavicaput
Arthropoda	Insecta	Lepidoptera	Erebidae	Diaphora	Diaphora mendica
Arthropoda	Insecta	Lepidoptera	Gracillariidae	Cameraria	Cameraria ohridella
Phylum	Class	Order	Family	Genus	Species

Arthropoda	Insecta	Lepidoptera	Hepialidae	Phymatopus	Phymatopus hecta
Arthropoda	Insecta	Lepidoptera	Lasiocampidae	Euthrix	Euthrix potatoria
Arthropoda	Insecta	Lepidoptera	Nepticulidae	Stigmella	Stigmella ulmivora
Arthropoda	Insecta	Lepidoptera	Noctuidae	Apamea	Apamea crenata
Arthropoda	Insecta	Lepidoptera	Noctuidae	Cerapteryx	Cerapteryx graminis
Arthropoda	Insecta	Lepidoptera	Noctuidae	Noctua	Noctua fimbriata
Arthropoda	Insecta	Lepidoptera	Noctuidae	Noctua	Noctua pronuba
Arthropoda	Insecta	Lepidoptera	Noctuidae	Phlogophora	Phlogophora meticulosa
Arthropoda	Insecta	Lepidoptera	Noctuidae	Xestia	Xestia xanthographa
Arthropoda	Insecta	Lepidoptera	Nymphalidae	Pararge	Pararge aegeria
Arthropoda	Insecta	Lepidoptera	Tortricidae	Ancylis	Ancylis achatana
Arthropoda	Insecta	Lepidoptera	Tortricidae	Epinotia	Epinotia nisella
Arthropoda	Insecta	Lepidoptera	Tortricidae	Gypsonoma	Gypsonoma dealbana
Arthropoda	Insecta	Mecoptera	Panorpidae	Panorpa	Panorpa germanica
Arthropoda	Insecta	Neuroptera	Hemerobiidae	Hemerobius	Hemerobius lutescens
Arthropoda	Insecta	Odonata	Aeshnidae	Anax	Anax imperator
Arthropoda	Insecta	Odonata	Coenagrionidae	Pyrrhosoma	Pyrrhosoma nymphula
Arthropoda	Insecta	Orthoptera	Acrididae	Chorthippus	Chorthippus binotatus
Arthropoda	Insecta	Plecoptera	Leuctridae	Leuctra	Leuctra hippopus
Arthropoda	Insecta	Plecoptera	Leuctridae	Leuctra	Leuctra inermis
Arthropoda	Insecta	Plecoptera	Leuctridae	Leuctra	Leuctra nigra
Arthropoda	Insecta	Psocoptera	Trichopsocidae	Trichopsocus	Trichopsocus sp. KY322
Arthropoda	Insecta	Thysanoptera	Thripidae	Anaphothrips	Anaphothrips obscurus
Arthropoda	Insecta	Thysanoptera	Thripidae	Aptinothrips	Aptinothrips rufus
Phylum	Class	Order	Family	Genus	Species

Arthropoda	Insecta	Trichoptera	Beraeidae	Beraea	Beraea pullata
Arthropoda	Insecta	Trichoptera	Hydroptilidae	Agraylea	Agraylea multipunctata
Arthropoda	Insecta	Trichoptera	Limnephilidae	Glyphotaelius	Glyphotaelius pellucidus
Arthropoda	Insecta	Trichoptera	Limnephilidae	Halesus	Halesus radiatus
Arthropoda	Malacostraca	Amphipoda	Crangonyctidae	Crangonyx	Crangonyx pseudogracilis
Arthropoda	Malacostraca	Isopoda	Asellidae	Asellus	Asellus aquaticus
Arthropoda	Malacostraca	Isopoda	Oniscidae	Oniscus	Oniscus asellus
Arthropoda	Malacostraca	Isopoda	Philosciidae	Philoscia	Philoscia muscorum
Arthropoda	Malacostraca	Isopoda	Porcellionidae	Porcellio	Porcellio scaber
Arthropoda	Ostracoda	Podocopida	Candonidae	Candona	Candona candida
Arthropoda	Ostracoda	Podocopida	Candonidae	Candona	Candona neglecta
Arthropoda	Ostracoda	Podocopida	Candonidae	Candonopsis	Candonopsis kingsleii
Arthropoda	Ostracoda	Podocopida	Cyprididae	Eucypris	Eucypris virens
Arthropoda	unknown_class	unknown_order	unknown_family	unknown_genus	Maxillopoda sp. BOLD:ACW5478
Arthropoda	unknown_class	unknown_order	unknown_family	unknown_genus	Maxillopoda sp. BOLD:ACW5664
Bryozoa	Phylactolaemata	unknown_order	Lophopodidae	Lophopus	Lophopus crystallinus
Bryozoa	Phylactolaemata	unknown_order	Plumatellidae	Plumatella	Plumatella fungosa
Chordata	Actinopteri	Cypriniformes	Leuciscidae	Rutilus	Rutilus rutilus
Chordata	Actinopteri	Cypriniformes	Leuciscidae	Scardinius	Scardinius erythrophthalmus
Chordata	Actinopteri	Esociformes	Esocidae	Esox	Esox lucius
Chordata	Actinopteri	Perciformes	Gasterosteidae	Gasterosteus	Gasterosteus aculeatus
Chordata	Amphibia	Anura	Ranidae	Rana	Rana temporaria
Chordata	Amphibia	Caudata	Salamandridae	Lissotriton	Lissotriton vulgaris
Chordata	Aves	Anseriformes	Anatidae	Anas	Anas platyrhynchos
Phylum	Class	Order	Family	Genus	Species

Chordata	Aves	Anseriformes	Anatidae	Cygnus	Cygnus olor
Chordata	Aves	Charadriiformes	Scolopacidae	Tringa	Tringa totanus
Chordata	Aves	Columbiformes	Columbidae	Columba	Columba oenas
Chordata	Aves	Columbiformes	Columbidae	Columba	Columba palumbus
Chordata	Aves	Gruiformes	Rallidae	Fulica	Fulica atra
Chordata	Aves	Gruiformes	Rallidae	Rallus	Rallus aquaticus
Chordata	Aves	Passeriformes	Corvidae	Coloeus	Coloeus monedula
Chordata	Aves	Passeriformes	Corvidae	Garrulus	Garrulus glandarius
Chordata	Aves	Passeriformes	Corvidae	Pica	Pica pica
Chordata	Aves	Passeriformes	Fringillidae	Chloris	Chloris chloris
Chordata	Aves	Passeriformes	Fringillidae	Fringilla	Fringilla coelebs
Chordata	Aves	Passeriformes	Fringillidae	Linaria	Linaria cannabina
Chordata	Aves	Passeriformes	Fringillidae	Pyrrhula	Pyrrhula pyrrhula
Chordata	Aves	Passeriformes	Sylviidae	Sylvia	Sylvia atricapilla
Chordata	Aves	Passeriformes	Turdidae	Turdus	Turdus iliacus
Chordata	Aves	Passeriformes	Turdidae	Turdus	Turdus philomelos
Chordata	Aves	Passeriformes	Turdidae	Turdus	Turdus pilaris
Chordata	Aves	Passeriformes	Turdidae	Turdus	Turdus viscivorus
Chordata	Mammalia	Artiodactyla	Bovidae	Bos	Bos taurus
Chordata	Mammalia	Artiodactyla	Bovidae	Ovis	Ovis aries
Chordata	Mammalia	Carnivora	Canidae	Canis	Canis lupus
Chordata	Mammalia	Eulipotyphla	Soricidae	Sorex	Sorex araneus
Chordata	Mammalia	Eulipotyphla	Soricidae	Sorex	Sorex minutus
Chordata	Mammalia	Perissodactyla	Equidae	Equus	Equus caballus
Phylum	Class	Order	Family	Genus	Species

Chordata	Mammalia	Primates	Hominidae	Homo	Homo sapiens
Chordata	Mammalia	Rodentia	Cricetidae	Microtus	Microtus agrestis
Chordata	Mammalia	Rodentia	Gliridae	Muscardinus	Muscardinus avellanarius
Chordata	Mammalia	Rodentia	Muridae	Mus	Mus musculus
Chordata	Mammalia	Rodentia	Muridae	Rattus	Rattus norvegicus
Cnidaria	Hydrozoa	Anthoathecata	Hydridae	Hydra	Hydra circumcincta
Cnidaria	Hydrozoa	Anthoathecata	Hydridae	Hydra	Hydra oligactis
Cnidaria	Hydrozoa	Anthoathecata	Hydridae	Hydra	Hydra viridissima
Cnidaria	Hydrozoa	Anthoathecata	Hydridae	Hydra	Hydra vulgaris
Gastrotricha	unknown_class	Chaetonotida	Chaetonotidae	Chaetonotus	Chaetonotus aff. persimilis MK-2019
Gastrotricha	unknown_class	Chaetonotida	Chaetonotidae	Chaetonotus	Chaetonotus aff. subtilis 4 MK-2019
Gastrotricha	unknown_class	Chaetonotida	Chaetonotidae	Chaetonotus	Chaetonotus borealis
Gastrotricha	unknown_class	Chaetonotida	Chaetonotidae	Chaetonotus	Chaetonotus jaceki
Gastrotricha	unknown_class	Chaetonotida	Chaetonotidae	Heterolepidoderma	Heterolepidoderma ocellatum
Gastrotricha	unknown_class	Chaetonotida	Chaetonotidae	Polymerurus	Polymerurus rhomboides
Gastrotricha	unknown_class	Chaetonotida	Dasydytidae	Stylochaeta	Stylochaeta scirtetica
Mollusca	Bivalvia	Galeommatida	Montacutidae	Kurtiella	Kurtiella bidentata
Mollusca	Bivalvia	Venerida	Sphaeriidae	Pisidium	Pisidium obtusale
Mollusca	Bivalvia	Venerida	Sphaeriidae	Pisidium	Pisidium subtruncatum
Mollusca	Bivalvia	Venerida	Sphaeriidae	Sphaerium	Sphaerium corneum
Mollusca	Bivalvia	Venerida	Sphaeriidae	Sphaerium	Sphaerium nucleus
Mollusca	Gastropoda	Littorinimorpha	Bithyniidae	Bithynia	Bithynia tentaculata
Mollusca	Gastropoda	Stylommatophora	Agriolimacidae	Deroceras	Deroceras invadens
Mollusca	Gastropoda	Stylommatophora	Agriolimacidae	Deroceras	Deroceras laeve
Phylum	Class	Order	Family	Genus	Species

Mollusca	Gastropoda	Stylommatophora	Arionidae	Arion	Arion hortensis	
Mollusca	Gastropoda	Stylommatophora	Arionidae	Arion	Arion intermedius	
Mollusca	Gastropoda	Stylommatophora	Arionidae	Arion	Arion subfuscus	
Mollusca	Gastropoda	Stylommatophora	Helicidae	Cepaea	Cepaea nemoralis	
Mollusca	Gastropoda	Stylommatophora	Hygromiidae	Monacha	Monacha cantiana	
Mollusca	Gastropoda	Stylommatophora	Hygromiidae	Zenobiellina	Zenobiellina subrufescens	
Mollusca	Gastropoda	Stylommatophora	Succineidae	Succinea	Succinea putris	
Mollusca	Gastropoda	Stylommatophora	Vitrinidae	Vitrina	Vitrina pellucida	
Mollusca	Gastropoda	unknown_order	Acroloxidae	Acroloxus	Acroloxus lacustris	
Mollusca	Gastropoda	unknown_order	Lymnaeidae	Ampullaceana	Ampullaceana balthica	
Mollusca	Gastropoda	unknown_order	Physidae	Aplexa	Aplexa hypnorum	
Mollusca	Gastropoda	unknown_order	Physidae	Physa	Physa fontinalis	
Mollusca	Gastropoda	unknown_order	Physidae	Physella	Physella ancillaria	
Mollusca	Gastropoda	unknown_order	Planorbidae	Anisus	Anisus cf. vortex P2333	
Mollusca	Gastropoda	unknown_order	Planorbidae	Hippeutis	Hippeutis complanatus	
Mollusca	Gastropoda	unknown_order	Planorbidae	Planorbis	Planorbis planorbis	
Nematoda	unknown_class	unknown_order	unknown_family	unknown_genus	unidentified nematode	
Nemertea	unknown_class	unknown_order	unknown_family	unknown_genus	Nemertean sp. NT000047	
Platyhelminthes	Catenulida	unknown_order	Stenostomidae	Stenostomum	Stenostomum cf. simplex AW-2018	
Platyhelminthes	Rhabditophora	Macrostomida	Microstomidae	Microstomum	Microstomum lineare	
Platyhelminthes	Rhabditophora	Tricladida	Dugesiidae	Schmidtea	Schmidtea polychroa	
Rotifera	Eurotatoria	Adinetida	Adinetidae	Adineta	Adineta sp. FR.5	
Rotifera	Eurotatoria	Adinetida	Adinetidae	Adineta	Adineta vaga	
Rotifera	Eurotatoria	Adinetida	Adinetidae	Adineta	Adineta vaga complex sp. B JFF-2016	
Phylum	Class	Order	Family	Genus	Species	

Rotifera	Eurotatoria	Philodinida	Habrotrochidae	Habrotrocha	Habrotrocha elusa
Rotifera	Eurotatoria	Philodinida	Habrotrochidae	Habrotrocha	Habrotrocha ligula
Rotifera	Eurotatoria	Philodinida	Philodinidae	Macrotrachela	Macrotrachela quadricornifera
Rotifera	Eurotatoria	Philodinida	Philodinidae	Philodina	Philodina citrina
Rotifera	Eurotatoria	Philodinida	Philodinidae	Philodina	Philodina sp. A459_PR6
Rotifera	Eurotatoria	Ploima	Asplanchnidae	Asplanchna	Asplanchna sieboldii
Rotifera	Eurotatoria	Ploima	Brachionidae	Brachionus	Brachionus calyciflorus
Rotifera	Eurotatoria	Ploima	Brachionidae	Euchlanis	Euchlanis dilatata
Rotifera	Eurotatoria	Ploima	Brachionidae	Mytilina	Mytilina mucronata
Rotifera	Eurotatoria	Ploima	Epiphanidae	Epiphanes	Epiphanes senta
Rotifera	Eurotatoria	Ploima	Gastropidae	Ascomorpha	Ascomorpha ecaudis
Rotifera	Eurotatoria	Ploima	Lecanidae	Lecane	Lecane closterocerca
Rotifera	Eurotatoria	Ploima	Proalidae	Proales	Proales daphnicola
Rotifera	Eurotatoria	Ploima	Synchaetidae	Synchaeta	Synchaeta cf. tremula/oblonga UO-2012
Rotifera	Eurotatoria	Ploima	Synchaetidae	Synchaeta	Synchaeta pectinata
Rotifera	Eurotatoria	unknown_order	unknown_family	Rotaria	Rotaria macroceros
Rotifera	Eurotatoria	unknown_order	unknown_family	Rotaria	Rotaria magnacalcarata
Rotifera	Eurotatoria	unknown_order	unknown_family	Rotaria	Rotaria rotatoria
Rotifera	Eurotatoria	unknown_order	unknown_family	Rotaria	Rotaria socialis
Rotifera	Eurotatoria	unknown_order	unknown_family	Rotaria	Rotaria sp. RotS1

# 7.1 Appendix i. Biological Monitoring Working Party (BMWP) Average Score per

## Taxon (ASPT) Scoring System

Taxonomic Class	Taxonomic Families	Score
Ephemeroptera	Ephemeridae	10
	Heptagoniidae	10
	Leptophlebiidae	10
	Pothamanthidae	10
	Siphonurridae	10
Plecoptera	Capniidae	10
	Chloroperlidae	10
	Leuctridae	10
	Perlidae	10
	Taeniopteterygidae	10
Hemiptera	Aphelochereididae	10
Trichoptera	Beraecidae	10
	Brachycentridae	10
	Goeridae	10
	Lepidostomatidae	10
	Leptoceridae	10
	Mollanidae	10
	Odontoceridae	10
	Phyrgancineidae	10
	Sericostomatidae	10
Ephemeroptera	Caenidae	7
Plecoptera	Nemouridae	7
Trichoptera	Rhyacophilidae	7
	Polycentropodidae	7
	Limnepphilidae	7
Mollusca	Neritidae	6

	Viviparidae	6
	Ancylidae	6
	Unionidae	6
Trichoptera	Hydroptilidae	6
Crustacea	Corophiidae	6
	Gammaridae	6
	Paleamonidae	6
Polychaeta	Nereidae	6
	Nephthyidae	6
Odonata	Plaqthycnemididae	6
	Coenagriidae	6
Hemiptera	Mesovelidae	5
	Hydrometridae	5
	Gerridae	5
	Nepidae	5
	Naucoridae	5
	Notonectidae	5
	Pletidae	5
Coleoptera	Chrysomelidae	5
	Corixidae	5
	Curculionidae	5
	Dryopidae	5
	Dytiscidae	5
	Eliminthidae	5
	Gyrinidae	5
	Haliplidae	5
	Helobidae	5
	Hydrophilidae	5
	Hygrobiidae	5
Phyrgancineidae	Hydropsychidae	5

Diptera	Tipulidae	5
	Simuliidae	5
Planaria	Planariidae	5
	Dendrocoelidae	5
Ephemeroptera	Baetidae	4
Megaloptera	Sialidae	4
Hirudinida	Piscicolidae	4
Mollusca	Valvatidae	3
	Hygrobiidae	3
	Lymnaeitidae	3
	Physidae	3
	Planorbidae	3
	Sphaeriidae	3
Hirudinida	Erpobdellidae	3
	Glossiphonidae	3
	Hirudidae	3
Others	Alderfly (meglaoptera, Sialidae)	4
	Shrimps (Caridea)	6
	Hoglice (Asellidae)	3
	Blackfly (Simuliidae)	5
	Cranefly (Tipulidae)	5
	Worms	1