Aquatic invertebrate community structure and phenology of the intermittent treed swamps of the semi-arid Paroo lowlands in Australia

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Abstract The middle Paroo lowlands in semi-arid western New South Wales support numerous intermittent wetlands of various types. Differences between them are promoted by three ecological drivers: salinity, turbidity and hydroperiod. Community structure and phenology of the two most common types, saline lakes and claypans, are known but similar ecologies are lacking for the third most common wetland, the treed swamps. These are of six subtypes distinguished by dominant tree species, geomorphology and hydroperiod, all with similar community structure and phenology, but with differing invertebrate diversities. Summed diversity is not as high as in local creek pools, the shorter hydroperiods and simpler geomorphology of the treed swamps being restrictive so that there is almost no replacement of species during the early dominance of branchiopods and later of insects. Such treed swamps are uncommon in the semi-arid zone, but much more speciose treed swamps are known under similar and seasonally dry Mediterranean climates of the Western Australian Wheatbelt where hydroperiods are more stable.

Keywords Branchiopod crustaceans · Insects · Diversity · Hydroperiod · Habitat heterogeneity

Introduction

Of all the intermittent wetlands of the world, most is known on those of Mediterranean lands (Boix et al. 2016) including of the two Mediterranean climate zones in Australia: southwest Western Australia and southeast South Australia-western Victoria (e.g. Bayly and Williams 1966, Davis et.al. 1993, Horwitz et al. 2009 and Pinder et al. 2004). The major environmental control in these is seasonal drying, which is shared with the dryland wetlands of the semiarid climates, though more intensely and less seasonally predictable. Adaptations to survive intermittent drying are similar in each (Williams 1985). These include resistant eggs of the branchiopod crustaceans, salinity tolerance of many crustaceans and insects, tolerance to high temperatures and good dispersal mechanisms. Introductions to Australia’s arid and semiarid wetlands are provided by Davis et al. 2013, 2018 and Timms and Boulton 2001. In the Australian semiarid zone, the middle and lower Paroo lowlands on a plainlands in north-western New South Wales contain many and varied wetlands, almost all unreliably flooded (Goodrick 1984; Timms and Boulton 2001) (Fig. 1). All occur in hollows due

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either to dunes blocking drainage lines, or to localised deflation. An exception, not considered here, are the riverine waterholes due to evorsion by water currents and containing water permanently.

The intermittent wetlands of this region are filled either by riverine floodwater or from local rainfall. Both types can support trees. Prominant among the former are lignum swamps (*Muehlenbeckia cunninghamii*) and Yapunyah (*Eucalyptus ochrophloia*) tree swamps. These are inundated for many months following a flood and have a distinctive invertebrate fauna including crabs and decapod shrimps (Timms 1997a) but are not included here. This paper focuses on the wetlands filled entirely from local rainfall which includes saline lakes (large and with bare surfaces, or small and with samphires (*Arthrocnemum* spp.), innumerable shallow claypans, many varieties of vegetated swamps and marshes, and a few temporary creek pools, small grassy pools, and shallow flood outs. The ecological drivers characterising these wetlands are salinity, turbidity and hydrology (Timms and Boulton 2001). For the treed swamps, other drivers may be associated with the trees, swamp structure affecting invertebrate diversity, changes in the water as it ages, and phenology.

Those wetlands filled by local rainfall have a distinctive variable hydrology and fauna. Gafney’s (1975) variability index (90% percentile—10% percentile divided by the 50% percentile of annual rainfall) is close to 1.50 for the area, which is not far from the extreme value of 1.75 for Australia. Of the aquatic fauna of the area, only the fauna of the saline lakes have been sufficiently studied to claim it is an inland one distinct from those of various groups of southern lakes (Timms 2007). The anostracan fauna is of eastern Australian character (Rogers and Timms 2014) and, judging from their listings in many regional studies, most of the inland insects are widespread across the continent.

The characteristics of many of these wetlands have now been elucidated. Foremost are the saline lakes (Timms 1993, 1998a, 2018), followed by claypans (Hancock and Timms 2002; Timms 2002), creek pools (Timms 2001), with less known of the large episodic terminal lakes, smaller intermittent freshwater lakes and permanent riverine waterholes (Timms and Boulton 2001). Least is known of the fauna, phenology and ecological drivers of the vegetated temporary swamps surrounded by trees and supporting a few trees within. This paper considers these, the last of the major Paroo wetlands to be studied.

The trees in these Paroo wetlands are probably not integral to their ecology. Certainly they are the most obvious feature seen from near and afar, but they are largely peripheral and their bases are only inundated when such a swamp is first flooded. They probably supply some leaf litter and more noticeably woody detritus which may add to invertebrate diversity (see later) and also perhaps shade the littoral from the fierce summer sun. Locally there are two dominant tree species, leading to apparently different wetland types: Black Box (*Eucalyptus largiflorens*) swamps and Poplar Box (*E. populnea*) flats. The Black Box swamps occur almost exclusively in the dune fields and the Poplar Black Box flats mainly in the red earth country; also Black Box swamps tend to be smaller and deeper than Poplar Box flats and with a longer hydroperiod maybe 4–6 months versus 2–5 months, but there are some exceptions. Both wetland types may have internal channels, a feature of some aridzone wetlands (Tooth and McCarthy 2007), but only Black Box swamps have uneven deposits of ex-terrestrial soils deposited during major inflows. Black Box trees also occur as tree cover in some lignum swamps, and in drainage lines across a wide

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**Fig. 1** Map of Bloodwood, Tredega and Muella Stations showing locations of the 18 treed swamps and four other sites. Coded to wetlands (all names unofficial except 21, 22 and used only locally): 1. Fence, 2. Far Homestead, 3. False Goat, 4. Reedy, 5. Washout, 6. Real Goat, 7. Utah, 8. George, 9. Donkey, 10. Carters, 11. Muella East South, 12. Muella East North, 13. Lismore, 14. Near Homestead, 15. Boggy Bend, 16. Quandong, 17. Tredega T/O, 18. Marsilea, 19. Ski Lake, 20. Turkey claypan, 21. Gidgee salt lake, 22. Horseshoe salt lake
area of the southern and central eastern inland (Cunningham et al. 1981).

Treed swamps in drylands are uncommon across the world with most apparently being in Australia and then being concentrated in the Middle Paroo catchment of northwestern New South Wales. In addition, the mediterranean zone of Western Australia has a few different types of wetlands dominated by trees, each with a different diversity of invertebrates, some halotolerant. Trees include Swamp Yate (*E. occidentalis*), and *Melaleuca* spp. Another concentration of treed swamps occurs in the sahel of Senegal, Africa (Lahr et al. 1999).

The purpose of this paper is to investigate of the basic ecological character of these treed swamps, one of the major dryland wetland types of the middle Paroo. In a comprehensive study of Paroo wetlands, they were left to last because of the difficulty in sampling soft and retreating shorelines and the perceived need to operate in a major flooding which occurs only occasionally. Besides characterising their flooding regime, variations in their conductivity, pH and turbidity were monitored. It is assumed their fresh waters are dominated by Na and Cl ions, as are almost all waters in the Paroo (Timms 1993). Invertebrate species diversity is an important discriminator between Paroo wetlands not only between major types but perhaps between subtypes as suspected among the treed swamps. So many questions arise, including 1. Are their distinctions between the subtypes? and 2. Are the ecological drivers in these temporary freshwater wetlands similar to most Australian arid-zone wetlands (Davis et al. 2018) and indeed the wider arid zone or can they be explained within those already known for other Paroo wetlands, i.e. salinity, turbidity and hydrology?

**Study site and methods**

The study area lies on three adjacent properties, Bloodwood, Tredega and Muella Stations, 130 km northwest of Bourke in western New South Wales and 50 km east of the Paroo floodplain. The study area is 250 km² and consists of two types of country—undulating low ridges on sedimentary and silcrete rocks covered by red earth soils to the east and dunefields of deep siliceous sands to the west (Hazelton and Johnson 1973).

Mean annual rainfall is 307 mm at Dungarvon Station immediately to their west (recorded by D. Leigo 1966 to 2016). Rainfall is maximal in summer (mean 98 mm) and minimal in winter (mean 60 mm) (C. Sives, pers. com.), with occasional large falls (e.g. 261 mm over two days in March 2021) and long droughts. Although there is some seasonality in the averages, the reality is rainfall is highly variable from month to month, and year to year. Evaporation is more reliable averaging about 3000 mm per year for this area (Warner 1977). What seems an anomaly, the amount of water in the bigger lakes is often maximal in winter and minimal in spring (C. Sives, pers.com) but this is due the overriding influence of seasonal variation in evaporation. Typically smaller wetlands studied here fill in mid to late summer and are dry again by winter or spring, but there exceptional years.

This study is based mainly on a major fill on March 3–5, 2020 resulting from 261 mm rainfall and very little afterwards. Eighteen treed swamps on three adjacent stations, Bloodwood, Tredega and Muella in the Middle Paroo were chosen and subsequently monitored monthly for six months (March 19–23; May 5–7; July 3–5; August 1–3; September, 6–8; but missed in June due to covid-19 travel restrictions (Figs. 1, 2). Also some data were collected from the same sites in more average flooding in May 2019, but it could only be followed for 6 weeks at fortnightly intervals. Also, comparative collections were taken from a nearby claypan, a freshwater lake and two saline lakes on Bloodwood. The treed swamps were grouped into six categories with an aim of three in each. The first criterion was the tree species present, and the second whether or not channels were present, and the third was size (> or < 0.5 ha), all factors which may be additional ecological drivers. Comments on overall filling frequency are based on field experience 1987–2021 inclusive.

In the main study, water samples from the 22 sites were collected regularly and conductivity measured with an Oakton ECTestr11 meter, pH with a Hanna pHep meter and turbidity with a Secchi disc tube. Maximal inundated area of each site was estimated from Google Earth maps by comparing ground extent in March 2021 with features visible on their 2015 image. Note that inundated areas vary with each fill as not one of the swamps overflow, but the levels of March 2021 were near maximal seen during 1987–2021. Depths were determined by a dumpy...
level from the vertical height of a marker installed at full level in March and the deepest place when the swamps dried. Habitat heterogeneity was subjectively measured equally by extent of bottom irregularity determined as three values—low, medium, high, the area estimated to be vegetated during the inundation period, again recorded as one of three values—low, medium, high, and the amount of inundated woody
died due to excessive water. a Reedy Black Box swamp showing the typical ring of trees around the swamp and often inundated on initial flooding. The dead trees relate back to an extended flooding when they

b Utah poplar box flat showing an extensive plain with a few trees but many at the far edge and a channel within the wetland
detritus across the floor of the swamp as counted when the site was dry, again as three values—low, medium, high. The result was a simple index ranging from 1 to 9, not detailed, but of sufficient robustness for purpose. Physicochemical measurements are summarised in SM1.

All biological sampling was done from the swamp edge, as it was dangerous to wander too far out due to unpredictable location of unconsolidated wetted sediments. So, while every effort was made to make the samples representative, this was not strictly possible, i.e. all samples were taken from different physical sites as each swamp dried. Zooplankton was collected with a net of 159 μm mesh and opening 25 × 15 cm and trawled for 2 min in each swamp on each visit. Littoral invertebrates were collected with a D-shaped pond net (25 × 23 cm) of mesh 1 mm trawled for 10 min altogether at 3 different sites in each swamp on each visit. The zooplankton collection was preserved in total in alcohol, but the littoral invertebrates were sorted in a tray and only representatives of each species retained in alcohol. Abundances of both zooplankters and littoral invertebrates were estimated on a log scale (1–2 individuals = 0.1; 3–5 = 0.3; 6–8 = 0.7; 9–20 = 1; 21–50 = 1.3, etc.). These estimations were quick and of sufficient robustness for the multivariate analysis.

Three levels of diversity were used: alpha diversity is simply the number of species instantaneous present on each collecting occasion averaged over each site within the six groups; beta diversity is the change in species diversity from one month to the next at each site; and gamma diversity is the total number of species encountered at each site through the study period.

The Pearson Correlation Coefficient in Excel was used to test a possible relationship between alpha diversity and both hydroperiod and habitat heterogeneity. It was assumed the data were linear. To test questions of faunal relationships between the six groups of swamps and flats, multivariate analyses were calculated using PRIMER (v5) (Clarke and Gorley 2001). Field and laboratory counts were already log transformed and a similarity matrix constructed based on the Bray–Curtis similarity coefficient. Non-metric multidimensional scaling (NMDS) using 999 permutations was then performed to visualise patterns in assemblage composition among the data sets. Data collected over the same time period on a nearby freshwater lake, a claypan and two saline lakes of widely different salinity ranges were added to ascertain their relationships to the treed swamps and flats.

**Results**

The treed sites, all ephemeral, varied greatly in size with the Poplar Box Flats generally larger than the Black Box swamps (Table 1). This is an artefact as all the large Black Box swamps on Bloodwood Station have been modified to hold permanent water for stock and hence not part of the study. Poplar Box flats generally were almost flat in profile, but sometimes with a distinctly deeper channel accounting for much (> 60%) of their recorded maximum depth, whereas the Black Box swamps typically were saucer shape, often again with a distinct channel but contributing less (< 40%) to their maximum depth. In the swamps and flats studied, those with channels were larger than those without (Table 1). In Black Box swamps a further irregularity was provided by randomly arranged humps of unstructured sediments, thought to be due to deltaic sedimentation in the past. In the March 2021 flooding, a new delta formed in the Washout Swamp. Particularly small sites of both types were considered separately as their hydroperiods were distinctly shorter (Table 1).

Water in all sites was fresh (< 250 μS/cm), with the larger, deeper swamps and flats concentrating more over time than the smaller or shallower sites (Table 1, SI 1). Also, pH varied though the hydroperiod, commencing about neutral and moving quickly to being moderately alkaline (i.e. ca 7.0 to ca. 8.3) (Table 1, SM 1). Water turbidities were low, always below 70 NTU and often < 30 NTU (Table 1, SM 1). The Black Box Swamps cleared somewhat during their hydroperiod, but the Poplar Box Flats increased in turbidity with time (Table 1). This is not natural as stock had access to the Poplar Box Flats on Muella Station but not the others.

Eightytwo taxa of invertebrates were encountered overall (SI 2), though 25 of these were collected fewer than three times. Few were unique to any one site or type of site, though *Eulimnadia hansomii*, *Eocyzius phytophyllus*, *Asplanchna* spp. and *Laninularia raccemovata* were restricted to Poplar Box flats. Black Box sites with channels were the most speciose,
followed by Poplar Box flats with channels, then Black Box sites without channels, Poplar Box sites without channels and the short hydroperiod sites last (Table 2). The channels add to habitat complexity and hence species diversity. This contrast in diversity patterns was evident also temporal patterns in momentary alpha diversity between the deepest sites with channels in which diversity increased with time and the shallowest nearly flat Poplar Box sites without channels in which diversity slowly decreased as they became shallower (Fig. 3a, b). Alpha diversity was significantly higher the longer the hydroperiod ($r^2 = 0.7921$, $P < 0.01$) and also was significantly influenced by habitat heterogeneity ($r^2 = 0.7583$, $P < 0.01$). Gamma diversities were also higher the longer the hydroperiod, but these were not tested statistically. Beta diversities indicated considerable changes between months for all sites. While this is influenced for the phenological change over between early dominance by large branchiopods to that by insects (see below), some is explained by rarity of many species and hence missing a record in some months.

There was a major change in communities with aging of sites (Table 4), with all sites dominated initially by large branchiopods then insects and changing slightly as species matured according to their life cycle characteristics. Zooplankton usually was dominated early and briefly by *Moina australiensis* then communities were dominated by either or both *Boeckella triarticulata* and *Daphnia carinata* s.l. Larger littoral invertebrates were sparse initially though *Triops australiensis*, larger clam shrimps (*Limnadopsis birchii*, *L. tatei* and *L. parvispinus*) soon

### Table 1 Physicochemical features of six types of treed swamps on Bloodwood and Muella Stations

| Sites                        | No. | Mean ± SE area (ha) | Mean ± SE maximum depths (cm) | Mean cond first-final (µS/cm) | Mean pH first-final | Mean turbidity first-final (NTU) |
|------------------------------|-----|---------------------|--------------------------------|------------------------------|---------------------|-------------------------------|
| Black box swamp with channels| 3   | 2.7 ± 0.4           | 90.0 ± 9.8                     | 57–250                       | 7.1–8.7             | 22–5                          |
| Black box swamps no channels | 3   | 1.2 ± 0.2           | 74.3 ± 14.2                    | 73–173                       | 7.1–8.2             | 34–14                         |
| Black box swamps small       | 2   | 0.3 ± 0.1           | 28.5 ± 3.5                     | 55–70                        | 6.8–8.4             | 25–10                         |
| Poplar box flats with channels| 3   | 7.1 ± 2.1           | 64.7 ± 11.2                    | 32–77                        | 7.4–8.3             | 45–58                         |
| Poplar box flats no channels | 4   | 6.3 ± 4.8           | 26.7 ± 4.3                     | 35–95                        | 7.0–8.4             | 39–68                         |
| Poplar box flats small       | 3   | 0.3 ± 0.1           | 23.8 ± 6.0                     | 73                            | 7.2                 | 10                            |

*These sites had water only once, therefore there is only one value for each

### Table 2 Diversity in six types of treed swamps on Bloodwood and Muella Stations

| Sites                        | Number | Alpha diversity mean ± SE | Beta diversity mean ± SE | Gamma diversity |
|------------------------------|--------|---------------------------|--------------------------|-----------------|
| Black box swamps with channels| 3      | 27.7 ± 1.3                | 18.3 ± 1.97              | 54              |
| Black box swamps no channels | 3      | 19.4 ± 0.7                | 14.0 ± 1.14              | 49              |
| Black box swamps small       | 2      | 16.0 ± 0.9                | 18.5 ± 2.35              | 32              |
| Poplar box flats with channels| 3      | 21.9 ± 1.4                | 18.6 ± 2.11              | 44              |
| Poplar box flats no channels | 4      | 14.1 ± 0.7                | 16.4 ± 2.94              | 40              |
| Poplar box flats very small  | 3      | 6.3 ± 0.3                 | (a)                       | 13              |

*No value as these flats were inundated only once, so comparisons between months not possible
were obvious as well as *Micronecta* sp. After two months these had largely disappeared to be replaced by various juvenile insects. These matured over the ensuing months, with almost no addition or replacement of species. Both the larger invertebrates and smaller planktonic invertebrates had many species considered uncommon (noted as u in SM 2) or in low numbers and not always present in one particular site or group of similar sites (marked as x in SM 2). In the main study of 2020, most branchiopods were restricted to the first sampling soon after filling and were not present at 2 months, but in 2019, when the sampling frequency was fortnightly, many persisted for 6 weeks. *Ozestheria packardi* sensu lato was unusual for a branchiopod, persisting right through the hydroperiod in some sites.

No data were collected on diets of the invertebrates, though from generalised information, most initial branchiopod colonizers were filter feeders on algae, and were replaced by more filter feeders in the form of copepods and cladocerans. Ostracods also ate algae, but by scraping or collecting them from plants and sediments. *Triops* was an early omnivore in most sites and *Eretes* larvae a prominent early carnivore in some sites. Interestingly their presence was patchy, varying in different fills (in some sites the 2019 populations

![Fig. 3](image-url)
were different from those in 2020). Detritus and partial omnivorous feeders were represented by *Micronecta*, *Agraptocorixa* and various beetle adults. Prominent carnivores towards drying were *Anisops* spp. and odonates. Strict vegetarians were uncommon, represented by limited and patchy populations of snails, though the periphyton on the plants (mainly *Chara*, also *Myriophyllum*) supported *Cloeon*, many ostracods and *Micronecta*.

An ordination of the 18 ephemeral treed swamps plus an episodic freshwater lake, a claypan, and two saline lakes as outgroups, is shown in Fig. 4. The dominant invertebrates are listed in Table 3. It suggests a close relationship between the three channelled Black Box Swamps, the three non-channelled Black Box Swamps, the three channelled Poplar Box Flats and the four non-channelled Poplar Box Flats. Thus, these four wetland types can be considered as one, the treed swamps. The smaller Black Box swamps are not far removed from this cluster, with the four small Poplar Box flats further removed. As far as is known, all species are widely distributed right across the inland, though a few branchiopods are restricted to the mid-eastern parts.

**Discussion**

This study is focused on the ecological characters that separate treed swamps from other wetland types in the middle Paroo lowlands. The physicochemical milieu and invertebrates of the saline lakes, claypans and creek pools have already been elucidated in some detail and some information is available on wetlands of lesser importance (Timms and Boulton 2001). Few other dryland wetland systems have such an array of wetland types and although invertebrate diversity is not remarkable in any one particular type, when considered together it is impressive, particularly for large branchiopods (eg. Timms 2015; Schwentner et al. 2015). Moreover besides the major environmental differences between wetland types, minor variations in habitat complexity contribute to invertebrate diversity, as explored in these treed swamps. This is a subject of many examples both in sets of wetlands in remote drylands (e.g. Hancock and Timms 2002; Timms 1998b) as well as in more accessible Mediterranean wetlands (e.g. Boix et al. 2016; Florencio et al. 2014a).

Over 34 years of casual monitoring at Bloodwood, the 2020 flooding was the greatest ever observed since white settlement, while the 2019 filling, though not measured, was more ‘average’ at about half the
inundated area measured in 2020. The great flooding of 2020 was akin to that in 1974–1976 (D. Leigo, pers. com.) and going back in the wider Paroo to 1885 of great floodings of Lake Wyara in 1890, 1959–1950 and 1974–1976 (A. McGrath pers. com; Timms 1998b) indicate a return interval for remarkable flooding rains of about 45–50 years.

In inland eastern Australia a major climate driver is the El-Nino – La Nina system of the Pacific Ocean. La Nina years are wetter and cooler and presently occur on a 10–11 year cycle (BOM 2021). Larger dryland lakes such as Lake Wyara in the southern Queensland Paroo flood and dry cyclically approximately every 10 years, the correlation coefficient between the Southern Oscillation Index and full and dry periods 1887–1994 is highly significant ($r = 0.622$, $n = 34$) (Bridgman and Timms 2012). Such a dominant driver also affects the treed swamps.

In general, Black Box swamps flood every 2–3 years, always in a La Nina climatic year and perhaps weakly two or rarely three times in between or two years in succession if La Nina events last two years. Smaller black box swamps such as Marsilea swamp floods to a maximum ca 30 cm depth only in La Nina years, with other wettings 5–10 cm deep and lasting only a few days to weeks. Poplar Box flats flood less frequently at about 3–5 year intervals, again in the same temporal pattern. In strong El Nino years, all swamps remain dry for years, perhaps up to 6 years, as in the Millennium drought of 2001–2006.

There are vast differences in habitat characteristics and faunal composition between these treed swamps

| Table 3 Phenology of dominant invertebrates in treed swamps on Bloodwood and Muella |
|---------------------------------|-----------------|-----------------|
| Two weeks | Two months | Four to six months |
| A. Poplar box flats | | |
| **Open waters** | | |
| *Moina australiensis* | A cyclopoid | A cyclopoid |
| *Branchinella spp.* | *Daphnia carinata* | rotifers |
| *Paralimnadia or Eulimnadia* | | |
| **Littoral** | | |
| *Micronecta sp.* | *Anisops spp. juv.* | *Anisops spp. adults* |
| *Agraptocorixa juv.* | *Agraptocorixa adults* | mature Odonata |
| *Triops australiensis* | Odonata juv. | |
| *Berosus larvae* | adult dytiscids | |
| *Eretes larvae* | | |
| *Sternopriscus adults* | | |
| *Cloeon larvae* | | |
| A mosquito | | |
| A. Black box swamps | | |
| **Open waters** | | |
| *Moina australiensis* | *Boeckella triarticulata* | *Boeckella triarticulata* |
| *Branchinella spp.* | *Daphnia carinata* | *Calamoecia lucasi* |
| *Limnadopsis spp.* | various ostracods | *Daphnia carinata* |
| *Paralimnadia queenslandicus* | *Ozestheria ‘packardi’* | |
| **Ozestheria spp.** | | |
| **Littoral** | | |
| *Triops australiensis* | *Anisops juv.* | *Anisops adults* |
| *Micronecta sp.* | *Agraptocorixa juv.* | *Agraptocorixa adults* |
| Clam shrimps as above | *Cloeon larvae* | mature *Cloeon larvae* |
| *Eretes larvae* | *Triplectides larvae* | mature *Triplectides larvae* |
| *Berosus larvae* | dytiscid adults | chironomid larvae |
| *Sternopriscus adults* | Odonata juv. | Mature Odonata |


and salt lakes and claypans, the other common wetlands of the Paroo (Fig. 4, Table 4; Timms and Boulton 2001). The salt lakes are most different: their saline waters have entirely different invertebrates including the ostracods *Diacypris* spp., *Mytilocypris splendidida* and *Trigonocypris globulosa,* the cladoceran *Daphniopsis queenslandensis* and the anostracan *Parartemia minuta* (Timms 1993, 2018). Claypans share some features with the treed swamps, mainly some insects and *Triops australiensis,* but their high turbidity is suitable for a range of branchiopods, such as *Branchinella affinis,* *B. budjiti,* *B. lyrifera* and *B. occidentalis* as well as spiniacaudatans *Ozestheria* spp. and *Eocyzicus argillaquus* (Hancock and Timms 2002). On the other hand, there are many similarities in habitat and fauna between these treed swamps and episodic freshwater lakes and creek pools (Fig. 3, Table 4; Timms and Boulton 2001). Paroo treed swamps are not as speciose as local freshwater lakes (Timms and Boulton 2001) and creek pools (Timms 2001), though the majority of invertebrates listed for the treed swamps also occur in the freshwater lakes and creek pools. The ecological drivers suggested for all Paroo wetlands also apply specifically to these treed swamps, namely water fresh (EC ≤ 250 μS/cm) (as against saline waters of the salt lakes and samphire lakes), water turbidity low (< 50 NTU) (as against high turbidity in claypans and crystal-clear saline waters and somewhat turbid waters of other fresh waters), and relatively short hydroperiods (< 6 months) (applies to many Paroo habitat types but some are permanent or episodic persisting a year or more).

In those treed wetlands with channels and the longest hydroperiods there is an increase in diversity with time and in all others a decrease (Figs. 3a, b). Similar trends have also been noticed in the Doñana wetlands with longer hydroperiods in Spain (Florenço et al. 2011), but local claypans exhibit a marked decrease in diversity with wetland wetting aging (Timms 2002). Explanations vary, but in the channelled tree swamps there may be addition of species from other drying wetlands to a habitat resource still available in the channels, or to more efficient collecting in the easier field conditions. For the simpler tree swamps or flats without channels, and the claypans, the less heterogenous environment as the wetland dries is a likely explanation.

Among the treed swamps and flats, it seems the trees themselves are not important in their ecology throughout a filling-drying cycle. Only in some floodings are the peripheral tree bases flooded, but when they do, some evidence suggests diversity is enhanced. This was seen occasionally by increased

| Table 4 Prominent invertebrates in five wetland types in the Paroo |
|---------------------|-------------------|-------------------|-------------------|-------------------|
| Treed swamps<sup>a</sup> | Salt lakes<sup>b</sup> | Claypans<sup>c</sup> | Episodic freshwaters<sup>d</sup> | Creek pools<sup>e</sup> |
| **Branchinella frondosa** | **Parartemia minuta** | **Branchinella lyrifera** | **Branchinella australiensis** | **Branchinella australiensis** |
| **Limnadopsis** spp. | **Daphniopsis queenslandicus** | **Branchinella affinis** | **Paralimnadia** spp. | **Limnadopsis** spp. |
| **Boeckella triarticulata** | **Trigonocypris globulosus** | **Ozestheria lutraria** | **Ozestheria parkardi** | **Boeckella triarticulata** |
| **Daphnia carinata** | **Cyprinotus edwardsi** | **Ozestheria packardi** | **Boeckella triarticulata** | **Ozestheria packardi** |
| **Micronecta** sp. | **Diacypris** sp. | **Ozestheria packardi** | **Daphnia carinata** | **Micronecta** sp. |
| **Anisops** spp. | **Micronecta** sp. | **Anisops stali** | **Anisops** spp. | **Anisops** spp. |
| **Odonates** | **Anisops** spp. | **Eretes australis** | **Dytiscid beetles** | **Dystiscid beetles** |
| **Dytiscid beetles** | **Tanytarsus barbitarsus** | **Other dytiscids** | **Physa acuta** | **Physa acuta** |

<sup>a</sup>Appendix SI2

<sup>b</sup>Timms (1993, 1998a, b, 2018)

<sup>c</sup>Hancock and Timms (2002), Timms (2002)

<sup>d</sup>Timms (1997a, b, 2001b)

<sup>e</sup>Timms (2001)
diversity of spinicaudatans (*Limnadopsis* spp., *Paralinadia queenslandicus*, *Eocyzius* spp.) experienced in those swamps most flooded in 2020 and also in previous floodings at Far Homestead Black Box swamp and Carters Poplar Box flat when some uncommon species were caught in 2000 and 2011 floodings (e.g. *Eoleptestheria* sp in 2000 and *Cycles-theria* nr *hislopi* in 2011) (Timms, 2009; Schwentner et al., 2013). Otherwise, the trees contribute woody detritus to the wetlands and this enhances species diversity, probably not as a food source, but as secure hiding places. Leaf litter from the trees was observed mainly in dry times and hence is processed terrestrially and does not contribute directly to the aquatic system. Hydrological period is also a contributor to diversity in these swamps and flats (Table 2); this is a common ecological driver in dryland wetlands (e.g. Florencio et al. 2011; Timms and Boulton 2001; Davis et al. 2018).

In a more comprehensive study of freshwater aquatic habitats across the Australian arid and semi-arid zones but excluding any sites in the Murray Darling basin, Davis et al. (2018) noted 10 types of habitats and major ecological drivers of latitude, connectivity, hydroperiod and biogeographic history, i.e. relic fauna in some isolated sites. Each of the drivers varied in importance with habitat type: latitude applied to most, connectivity to the riverine systems and isolation to desert rockholes, some waters were permanent and hydrologically stable, others temporary and variable. These authors did not look at the influence of water chemistry or related parameters, but had they done so, then (a) salinity as an ecological driver would have featured, as it does in the Paroo, (b) turbidity featured only for the uncommon claypan sites across the arid inland whereas in the Paroo these are common and there are minor variations in water clarity in other Paroo habitat types, and (c) hydrological variability is arguably the most important driver everywhere.

The factor of hydrological variability is shared with Mediterranean wetlands, though in these there is seasonal reliability (Boix et al. 2016), important for at least some niche separation of species, and hence increased diversity. Perceived diversity in the Paroo is artificially low because some groups have hardly been studied (rotifers) or have probably been incompletely collected (e.g. chironomids). However, this is not enough to account for the lower species richness in the Paroo treed pools when compared to the generally speciose Mediterranean pools or the temporary wetland ponds of temperate regions—gamma diversity > 100 species (Boix et al. 2016; Lake et al. 1989; Pinder et al. 2004; Jeffries et al. 2016). The reason is probably largely due to the unreliable filling of the semiarid Paroo treed swamps, but short hydroperiods and simpler habitat structure may contribute (Florenceo et al. 2014a, b). These factors probably also contribute to the lack of any detailed seasonal succession, except those determined by phylogenetic characteristics (e.g. branchiopods early in succession, predatory insects including odonatans maturing late).

Treed swamps also occur in the wheatbelt of Western Australia, including Southern Yate (*Eucalyptus occidentalis*) and *Melaleuca* spp. swamps. Such sites generally have abundant vegetation of sedges and a greater diversity of invertebrate assemblages (Pinder et al. 2004) but fewer branchiopods (author unpublished data) compared with Paroo treed swamps. Gross differences could be explained by lower vegetation density and greater environmental harshness of the Paroo treed swamps. Phenological sequences in the Western Australian swamps have not been explored.

The most similar study of a treed swamp in a semiarid climate is by Lahr et al. (1999) of a sahelian temporary pond in Senegal. It had an eight-month hydroperiod and similar fluctuations over time in conductivity, turbidity and pH as in Paroo treed swamps. Its fauna was dominated early by crustaceans, mainly branchiopods, then later by insects, particularly hemipterans and dytiscids, but with some specific separation in abundances perhaps explained by its longer hydroperiod. Striking similarities include the early dominance of the zooplanker *Moina*, the early abundance of two species of *Anostraca*, and the later dominance of the insects *Micronecta* and *Anisops* and *Eretes*. Significant differences include the variety of clam shrimps and ostracods in the Paroo sites in keeping with their high diversity in inland Australia (Schwentner et al. 2015; De Deckker 1983; Halse and Martens 2019), and the greater importance of some lesser insect groups (odonatans, ephemeropterans and trichopterans) in Paroo sites.

Perhaps the greatest unknown of these treed swamps is the ecological role of the trees which define them. So far it seems they are a collateral presence only contributing during super full phases,
and then only to minor extent. Nutrient flows have not been studied, but where they have in nearby claypans, nutrients were maximal when first full and then soon become depleted (Hancock and Timms 2002). If similar in these treed swamps, such changes would account for the abundance of branchiopods in the first flush and the general scarcity of insects later on. Maybe a nutrient problem could contribute via the paucity of invertebrates, except initially, and of plants to the limited use of waterbirds of treed swamps, compared to their abundance in the saline lakes (Kingsford 1999). Given the abundance of wetlands in the middle Paroo (Kingsford 1999) and hence possibilities for species exchanges, dispersal of branchiopods and insects to these wetlands does not seem to be a problem to rebuild invertebrate populations on re-flooding, thus providing a future comparative study where dryland wetlands are scarce.

Management

These treed swamps are unmanaged on Bloodwood, Tredega and Muella as elsewhere in the Paroo, though some larger Black Box swamps have long been much modified ecologically by the excavation of dams in them. Many decades of sheep grazing are believed not to have affected the wetlands as sheep keep out water, but same cannot be said for more recent cattle grazing as cattle pug the wet surfaces. The ecological effect of this is unknown, but thought to be minor.

Climate change predictions for the next few decades for the eastern inland of Australia suggest an increase of temperatures by a few degrees, and, though a 10–20% decrease in annual rainfall, an increase in intensity of summer rainfalls and their unreliability (CSIRO 2021; Huang et al. 2017; IPCC 2014). Neither should influence the robust aquatic fauna of the Paroo. Experiments by Sives (pers. com.) in the hatching of branchiopod eggs from some Paroo wetlands found little change with increased temperatures and the aquatic faunas already successfully deal with the unreliable rainfall. It is possible, however, that seasonal changes in fauna may change with insects more common in winter than they are now (Timms 2018). Given the present rarity of water still present in the treed swamps in winter, response to increase in winter temperatures would be minor. What is already seen, however, is some tropical taxa, such as Cyclestheria moving southwards (Schwentner et al. 2013). Maybe a few other tropical anostracans and spinicaudatans will also move south. Huang et al. 2017

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Data availability

The detailed datasets generated during this study, besides that reported in the paper and appendices, are available from the author on a reasonable request.

Declarations

Conflict of interest

The author declares that he has have no conflicts of interest.

Ethical approval

No ethics approval was necessary to study invertebrates in the jurisdiction of the field sites.

Informed consent

As noted in the acknowledgements I had permission from the owners to study the treed swamps on their lands. As a retiree under my own auspices I do not need consent for publication.

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