Pilbara stygofauna: deep groundwater of an arid landscape contains globally significant radiation of biodiversity

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Abstract – The Pilbara region was surveyed for stygofauna between 2002 and 2005 with the aims of setting nature conservation priorities in relation to stygofauna, improving the understanding of factors affecting invertebrate stygofauna distribution and sampling yields, and providing a framework for assessing stygofauna species and community significance in the environmental impact assessment process.

Approximately 350 species of stygofauna were collected during the survey and extrapolation suggests that 500–550 actually occur in the Pilbara, although taxonomic resolution among some groups of stygofauna is poor and species richness is likely to have been substantially underestimated. More than 50 species were found in a single bore. Even though species richness was underestimated, it is clear that the Pilbara is a globally important region for stygofauna, supporting species densities greater than anywhere other than the Dinaric karst of Europe. This is in part because of a remarkable radiation of candonid ostracods in the Pilbara. Ostracods are the dominant stygofaunal group in terms of both species richness and animal abundance. Together, ostracods, copepods, amphipods and oligochates comprised 77% of species and 96% of animals collected.

Stygofauna were found in 72% of samples collected and 81% of wells sampled. The average sample (including those without stygofauna) contained 3.2 ± 0.1 species. A feature of the Pilbara is that stygofauna occur across most of the landscape, often where the depth to groundwater is considerable, although yields were low where depth to groundwater was >30 m. Another feature is high endemicity: on the basis of current taxonomy 98% of the stygobites and 83% of the other groundwater species occur only within the region. Few factors affecting stygofauna occurrence could be identified, however. Numbers of specimens and species collected were positively related to well diameter and negatively related to depth to groundwater. Numbers of species declined in inland sub-regions, although variability within sub-regions was high.

While a range of freshwater chemistries occurred, 79% of water samples were weakly saline and NaCl dominated. Profiling and purging of wells suggested that water quality measurements reflected aquifer conditions in most situations. Water chemistry appeared to have limited influence on stygofauna occurrence in the Pilbara. Geology also appeared to have little effect on stygofauna occurrence but this may have been the result of non-random siting of wells; there was a bias towards wells being in transmissive locations that were not necessarily typical of the geology in which each well occurred.

No potential reserves for stygofauna are recommended in this paper but nine areas of high stygofauna richness were identified, including the listed Ethel Gorge stygofauna community. Theoretical analysis of species ranges suggested that half of the species found only in the vicinity of development projects will have ranges less than 680 km². Consequently, projects involving extensive groundwater drawdown (sometimes through the interaction of de-watering operations at adjacent projects) have the potential to affect a large proportion of the population of a restricted species or to threaten the persistence of species with particularly small ranges.

Keywords – subterranean fauna, stygobite, sampling, groundwater, distributions, Western Australia
INTRODUCTION

The global volume of groundwater is approximately 30 times greater than the volume of non-marine surface water (Gleick 1993). This disparity is even greater in arid regions, such as the Pilbara of Western Australia, where there is relatively little surface water but vast amounts of fresh groundwater (Johnson and Wright 2001). Consequently, groundwater comprises by volume almost all of the aquatic habitat of the Pilbara, although surface claypans, river pools and springs are highly biodiverse (Pinder et al. 2010).

Some surface-water animals make use of groundwater where it discharges from springs or seeps into river pools through the streambed. However, animals using groundwater at greater depths are referred to as stygofauna and most show some modifications to subterranean life and the absence of light. Classically, the adaptations include loss of eyes and pigment, elongation of appendages and sensory structures, and a veriform body shape (Christiansen 1961). Nearly all stygofauna are invertebrates (Gibert et al. 1994) and they are often classified into three broad categories according to their dependence on groundwater (Camacho 1992; Sket 2008). Stygobites spend their full life cycle in groundwater; stygophiles either have a life stage in epigean habitats or some of their populations occur in surface water; stygoxenes are facultative users of groundwater that are found mostly in epigean habitats.

The early history of stygofauna research in Western Australia is similar to that in other parts of the world, with the species discovered prior to the 1990s mostly being collected by speleologists in caves, or from wells in highly karstic environments (e.g. Holthuis 1960; Whitely 1945). However, the direction of research changed in the early 1990s, when W.F. Humphreys began extensive survey work at the Cape Range, Barrow Island and then in the Pilbara region (Adams and Humphreys 1993; Humphreys 2000, 2001a, 2008). This pioneering work, which mostly involved sampling pastoral wells, showed that the Pilbara of Western Australia contains a rich, diverse array of stygofauna species across the landscape matrix rather than in caves (Eberhard et al. 2005a). It was found that relictual species from both previous Tethyan (e.g. Poore and Humphreys 1992, 1998) and Gondwanan (e.g. Knott and Halse 1999; Karanovic, T. 2006) alignments of the Pilbara persist in the region’s current stygofauna communities. In addition, the Pilbara stygofauna community contains some highly distinctive and endemic elements (e.g. Karanovic, I. 2007).

Since the late 1990s, there has been growing awareness that some species in the rich Pilbara stygofauna communities may be threatened by mining developments. Western Australia is historically the third largest producer of iron ore (17% of global production) after China (25%) and Brazil (20%) (DMPR 2001), with substantial increases in production underway. More than 95% of Western Australian iron ore is extracted from the Pilbara in large, open-pit mining operations that usually require removal of groundwater (de-watering) to provide access to ore below the water table (Johnson and Wright 2001). De-watering may result in groundwater drawdown of several hundred metres (e.g. EPA 2009). This threat to stygofauna has been recognised by the Environmental Protection Authority and a requirement to consider impacts on stygofauna is incorporated into the environmental assessment process for new mines and other developments affecting groundwater (EPA 2003, 2013).

Evaluating the likely impact of a development on stygofauna is much easier if the distribution of stygofauna species and the factors affecting their occurrence are known. Accordingly, stygofauna were included as a survey element of the Pilbara Biodiversity Survey (McKenzie et al. 2009). As a program, the Pilbara Biodiversity Survey had the following objectives:

1. To collect systematic baseline data on the current distribution of biota across the region to provide a regional perspective on nature conservation priorities and a framework for future monitoring of regional-scale trends in occurrence.

2. To improve understanding of the factors influencing plant and animal distributions and to provide a context for assessing the conservation status of species and communities and the significance of localised occurrences of species.

3. To identify gaps in the coverage of the reserve system for species and communities and to identify areas where reservation will efficiently improve the reserve system according to three criteria: comprehensiveness (i.e. all regional-scale ecosystems included), adequacy (i.e. sufficient areas reserved to be viable) and representativeness (i.e. fine scale variability within ecosystems incorporated into reserves).

The stygofauna survey program focused on the first two objectives of the Pilbara Biodiversity Survey (collection of baseline data and providing context for assessment). At this stage, the third objective (improving the reserve system) is difficult to achieve for stygofauna, partly because stygofauna can be sampled only after wells or drill...
holes are in place. These are most abundant in areas intended for development.

The survey was directed at invertebrate stygofauna, which were the only stygofauna species known from the Pilbara when the survey was planned. Subsequently the blind eel, *Ophisternon candidum*, was collected near Bungaroo Creek by Biota Environmental Sciences (EPA 2012), although this record has not been formally published and evaluated. The blind eel also occurs in the Cape Range, together with the blind cave gudgeon, *Milyeringa veritas* (Humphreys 2001a). A species of blind eel and another species of blind gudgeon, *M. justitia*, have been collected from Barrow Island off the Pilbara coast (Larson et al. 2013).

**METHODS**

**Study area**

For the purposes of the stygofauna survey, the Pilbara region was considered to be bounded to the south by the main channel of the Ashburton River and to the north-east by the De Grey/Oakover River and its tributaries (Figure 1). This area of about 261,144 km² includes all the Pilbara region as recognised in the Interim Biogeographic Regionalisation for Australia (IBRA) and parts of the Augustus and Ashburton subregions of the northern Gascoyne IBRA region (Environment Australia 2008). McKenzie et al. (2009) provide a summary of the region’s landforms, geology and climate.

Figure 1 Surface geology of the Pilbara, stygofauna survey boundary and selected locations.
The Pilbara is hot and dry, with the annual rainfall in most areas (250–300 mm, decreasing inland) outweighed 10:1 by evaporation. Most rainfall occurs in monsoonal thunderstorm or cyclonic events during summer, although winter rainfall is sometimes significant in coastal areas. Mean summer minimum and maximum temperatures are 25°C and 36°C, respectively, while mean winter temperatures are 12°C and 27°C.

The Pilbara Craton, one of Australia’s major geological blocks (Geological Survey of Western Australia 1990), largely coincides with the Pilbara region. The Craton is characterised by hard rock landscapes that were laid down in Archaean times, and part of the Craton formed the earliest known emergent landmass about 3.5 billion years ago (Buick et al. 1995). The Archaean basement rock is exposed as granite and greenstone terrain in the northern Pilbara but is overlain by rugged sedimentary strata, volcanic flows and lateritised caps in the southern Pilbara. Importantly from a biogeographic viewpoint, most of the Craton has been emergent throughout the earth’s history of continent formation and break-up (Johnson 2004).

There is little permanent surface water in the Pilbara and all rivers are ephemeral (Pinder et al. 2010). Nevertheless, river systems are extensive, with five drainage basins recognised: the Ashburton, Robe, Fortescue, Port Hedland Coast and De Grey (Figure 1). These basins mostly contain three types of groundwater aquifers that potentially provide habitat for stygofauna: (1) unconsolidated sedimentary aquifers, including those in recent valley-fill alluvium and colluvium, and coastal deposits; (2) aquifers in chemically-deposited calcrites and pisoliths within Tertiary drainage channels; and (3) fractured-rock aquifers in dolomites, banded-iron formations (BIF), granite and other rocks (Johnson & Wright 2001). Aquifers in calcrete and pisolite are often associated with alluvial aquifers and alluvium is a prominent feature of all drainage channels.

Groundwater in the Pilbara is mostly fresh (200–1500 mg L⁻¹) and often bicarbonate-dominated, although NaCl-rich waters are common in both the coastal and arid eastern margins. Water under the Fortescue Marsh in the central part of the Fortescue basin is hypersaline and dominated by NaCl. Recharge to the groundwater aquifers is principally via cyclonic rain (Dogramaci et al. 2012) but there is also some seepage during peak flow periods from rivers and creeks to alluvial aquifers. Depth to groundwater is highly variable, with groundwater discharging to the surface at springs and in the beds of many large river pools, although the latter is usually water stored in the channel alluvium rather than groundwater from regional aquifers (Fellman et al. 2011). Away from watercourses, groundwater is mostly 5–20 m below ground level (bgl) across the coastal plain and often more than 40 m bgl in the Hamersley and Chichester Ranges.

A fuller account of groundwater conditions is provided by Reeves et al. (2007).

The occurrence of stygofauna in an aquifer is regarded as being controlled largely by the types of voids and interstitial spaces present, and by groundwater chemistry (Danielopol et al. 2003). Both these attributes are influenced by the host geology of the aquifer, the amount of landscape weathering, and local chemical and hydrological processes (Reeves et al. 2007).

Survey design and methods

Stygofauna survey began in 2002 and continued until 2005. A total of 1053 samples was collected from 507 wells and drill holes across the Pilbara that intersected groundwater (Figure 2, Appendix 1). These holes were mostly wells installed by government agencies, pastoralists or mining companies. Thirty-six wells were sampled once, 441 were sampled twice, and 30 wells were sampled more frequently to examine the efficiency of sampling methods (see Eberhard et al. 2009) or in an attempt to collect particular species.

Eleven ‘sub-regions’ across the Pilbara were identified for the purpose of ensuring stygofauna sampling effort was geographically well spread (Figure 2). The sub-regions were based on the major river basins, although the Robe basin was combined with the Fortescue to reflect the past connection of these basins. The diversion of the lower Fortescue away from the Robe River most likely occurred since the Last Glacial Maximum (Barnett and Commander 1985). The identification of sub-regions within the basins was based on distance from the coast, topography and geology. Insofar as possible, sampling effort was distributed evenly across the sub-regions (65–109 samples in each). In addition to the 11 sub-regions, small areas to the north (Eighty Mile Beach), east (Great Sandy Desert), south (north-east Gascoyne) and south-west (Carnarvon) were sampled to provide some biogeographic context for the results of the Pilbara sampling. Twelve to 27 samples were collected from each of these areas.

While an attempt was made to include as many different geologies as possible in the survey, the use of existing wells biased the sampling towards aquifers that yielded high volumes of fresh water. Consequently, more samples were collected from alluvium, colluvium and calcrite than would be expected from the spatial occurrence of these geologies in the Pilbara.

Wells

Nearly all the wells sampled for stygofauna were cased and slotted because uncased wells and drill holes caused frequent loss of equipment.
Wells ranged in diameter from 50 mm to about 2 m. The larger wells were mostly older, located on shallow watertables, hand-dug and lined with rock. Narrower wells were cased either with polyvinyl chloride (PVC) or steel and slotted below the watertable in a variety of ways to enable exchange of water between the bore and the surrounding aquifer. A high proportion of the wells sampled were either monitoring or production wells and were slotted for most of their underwater length. A small proportion were piezometers and, thus, were slotted at particular depths. Details of slotting and the screens used in well construction were usually not available and no attempt was made to evaluate whether slot and screen sizes affected faunal yields, other than noting that large animals were frequently present in wells with standard (0.5–1 mm) slotting. All the wells sampled were more than 6 months old and most were capped (i.e. the casing or a collar at the top of the well was sealed at the surface to prevent animals and material falling into the well).

**Sampling methods**

Prior to fauna sampling, electrical conductivity (EC), pH, redox and dissolved oxygen (DO) were measured in each well using a Yeo-Kal 611 water quality analyser lowered to 1 m below the watertable (referred to as standing water level, SWL). Depths to SWL and the base of the well were measured with a Richter Electronic Depth Gauge or weighted Lufkin tape measure.

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**Figure 2** Locations of wells sampled across the Pilbara, with sampling sub-regions outlined (see text for explanation of sub-regions).
Water was also collected from 1 m below SWL using a sterile bailer (Clearwater PVC disposable 38 × 914 mm) and a 250 ml sample was collected for analysis of total dissolved solids (TDS), ionic composition, alkalinity, colour and turbidity. A second 125 ml sample was collected, and frozen in the field, for analysis of total soluble nitrogen, nitrate/nitrite, total soluble phosphorous and soluble reactive phosphorous by ChemCentre using the laboratory methods of APHA (1995). All analyses occurred within one month of collection.

Stygofauna were collected with small haul nets, made from either 50 or 150 μm mesh, with a glass collecting vial at the base within a brass weight (Eberhard et al. 2005b). The base of the glass vial was removed and replaced with 50 μm mesh to improve water flow through the net. At each well, a stygofauna sample was collected by lowering the net to the end of the well, bouncing the net several times to stir up sediment and slowly retrieving it. The contents of the vial were then transferred into 100% ethanol. Three hauls with the 50 μm mesh net and three hauls with the 150 μm mesh net were made for each sample.

After each sampling, nets were washed in Decon90, rinsed in deionised water and air-dried to prevent the transfer of stygofauna between bores during the survey.

**Stygofauna sample processing and identification**

Prior to sorting under a dissecting microscope, samples were separated in the laboratory into three size fractions by sieving through 250, 90 and 53 mm Endecott sieves to facilitate searching for preserved stygofauna. All stygofauna taxa collected were identified to the lowest taxonomic rank possible using published and informal keys, the aim being to achieve species or morpho-species identification. Where necessary, animals were dissected and examined under a compound microscope to achieve identification. The numbers of individuals of each taxon present were recorded using a logarithmic scale (1 = 1–10 animals, 2 = 11–100 animals etc.) and adjusted to midpoint log values of 0.7, 1.7 etc. for analyses involving abundance.

As part of the survey, and to facilitate identification, taxonomic work was instigated on copepods (Karanovic, I. 2006; Tang et al. 2008), ostracods (Karanovic, I. 2007) and isopods (Keable and Wilson 2006; Bruce 2008) and water mites (M.S. Harvey unpublished). Existing work on amphipod taxonomy was supported (Finston et al. 2007, 2011).

**Profiling and purging wells**

The correspondence between the physicochemical data collected 1 m below SWL in wells and the surrounding aquifer was investigated in 34 bores in the western half of the Pilbara during 2005 and 2006. Approximately half of the wells were investigated each year (Table 1). Three types of data were collected:

1. Measurements of EC, pH, temperature and DO were made at approximately 1m intervals for the length of the water column in all wells. The purpose of this profiling was to determine whether measurements made 1 m below SWL were representative of conditions throughout the water column.

2. Data on EC, pH, ionic composition and nutrients were collected from six wells 1 m below SWL before and after these bores were purged. Purging consisted of pumping out three times the volume of water occurring in the well so that the water column was entirely replaced by water from the surrounding aquifer. The purpose of these measurements was to determine whether water in the well was representative of water in the surrounding aquifer or whether atypical conditions exist within the well that may have favoured a few stygofauna species.

3. In 2005 profiling was repeated after an interval of four months at three wells that exhibited marked vertical variation in their profiles to determine whether profiles were stable across seasons. The profiles of two wells were re-examined in 2006.

**Data analysis**

Unless stated otherwise, all analyses were based on the first two samples taken from each of 471 wells, plus the single samples from 36 wells. This was done to prevent analyses being biased by wells with unusual characteristics that were sampled on multiple occasions (Eberhard et al. 2009).

**Water chemistry**

Water samples were classified based on values of the parameters EC, pH, and major ions (Na, K, Ca, Mg, Mn, Cl, HCO$_3$–CO$_3$, SO$_4$ expressed in milliequivalents L$^{-1}$). Samples with more than 15% difference between the milliequivalent sums of cations and anions were excluded from analysis (APHA 1995). Parameters were range standardised and dissimilarity between samples was calculated using the Gower metric. The UPGMA option with default settings in the PATN multivariate analysis package (v. 3.12, http://www.patn.com.au) was used to perform the classification. Results were examined in an ordination plot using the SSH procedure and the PCC option was used to show correlations of the water chemistry parameters with ordination space.

Differences in water chemistry among the
Table 1  Wells in which water chemistry profiles, or chemical composition before and after purging, were measured. WC, length of water column; N, number of occasions profiled; R = profiled, U = purged.

| Code          | Name               | Basin      | Coordinates                  | Diam. (mm) | Casing  | Depth (m) | WC (m) | N  | Type |
|---------------|--------------------|------------|------------------------------|------------|---------|-----------|--------|----|------|
| Karratha2     | Boundary Well      | Fortescue  | 20°30'51" S, 119°54'42"    | 1200       | Concrete| 6         | 4      | 1  | R    |
| NWSLK220A     | Hardey River       | Ashburton  | 23°21'00" S, 117°49'32"    | 150        | PVC     | 17        | 13     | 1  | R    |
| NWSLK220B     | Hardey River       | Ashburton  | 23°20'52" S, 117°47'56"    | 150        | PVC     | 17        | 13     | 1  | R    |
| Pyramid10     | Cup of Tea Well    | Port Hedland Coastal | 21°13'18" S, 116°06'30" | 1000       | Steel   | 10        | 4      | 1  | R    |
| Pyramid11     | Middle Well        | Port Hedland Coastal | 24°57'10" S, 119°24'44" | 1000       | None    | 9         | 5      | 1  | R    |
| Pyramid6      | Minson Well        | Port Hedland Coastal | 21°57'06" S, 119°39'28" | 1000       | PVC     | 8         | 2      | 1  | R    |
| Pyramid8      | Johannie Walker    | Port Hedland Coastal | 22°09'08" S, 119°31'49" | 1000       | PVC     | 14        | 8      | 1  | R    |
| T176          | Coon Siding        | Port Hedland Coastal | 21°05'49" S, 119°21'59" | 150        | None    | 37        | 26     | 1  | R    |
| T243          | Cowra Siding       | Fortescue  | 21°19'51" S, 120°22'32"    | 150        | Unknown | 48        | 40     | 2  | R    |
| T255          | Cowra Siding 2     | Fortescue  | 21°40'44" S, 115°21'42"    | 150        | Steel   | 42        | 38     | 3  | R    |
| T274A         | Gidgi Siding       | Fortescue  | 21°19'51" S, 120°22'32"    | 150        | Unknown | 41        | 31     | 2  | R    |
| T359B         | Mindy Siding       | Fortescue  | 22°25'31" S, 117°18'38"    | 150        | PVC     | 40        | 15     | 2  | R    |
| T90           | Gillam Siding      | Port Hedland Coastal | 20°56'45" S, 117°37'49" | 150        | PVC     | 37        | 30     | 1  | R    |
| 126/4         | Ragged Hills Mine  | Great Sandy Desert | 22°01'52" S, 116°06'19" | 150        | None    | 35        | 33     | 1  | R    |
| COOL1         | Nr Tampathanna Pool| Fortescue  | 22°32'10" S, 120°9'22"     | 150        | PVC     | 11        | 10     | 1  | R    |
| G70830104     | Fortescue 3A       | Fortescue  | 22°48'49.2" S, 115°24'13.3" | 50        | PVC     | 20        | 14     | 1  | R, U |
| Code       | Name                  | Basin            | Coordinates                  | Diam. (mm) | Casing     | Depth (m) | WC (m) | N | Type |
|------------|-----------------------|------------------|------------------------------|------------|------------|-----------|--------|---|------|
| GSORC148   | Woodie Woodie Mine    | Great Sandy Desert | 22°07'11.3" S, 116°03'53.4" E | 50         | Concrete   | 29        | 10     | 1 | R    |
| MBSLK356A  | Carlinde Station      | De Grey          | 21°19'24.8" S, 117°52'47.1" E | 150        | Concrete   | 32        | 20     | 1 | R    |
| NWSLK304   | Warp2                 | Ashburton        | 21°49'03.1" S, 116°42'29.7" E | 150        | Unknown    | 85        | 73     | 1 | R    |
| OnslowSLK8 | MINNIE 2              | Robe             | 21°25'37.8" S, 118°28'27.3" E | 150        | Unknown    | 34        | 26     | 1 | R    |
| PannaSLK24 | Yarraloola Well       | Robe             | 21°25'37.8" S, 118°28'27.3" E | 150        | Unknown    | 38        | 23     | 1 | R    |
| PannaSLK4B | Yarraloola Station    | Robe             | 22°28'48.0" S, 116°28'16.3" E | 150        | Unknown    | 22        | 17     | 1 | R    |
| PSPRSLK48  | Seven Mile Creek      | Ashburton        | 22°47'10.8" S, 114°58'02.5" E | 150        | Concrete   | 42        | 41     | 1 | R    |
| ROY HILL 1 | Tuccamunna            | Fortescue        | 22°14'51.3" S, 116°08'01.3" E | 150        | PVC        | 25        | 21     | 1 | R    |
| RWSLK6     | South Fortescue Marsh | Fortescue        | 22°47'10.8" S, 114°58'02.5" E | 150        | Concrete   | 34        | 33     | 1 | R    |
| W260       | At Production Bore K31 | Fortescue     | 22°14'51.3" S, 116°08'01.3" E | 50         | PVC        | 22        | 20     | 1 | R    |
| WW8-4      | RHR Road              | De Grey          | 22°01'51.6" S, 116°06'19.1" E | 150        | None       | 17        | 15     | 1 | R    |
| BH15       | Weeli Wolli           | Fortescue        | 22°56'29.9" S, 119°09'58.9" E | 50         | PVC        | 45        | 37     | 1 | R    |
| BH18D      | Weeli Wolli           | Fortescue        | 22°55'27.8" S, 119°11'49" E  | 50         | PVC        | 70        | 60     | 1 | R    |
| G70730101  | Robe 1A               | Robe             | 21°34'31.5" S, 115°52'57.6" E | 75         | PVC        | 23        | 17     | P | U    |
| G70730103  | Robe 3A               | Robe             | 21°32'58.8" S, 115°51'50.2" E | 100        | PVC        | 16        | 10     | P | U    |
| G70730104  | Robe 4A               | Robe             | 21°34'06.1" S, 115°50'43.6" E | 100        | PVC        | 28        | 21     | P | U    |
| G70830035  | Fortescue 32A         | Fortescue        | 21°13'18.5" S, 116°06'30.1" E | 150        | Steel      | 16        | 10     | P | U    |
| G70830105  | Fortescue 4A          | Fortescue        | 21°11'57.2" S, 116°03'6.5" E  | 80         | PVC        | 25        | 19     | P | U    |
main groups of samples were examined by one-way ANOVA for each parameter. When necessary, data were log-transformed to achieve homoscedasticity and approximately normal distributions. Manganese was omitted from this analysis because it is not a major ion.

Factors affecting species occurrence

The relationship between overall stygofauna abundance in a sample and the number of species present was examined by assigning samples to the log abundance category of the most abundant species in the sample. A one-way ANOVA was used to test whether there were overall differences in species richness across the log abundance categories and a Tukey’s HSD range test was used to examine the significance of differences between log abundance categories.

Differences in numbers of species and animals in samples belonging to different sub-regions, water chemistry groups and geologies were examined by one-way ANOVA. The seven broad categories of geology used in this analysis represent an aggregation of the units used on 1:250,000 geological map sheets by independent geological experts with the aim of amalgamating units possessing similar structure and geological history (Table 2).

Relationships between species richness in wells and their diameter, casing, distance from the coast, depth to SWL, salinity, nutrient concentration and DO were examined by one-way ANOVA or correlation analysis. The relationship between presence of casing and the numbers of species collected from wells was examined in the Fortescue basin using a small sub-set of survey wells and larger sets of wells sampled after completion of the survey by Bennelongia Environmental Consultants using the same collecting techniques and staff as the stygofauna survey (Table 3).

Temporal variability

Variation in the number of stygofauna species collected by the two samples from a well was examined in two ways. First, the number of species collected in the second sample was plotted against the number in the first sample to see how well species richness could be predicted from the first sample. Second, the number of species collected per sample in autumn and spring was compared using one-way ANOVA. Autumn was defined as April to June and represented the period when summer and autumn cyclonic and monsoonal rainfall was likely to be recharging the aquifer. Spring was defined as September to November and represented a period with dry soil and declining groundwater levels. Samples collected in July and August were omitted from the second analysis.

Table 2

Geological categories used in analysis of stygofauna abundance and richness. Numbers of wells in each category are shown.

| Geology                  | Count |
|--------------------------|-------|
| BIF                      | 16    |
| Calcrete                 | 55    |
| Granitic intrusives      | 31    |
| Mafic volcanics          | 29    |
| Quaternary alluvials     | 507   |
| Sedimentary              | 49    |
| Tertiary detritals       | 172   |

Table 3

Number of wells sampled in the vicinity of the Fortescue Lower and Fortescue Middle sub-regions to examine effect of casing on species yield during sampling. Bennelongia Environmental Consultants’ (BEC) data represent collecting in 2007 around Fortescue Lower and from 2007–2010 around Fortescue Middle in wells of 50–100 mm diameter.

| Sub-region     | Survey | BEC data | Survey | BEC data |
|----------------|--------|----------|--------|----------|
| Fortescue Lower|        |          |        |          |
| Uncased        | 0      | 28       | 2      | 35       |
| Cased          | 24     | 19       | 95     | 202      |
| Total          | 24     | 47       | 97     | 237      |
| Fortescue Middle|       |          |        |          |
| Survey         |        |          |        |          |
| BEC data       |        |          |        |          |
Community composition

In order to examine whether various, distinctly different stygofauna communities occur across the Pilbara, stygofauna samples were displayed in a three-dimensional ordination plot using order level taxonomy, the Bray Curtis dissimilarity measure and the SSH procedure in PATN. Initial species level ordination plots revealed no pattern and many sites with stygofauna had to be deleted because they shared no species with other sites. In the order level ordination, samples containing either no stygofauna or only one order of stygofauna were omitted from analysis, leaving 545 samples. Prior to ordination, the samples were classified into 10 groups of ‘similar’ composition using UPGMA, with the Bray Curtis dissimilarity measure and β = -0.1.

Species distributions

The likely range of a species is an important issue in environmental assessment because the risk of a species being made extinct by anthropogenic change is considered to be much higher for species with small ranges (Ponder and Colgan 2002; Payne and Finnegan 2007). Both the low sampling density and the fact that the wells sampled were often clustered in nodes (Figure 2) may have caused ranges of many species to be underestimated in this, and other, surveys and their conservation significance is consequently overstated. To obtain an alternative view of the likely spatial extent of a species restricted to a single sub-region, we assumed the sub-region and species’ range were squares with sides of S km and R km, respectively, and then solved the following equation for R when p = 50.

\[(S - R)^2 / (S + R)^2 = p / 100 \]  

This solution provides the median range of a species restricted to a single sub-region. Fifteen sub-regions were recognised in this analysis; 11 in the Pilbara and four on the periphery (Figure 2). We concede that some of the assumptions behind the calculation may be considered unrealistic, in particular that species ranges are not determined by hydrological or geological boundaries, but defend the calculation as an effort to obtain range information in a way that was independent of the bias in much of our sampling. We suggest the result provides useful background information about species ranges for environmental impact assessment in the Pilbara.

RESULTS

Water chemistry

PATN analyses indicated that groundwater samples from wells were best classified into 10 groups along gradients of salinity, pH and ionic composition (Figure 4, Table 4). Samples ranged from being weakly saline (WG1) to very fresh (WG8, WG10). Only two groups of samples were NaCl-dominated (WG1, WG2), although Na+ was the dominant cation in an extra two groups (WG4, WG7). Ca²⁺ was dominant in WG3, and co-dominant with Mg²⁺ in WG8, whereas Mg²⁺ was dominant in WG9 and, to a less extent, in WG5. In addition to WG1 and WG2, Cl⁻ was the dominant anion in WG3 and co-dominant with HCO₃⁻ in WG4. HCO₃⁻ was dominant in WG7-10, with a significant amount of CO₃²⁻ present in WG9. In WG5 and WG6, SO₄²⁻ was the dominant anion. WG6 contained a single sample of fresh water from well MBSKL124, south of Nullagine in the upper Fortescue catchment, and was distinct from other samples with a pH of 4.85 (Table 5).
Figure 3  Areas of intensive sampling in Fortescue Lower and Fortescue Middle sub-regions.
### Table 4

Mean values (± SE) of salinity, pH and ionic composition parameters for the four main groups of wells identified by water chemistry. *N* = number of water samples.

|         | WG1     | WG2     | WG3     | WG4     | WG5     | WG6     | WG7     | WG8     | WG9     | WG10    |
|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| **N**   | 19      | 208     | 16      | 243     | 16      | 1       | 77      | 34      | 9       | 238     |
| TDS g/L | 6.30±2.46| 2.68±0.18| 0.85±0.13| 0.87±0.02| 1.53±0.23| 0.32    | 0.78±0.07| 0.39±0.02| 0.72±0.15| 0.51±0.01|
| pH      | 6.88±0.15| 7.11±0.04| 6.88±1.78| 7.07±0.03| 6.45±0.13| 4.83    | 7.40±0.07| 6.90±0.06| 6.88±0.13| 6.93±0.03|
| Na mg/L | 54.5±2.5 | 58.0±1.0 | 24.7±6.4 | 46.6±0.8 | 30.1±1.6 | 8.5     | 71.9±1.5 | 16.0±1.6 | 13.4±3.9 | 32.7±0.5 |
| Ca mg/L | 16.5±1.5 | 14.5±0.6 | 39.5±10.2| 20.2±0.6 | 24.1±1.4 | 77.3    | 10.4±0.8 | 44.1±2.0 | 19.0±3.3 | 29.0±0.5 |
| Mg mg/L | 25.2±1.6 | 25.8±0.7 | 29.3±7.6 | 31.2±0.5 | 43.3±2.3 | 6.1     | 15.8±1.0 | 38.0±2.2 | 66.4±2.6 | 36.2±0.5 |
| K mg/L  | 3.4±0.9 | 1.8±0.1 | 6.5±1.7 | 1.9±0.1 | 2.4±0.7 | 8.0     | 1.9±0.2 | 1.8±0.2 | 1.2±0.6 | 2.1±0.2 |
| Mn mg/L | 0.4±0.1 | 0.0±0.0 | 0.1±0.0 | 0.0±0.0 | 0.1±0.1 | 0.2     | 0.0±0.0 | 0.1±0.0 | 0.0±0.0 | 0.0±0.0 |
| Cl mg/L | 91.0±1.3 | 62.3±0.7 | 68.2±17.6| 43.7±0.5 | 27.8±2.5 | 7.2     | 31.3±1.4 | 12.9±1.0 | 27.4±8.3 | 28.1±0.5 |
| HCO₃ mg/L | 5.2±1.1 | 18.3±0.5 | 27.1±7.0 | 41.2±0.5 | 23.3±3.1 | 22.6    | 59.3±1.4 | 78.0±1.8 | 57.6±8.7 | 61.6±0.6 |
| CO₂ mg/L | 0.3±0.1 | 1.5±0.2 | 0.5±0.1 | 2.1±0.3 | 1.1±0.8 | 1.0     | 4.8±0.9 | 1.9±0.6 | 10.9±4.6 | 2.0±0.3 |
| SO₄ mg/L | 3.5±1.1 | 18.0±0.5 | 4.2±1.1 | 13.0±0.4 | 47.8±3.2 | 69.3    | 4.6±0.4 | 7.2±1.1 | 4.1±0.9 | 8.3±0.3 |
Figure 4  Three-dimensional SSH ordination of water samples. A, all samples; B, group centroids. Stress = 0.09. Circle-size indicates the third dimension.

Table 5  Proportion of samples in each sub-region belonging to different water chemistry groups. Cells are shaded grey if ≥20%, or dark grey if ≥50%, of samples from a sub-region belong to a water chemistry group.
Figure 5  Physico-chemical profiles of selected wells. Wells slotted at dashed depths (slotting data not always available). A, Cowra T243 13.vi.2005; B, Cowra T243 27.ix.2005; C, Cowra T243 29.vii.2006; D, Gidgi T274B 12.vi.2005; E, Gidgi T274B 27.ix.2005; F, Weeli Wolli BH15 11.v.2005; G, Mindi T359B 12.vi.2005; H, NWSLK220B 25.ix.2005; I, MBSLK356A 25.vii.2006.
Despite the range of water chemistries (Appendix 2), 79% of samples in the Pilbara belonged to the groups WG2 (weakly saline, NaCl dominated), WG4 (fresh, Na⁺ dominant; Cl⁻–HCO₃⁻ equivalent) and WG10 (very fresh, Na⁺-Mg²⁺-Ca²⁺ equivalent; HCO₃⁻ dominant), with another 9% of samples belonging to WG7 (fresh, NaHCO₃ dominated) (Table 5). All groups other than WG6 had widespread occurrence but there was a tendency for samples from the middle and upper Fortescue and lower Port Hedland Coast catchments to belong to WG10, while the lower Ashburton, Fortescue and De Grey catchments had a high proportion of more Na⁺-dominated samples (WG4 and WG2). The upper Port Hedland Coast predominantly contained samples in the WG7 and WG10 groups. Water samples outside the Pilbara mostly belonged to WG2 and WG4, with the wells in the Great Sandy Desert almost all belonging to WG2.

**Groundwater profiles and purged bores**

Profiling showed little change with depth in most physico-chemical parameters in the wells profiled (Figure 5F, I), except around Fortescue Marsh where a lens of relatively fresh water overlaid more saline water at depth (Figure 5A–E). Dissolved oxygen was the most variable parameter, and measurements 1 m below SWL did not predict DO concentrations in the remainder of the water column for 34% of wells and were only marginally accurate for a further 7%. In some wells DO concentrations increased in the slotted section of the well, perhaps implying that water above and below the slots was stagnant and depleted of oxygen (Figure 5A–C); whereas in other wells DO concentrations either increased or declined with depth (e.g. Figure 5G), perhaps reflecting concentrations in different aquifers. Profiles appeared to be temporally stable, and wells around Fortescue Marsh that were profiled in 2005 and 2006 showed similar profiles across seasons in 2005 and between 2005 and 2006 (Figure 5A–E).

Further evidence that water samples usually reflected conditions in the local aquifer, with the possible exception of DO, was provided by the results of purging six wells. Pre- and post-purging measurements of salinity, pH and ionic composition showed almost no differences from a faunal perspective (Figure 6). One of the purged wells was profiled prior to purging (Fortescue 32A) and the sample 1 m below SWL was representative of water column concentrations of all parameters other than DO in the bottom 3–4 m of the column.

**Characteristics of Pilbara stygofauna**

At least 350 recognisable species or morphospecies of stygofauna were collected during the survey (Plate 1, Appendix 3), with 314 of these collected from the 973 samples on which most analyses were based. The additional 36 species were collected in extra sampling associated with studies of sampling efficiency (Eberhard et al. 2009) or targeted sampling. Another 10 or so described species are known to occur in the Pilbara but were not re-collected during the survey.

Sixteen broad groups of stygofauna were collected, representing at least 43 families. Among the 350 species, ostracods were the most speciose group, followed by copepods (Figure 7A). Ostracods, copepods, amphipods and oligochaetes comprised about 77% of species, with many species of nematodes, water mites, isopods and syncarids also recognised despite the lack of taxonomic work in these groups (Appendix 3). A more exaggerated pattern of ostracod and copepod dominance was observed when animal abundance was examined using the dataset of 314 species, which represented occurrence patterns better because sampling effort was more even among wells (Figure 7B). Ostracods, copepods, amphipods and oligochaetes comprised more than 96% of all animals collected.

Using the dataset for the 314 species, it appears that community composition varied considerably across sub-regions (Figure 8), with ostracods comprising more than three-quarters of the fauna in all De Grey sub-regions, about half in

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**Figure 6** Measured salinity (mg L⁻¹ TDS) and pH of selected wells before and after purging. Blue, before; red, after.
Plate 1  Stygofauna A, snail, Hydrobiidae; B, amphipod, Neoniphargidae; C, ostracod, Pilbaracandona rhabdote; D, ostracod, Gomphodella yandi; E, syncarid, Billibathynella; F, amphipod, Bogidiellidae; G, isopod, Pygolabis humphreysi; H, oligochaete, Enchytraeidae; I, copepod, Elaphoidella humphreysi.
Figure 7  Taxonomic composition of Pilbara stygofauna. A, Number of species; B, Number of animals.

Figure 8  Taxonomic composition of stygofauna in sub-regions of the Pilbara and elsewhere (Carnarvon, Eighty Mile Beach, Great Sandy Desert and north-east Gascoyne).
Figure 9  Pleotelsons, furcal rami and uropods of four possible species within *Atopobathynella*.
the Fortescue sub-regions (although with fewer animals inland), and about one-third in the Ashburton and Port Hedland Coastal (with more animals in the inland Ashburton). Copepods varied from about one-quarter to more than half the fauna in different sub-regions without an obvious pattern to the distribution. The proportion of amphipods was greatest in the lower and middle Ashburton sub-regions, where the group comprised about one-third of the fauna. The lower Ashburton sub-region was comparatively rich in oligochaetes.

Assignment to the different categories of dependence on groundwater was attempted for 290 species (the others were mostly identifications at higher taxonomic levels for which no reasonable prediction about groundwater dependence could be made). About 83% of species were considered likely to be stygobites, 10% stygophiles and 7% stygoxenes.

**Taxonomic limitations**

The 314 stygofauna species used in analyses included 136 formally named species and 153 informally recognised morphospecies, as well as 25 higher-level identifications (genus or above) that were treated as single species in analyses. Another 11 described species, 33 morphospecies and 10 higher-level identifications were recognised in the dataset of 350 species.

The higher-level identifications include some groups where more species are known, or are likely, to exist. Darwinulid ostracods were identified only to family level but subsequent work showed that four species occur in the Pilbara (Schon *et al.* 2010). Syncarids were mostly identified only to genus but genera have been shown to consist of multiple species in the Kimberley and Yilgarn (Cho *et al.* 2005; Guzik *et al.* 2008) and this is also the case in the Pilbara. The four species of *Atopobathynella* illustrated in Figure 9 were not distinguished during the survey, despite possessing some distinctive characters, because of the poorly developed state of syncarid taxonomy in Western Australia at the time when survey samples were processed.

Discrepancies between survey identifications and species boundaries were perhaps greatest for amphipods. Genetic work has shown that the paramelitid amphipod genera *Chydaekata* and *Pilbarus* contain more species than recognised morphologically during the survey (Finston *et al.* 2007). Probably the number of amphipod lineages and, most certainly, the numbers of species within each lineage were underestimated during the survey, so that overall richness of amphipods was substantially underestimated. For example, recent unpublished morphological and genetic work has shown that the lineage identified as Paramelitidae sp. 2 (PSS) during the survey is a complex of at least eight species, including three species in the Weeli Wolli Creek catchment (Figure 10).

**Sample richness and abundance**

Stygofauna were collected in 72% of samples. The average sample (including those without stygofauna) contained 3.2 ± 0.1 species and 16% of samples yielded ≥6 species of stygofauna. The average number of animals per sample was 48 ± 5.3. There was a significant relationship between the number of animals collected and the number

![Figure 10](image-url)  
Some telsons of different species within the Paramelitidae sp. 2 (PSS) lineage from the Weeli Wolli Creek catchment. A, Paramelitidae genus 2 species 2; B, Paramelitidae genus 2 species 1; C, Paramelitidae genus 2 species 3.
of species present, but this relationship was driven by the constraining influence of small numbers of animals on the number of species present, and species yield did not increase substantially at animal abundances >100 (Figure 11).

The average number of species per sample varied geographically, although differences were relatively small in most cases (maximum factor of 2.5), with Tukey’s HSD tests showing that downstream sub-regions tended to have higher richness than headwater sub-regions in the northern Pilbara (Figure 12). However, richness was variable within all sub-regions and the richness of individual samples showed overlap between most sub-regions.

The differences between Pilbara and adjacent sub-regions were of similar magnitude to differences within the Pilbara.

Identification of communities

Ten groups of samples were recognised in a classification based on order level composition of the animals in two samples from each well (Figure 13, Table 6). There was very little biogeographic signal in the classification, with the largest classification group (SG2) being the most common in all sub-regions (28–61% of samples) and the next largest groups also occurring in all sub-regions (SG4 3–23%, SG10 2–21% and SG9 3–17%,
respectively). The lack of biogeographic signal probably reflects that most stygofauna orders occur across the whole Pilbara and that site-specific factors have a greater role than geography in determining the richness and composition of the fauna at individual wells.

Most classification groups were dominated by ostracods (a dominant order in all but SG1, SG 3 and SG6, Table 6), copepods (all but SG1 and SG5) and amphipods (all but SG5, SG7 and SG9), which reflects the prevalence of these three orders in samples. The absence of ostracods from classification groups SG1, SG3 and SG6 reflects low HCO$_3^-$ concentrations, while their abundance in SG4 and SG5 reflects higher HCO$_3^-$ concentrations (Figure 13).

All classification groups other than SG2 occupied constrained areas of ordination space. SG2 contained 41% of all samples and contained the most speciose samples with diverse taxonomic composition, although SG7 samples contained almost twice as many animals on average and SG4 and SG10 samples contained 7–16% more animals than SG2 (Table 6). The very preliminary analyses conducted here suggest that, other than for ostracods, community structure is influenced less by water chemistry than by other habitat factors.

**Species distributions**

Sixty-nine per cent of the 154 stygofauna taxa collected at least twice, and representing named species or well-defined morphospecies, were collected from three or fewer sub-regions (Figure 14). This might be interpreted as indicating that the species are confined to a river basin or the catchment of a large tributary river (Figure 2). However, when the actual sub-regions occupied are examined, 79% of the species collected from two or three sub-regions were collected from more than one basin, which suggests that river basins and other surface hydrological features define the limit of a species range less well than previously thought. An alternative explanation is that many taxa recorded as species may be species complexes, with different species in different basins.

Somewhat surprisingly, given the expectation that many subterranean species will have tightly restricted distributions, only 23% of species recorded at least twice were collected from a single sub-region. The median range of those species that are actually restricted to a single sub-region was estimated using equation (1) as 682 km$^2$. This calculation assumes that ranges are characteristic of a species rather than being determined by major physical constraints associated with sub-regional boundaries, such as catchment divides. The calculation highlights the likelihood of groundwater impacts extending over distances of 20–30 km, such as may occur with large-scale mine dewatering, threatening the persistence of restricted species.

Documented species ranges, measured as number of sub-regions occupied, differed between various groups of stygofauna (Table 7). This

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### Table 6: Characteristics of groups in the stygofauna classification. Where percentage composition is not stated, orders shown are represented at 100% of sites. Rich., average number of species in a well; Abund., average number of specimens.

| N  | Rich. | Abund. | Dominant orders                          | TDS | pH |
|----|-------|--------|-----------------------------------------|-----|----|
| SG1| 18    | 2.6    | 7.1 Amphipods 67%, oligochaetes 47%, isopods 32% | 0.8 | 6.9 |
| SG2| 226   | 6.6    | 83.4 Copepods 86%, ostracods 79%, oligochaetes 59%, amphipods 57% | 1.1 | 7.0 |
| SG3| 13    | 4.2    | 19.3 Amphipods, copepods, oligochaetes     | 0.8 | 6.9 |
| SG4| 65    | 6.0    | 96.8 Amphipods 100%, ostracods 100%, copepods 85% | 1.1 | 7.0 |
| SG5| 21    | 2.8    | 37 Ostracods, oligochaetes                 | 0.9 | 7.1 |
| SG6| 25    | 2.7    | 12.1 Amphipods, copepods                  | 0.9 | 7.0 |
| SG7| 32    | 4.4    | 158.3 Ostracods, copepods, oligochaetes    | 1.3 | 7.1 |
| SG8| 37    | 6.2    | 76.5 Amphipods, ostracods, copepods, oligochaetes | 1.5 | 6.9 |
| SG9| 49    | 2.8    | 81 Ostracods, copepods                    | 1.2 | 7.1 |
| SG10| 59   | 5.0    | 89.5 Amphipods, ostracods, copepods       | 1.0 | 6.9 |
reflected either variable patterns of distribution among groups or different approaches to species delineation (recognised species may have been more likely to consist of multiple taxa in some groups and species ranges in these groups were probably overestimated). The taxonomy of larger isopods and copepods is likely to be moderately robust and the larger isopod species occupied only one or two sub-regions, whereas copepods had a median range covering four sub-regions. There was, however, considerable variation between copepod species. The widespread stygobitic species included *Stygoridgwayia trispinosa* (11 sub-regions), *Megastygonitocrella* species (*M. trispinosa*, *M. bispinosa*, *M. unispinosa*, 6–8), *Elaphoidella humphreysi* (9), *Parastenocaris jane* (7) and several *Diacyclops* species (*D. humphreysi humphreysi*, *D. sobeprolatus*, *D. cockingi*, *D. scanloni*, 13–8). Genetic analysis has excluded the possibility that occurrence of *Diacyclops* species across many sub-regions is the result of multiple cryptic species (Karanovic, T. and Krajicek 2012). Several species found widely
in surface waters as well as groundwater were also collected from multiple sub-regions, including the cosmopolitan *Microcyclops varicans* (12) and *Mesocyclops brooksi* (13), which is widespread in southern Australia.

In contrast to the copepod pattern, stygobitic ostracod species were confined to single sub-regions, with the exceptions of *Areacandona scanlonii* (7 sub-regions) and *Gomphodella hirsuta* (5), whereas species also known from surface waters occurred in multiple sub-regions. Surface species included *Candonopsis tenuis*, which occurs throughout Australia with strong groundwater affinity (4 sub-regions), *Cyprinotus kimberleyensis* (6), *Limnocythere stationis* (8) and *Cypretta seurati* (10). The names of the latter two species, which are known from outside Australia, may have been applied incorrectly to Australian endemics but the important issue is that species also found in surface water have wide ranges compared with those species occurring only in deeper groundwater. It is likely that the widespread groundwater ostracod *Gomphodella hirsuta* consists of more than one species (in addition, contradictory distributions of *Gomphodella* species have contributed to confused identifications in the genus – see Karanovic, T. 2006, Karanovic, I. 2009).

Unlike larger isopods, the taxonomic framework used when identifying amphipods suggested that nearly all species had large ranges and were found in multiple sub-regions. Information collected subsequently has shown that this range information is misleading and is the result of the taxonomic framework for Pilbara amphipods being poorly developed when identifications were made. Subsequent work has shown that most ‘species’ recognised in this paper in fact represent genera or species complexes. For example, *Paramelidae* sp. 2 (PSS) which was collected from 11 sub-regions consists of at least eight species, *Melitidae* sp. 1 (PSS) which was collected from 10 sub-regions consists of at least four species (King *et al*. 2012b), and *Pilbara millsi* which was collected from five sub-regions is probably a species complex (Finston *et al*. 2007). The work by Finston *et al*. (2007, 2011) suggests that amphipods are likely to be confined to single sub-regions.

Worms and mites were mostly widespread (Table 7). A high proportion of worm species are found in surface as well as groundwater, such as *Pristina longiseta* (10 sub-regions) and *P. aquiseta* (7). Stygofaunal mite taxonomy is not well developed but morphological examination and known life history information of the group (M.S. Harvey personal communication) suggest that most stygobitic mites in the Pilbara are moderately widespread.

**Factors affecting species occurrence**

Unsurprisingly, given the wide distribution of stygofauna communities across the Pilbara, few factors that affected sample yields strongly could be identified. The exception was well diameter; both species richness and animal abundance were greater in large wells than those with diameter <750 mm (Figure 15, *P* <0.001 ANOVA and Tukey’s HSD range tests). Casing had relatively little effect on the numbers of animals collected from wells although, in one of the two areas (Fortescue Lower and Middle sub-regions) where comparisons were made, species richness was greater in wells with casing.

**Table 7** Median number and range of sub-catchments occupied by species in various groups of stygofauna. *N* = number of species used in calculation.

| Species            | N  | Median | Range  |
|--------------------|----|--------|--------|
| Aphanura           | 4  | 3.5    | 2–13   |
| Oligochaeta        | 20 | 3      | 1–10   |
| Acarina            | 7  | 2      | 1–5    |
| Ostracoda          | 62 | 2      | 1–10   |
| Copepoda           | 32 | 4      | 1–13   |
| Syncarida          | 1  | 2      | -      |
| Thermosbaenacea    | 1  | 3      | -      |
| Amphipoda          | 18 | 3      | 1–11   |
| Isopoda            | 9  | 1      | 1–2    |
made, more species were collected from cased than uncased wells (Figure 16A, \( t = 3.19, P = 0.003 \)). Among the cased wells, those with a PVC lining yielded about 50% more animals than steel casing (Figure 16C, \( t = 2.64, P = 0.008 \)). The reasons for this are not apparent. The standard slot sizes in PVC and steel casing are the same (0.5 or 1 mm, with 1 mm used more in hard rock where no fine sediment is expected).

Depth to the water table was correlated negatively with both species richness and abundance in wells, although the relationships had very little explanatory power (7–9%, Figure 17) and depth is better viewed as a constraint on the number of species and animals that may be collected rather than a predictor of these values. Large numbers of animals (\( > 50 \)) were recorded only from depths less than 32 m and speciose samples (\( \geq 8 \) species) only from depths less than 19 m. Small numbers of animals were recorded from depths to groundwater of up to 88 m, which was the maximum depth sampled.

Stygofauna occurrence was positively correlated with DO in the top metre of groundwater but the relationship had very little explanatory power for either species richness or abundance (6–7%, Figure 18). There was little variation in the maximum number of animals or species collected across the recorded range of DO (0–100%) and the positive relationship was driven by more records of low animal and species numbers at very low DO rather than by higher numbers at high DO. Number of species collected was negatively correlated with salinity but there was no relationship for animal abundance (Figure 18). As with depth to the watertable, salinity appeared to constrain the

![Figure 15](image1.png) Mean number (±SE) of species and animals collected in samples in relation to diameter of wells. Comparisons between diameters, \( F = 20.15, P < 0.001 \) for species and \( F = 56.70, P < 0.001 \) for abundance, abundance data log-transformed.

![Figure 16](image2.png) Mean (±SE) numbers of species and animals collected according to well casing. Raw data plotted with number of species shown in left part of bar. Outliers were removed and number of animals log-transformed prior to statistical tests. A, Fortescue Lower (\( t = 3.19, P = 0.003 \)); B, Fortescue Middle; C, all survey data (\( t = 2.64, P = 0.008 \)).
Figure 17  Relationship between stygofauna collected and depth to groundwater. A, number of species ($r = -0.31$, $P < 0.001$); B, log-transformed number of animals ($r = -0.26$, $P < 0.001$).

Figure 18  Relationship between stygofauna and percentage dissolved oxygen and salinity in top metre of water. A, number of species and DO ($r = 0.24$, $P < 0.001$); B, log-transformed number of animals and DO ($r = 0.27$, $P < 0.001$); C, number of species and TDS ($r = -0.16$, $P < 0.001$); D, log-transformed number of animals and TDS ($r = 0.008$, NS). TDS values log-transformed for correlations.
maximum number of species rather determine the actual number. No stygofauna were collected at salinities >14 mg/L TDS (Figure 18) but this may merely reflect lack of samples from more saline areas. Distance to the coast explained little more than 1% of variation in richness and abundance, with numbers of animals and species declining somewhat 200 km and 400 km, respectively, from the coast.

Geology, based on amalgamated 1:250,000 mapping units, influenced species richness and animal abundance less than expected. Numbers of animals collected were highly variable in aquifers of all geologies, although this variability was masked when applying standard errors to yields from Quaternary alluvials and Tertiary detritals by large sample sizes (Figure 19B). Sedimentary rock aquifers (mostly sandstone and dolomite), Quaternary alluvials, mafic volcanics (representing some lower altitude sites in the northern half of the Pilbara), granitic intrusives (representing greenstone areas of the northern Pilbara) yielded more animals than BIF or Tertiary detritals, although the overall statistical difference in animal yield between geologies was only marginally significant (driven by the low yield from Tertiary Detritals).

Surprisingly, aquifers in mafic volcanics supported more species than other geologies, although the average number of species per sample in mafic volcanics was little more than twice the number in the least prospective geologies (sedimentary rock and BIF) (Figure 19A). Some of the differences in species richness among the geologies may relate to the pattern of animal abundance but the high species yield from Tertiary detritals compared with number of animals present suggests species richness and animal abundance may behave differently.

Neither total N nor total P concentrations affected the numbers of stygofaunal species or animals collected ($P >0.99$ in all cases).

**Temporal variation in yield**

While the number of species collected in the first and second samples from a well were correlated, slightly less than half of the variation in species richness of the second sample could be predicted from the first (Figure 20). At times the pairs of samples from a well showed large variations in yield, with differences of up to 11 species, but it was unusual for one sampling event to yield a large number of species if the other yielded none. Overall, there was no meaningful difference in average yield of the first and second samples ($3.3 \pm 0.2$ v. $3.4 \pm 0.2$). Wells sampled only once (excluded from this analysis) tended to have lower numbers of species.

Variations in the number of species collected from a well appeared to be largely stochastic. There was no overall difference between autumn and spring in the number of species collected per sample ($t = -0.14, 674 \text{ df}, P = 0.89$) and, while the mean richness per sample varied by about 30% across individual seasons, these differences were not significant (Table 8).

**Interpolation analysis**

Analyses of sub-regional patterns and the site characteristics that affect stygofauna occurrence indicate that stygofauna are distributed across the whole Pilbara, although approximately 19% of wells yielded no stygofauna. The absence of stygofauna in samples from these wells may have been the result of local conditions being unsuitable for stygofauna, well construction being unsuitable for colonisation, or sampling error whereby the animals present in the well were not collected.
Fitting a kriged surface to species richness values of individual wells (based on the average richness of samples from the wells) indicated that pockets of high species richness occurred in all parts of the Pilbara except the south-east (Figure 21). There appeared to be six extensive areas of high stygofauna richness, namely: (1) southwards from the Robe River valley in the southern part of the Fortescue Lower sub-region along the boundary of the Ashburton Lower and Middle sub-regions; (2) around the Sherlock River on the boundary of the Port Hedland Coast Lower and Upper sub-regions, extending to the northern side of the Fortescue Middle sub-region; (3) around Paraburdoo at the eastern end of the Ashburton Middle sub-region; (4) around the Coongan River and De Grey Rivers north of Marble Bar in the eastern part of the De Grey Lower sub-region; (5) around the Strelley River in the western part of the De Grey sub-region; and (6) a less well developed area of richness in the headwaters of the Nullagine River in the De Grey Middle sub-region, with a nearby small area of richness in the northern Fortescue Upper sub-region. There were a further three smaller areas of notable stygofauna richness: namely (7) the mouth of the Fortescue River in the Fortescue Lower sub-region; (8) in Ethel Gorge near Newman in the Fortescue Upper sub-region; and (9) in the western Fortescue Plain near Millstream in the Fortescue Middle sub-region (Table 9, Figure 21).

Despite identification of the afore-mentioned areas of high stygofauna richness, kriging appeared to provide limited information about the occurrence of stygofauna beyond that obtained from direct sampling results. This was perhaps because kriging inferred intermediate richness values wherever sampling had not occurred. More regularly spaced sampling points would probably have improved the capacity of kriging to infer true species richness in unsampled areas but logistical constraints (mostly the absence of suitable existing wells but also a requirement to obtain permission to access wells) prevented the selection of more regularly located wells as well as the alternative strategy of more randomly selected wells.

**Intensive sampling**

Intensive sampling within the Fortescue Lower sub-region, after the survey was completed, collected an additional 19 species that were not recorded in the area during the survey itself (a 42% increase in the fauna list, Table 10). A similar result was obtained from intensive sampling in.

**Table 8** Mean number of species (±SE) collected from samples each spring and autumn. *N* = number of samples. Between all seasons comparison, *F* = 1.73, *P* >0.1.

| Season       | N  | Mean  |
|--------------|----|-------|
| Spring 2002  | 60 | 3.6±0.4 |
| Autumn 2003 | 132| 3.5±0.3 |
| Spring 2003  | 78 | 3.1±0.4 |
| Autumn 2004 | 108| 2.6±0.3 |
| Spring 2004  | 125| 3.3±0.3 |
| Autumn 2005 | 127| 3.7±0.3 |
| Spring 2005  | 46 | 2.5±0.4 |

Figure 20 Relationship between number of stygofauna species collected in first and second samples from a well (*r* = 0.67).
Figure 21  Interpolated species richness surface based on ordinary kriging. Number of species increases with density of shading. A, all bores sampled showing how there is little information about richness in unsampled areas, B, bores with >9 species per sample, areas of high stygofauna richness are numbered, see text for details of areas.
the Fortescue Middle sub-region, where 24 extra species (33%) were recorded.

Although in both sub-regions intensive sampling collected species not previously known from those sub-regions, the overall yields from intensive sampling were 18% and 26% lower than in the survey itself, despite sampling effort being about 2–2.4 times higher (Table 10). Sixteen species (36% of the list) collected during survey in the Fortescue Lower sub-region and 34 species (44%) in Fortescue Middle were not recollected during intensive sampling. This was probably mostly a consequence of the intensive sampling covering smaller areas than the comparable units covered by the survey (see Figure 3). Species accumulation curves showed that, while the survey collected only 70–75% of the species known to occur in the two sub-regions, it accumulated species much faster than the two intensive sampling programs (Figure 22). The stratified sampling design used in the survey

| Area               | Description                                                                 | Geology                  |
|--------------------|------------------------------------------------------------------------------|--------------------------|
| 1 Robe Valley      | Most extensive area of high-yielding wells, associated with Robe and Ashburton Rivers | Quaternary alluvials     |
| 2 Sherlock River   | Area of high abundance from coastal plain to headwaters of the Sherlock River, including some George and Yule River wells | Quaternary alluvials, mafic volcanics |
| 3 Paraburdoo       | Discrete area of richness extending west from Paraburdoo wellfield           | Quaternary alluvials     |
| 4 Coongan River    | Area of moderate richness between Coongan and De Grey Rivers north of Marble Bar | Quaternary alluvials     |
| 5 Strelley River   | Discrete area of moderate richness on the Strelley River                      | Grantic intrusives       |
| 6 Nullagine River  | Diffuse area of moderate richness in headwaters of Nullagine near some areas where stygofauna appear to be absent | Quaternary alluvials     |
| 7 Fortescue Mouth  | Discrete area of high abundance associated with Fortescue River mouth        | Quaternary alluvials     |
| 8 Ethel Gorge      | Discrete Threatened Ecological Community near Newman                          | Quaternary alluvials     |
| 9 Western Fortescue Plain | Discrete area of high abundance on the Fortescue River east of Millstream | Calcrite, Quaternary alluvials |

Figure 22  Mean number (±SE) of species, and animals collected from samples in different geologies. Predicted species richness using ICE algorithm is shown. Bennelongia Environmental Consultants (BEC). A, Fortescue Lower; B, Fortescue Middle. Horizontal lines show number of species actually collected in each area from the combined sampling programs (Table 8).
provided more information about the composition of the sub-regional and regional fauna than would be obtained by combining the results of a series of intensive surveys of similar total sampling effort.

**DISCUSSION**

The collection of at least 350 recognisable species or morphospecies of stygofauna is a considerable underestimate of the richness of the Pilbara. Using the dataset reported here, Eberhard et al. (2009) calculated that between 500 and 550 species occur in the Pilbara, assuming that the taxonomy used was sound. The additional sampling undertaken in the Fortescue Lower and Middle sub-regions subsequently showed the survey under-collected species in these sub-regions by at least 42% and 33%, respectively (Table 10). This coincides well with Eberhard et al.’s (2009) estimate of 500–550 species occurring in the Pilbara as a whole, which implied under-collection by up to 36%.

The actual number of species in the Pilbara is likely to be substantially higher than 500–550 species (see Guzik et al. 2010) because, while the taxonomy used in the survey may have been reasonably sound for some groups, it was not for amphipods, syncarids, microcerberid isopods and molluscs. Nor was it sound for nematodes, bdelloid rotifers and other groups that are poorly known even in surface water systems. For amphipods, little effort was made to distinguish between the various melitid species and, although a framework was

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**Table 10**  Number of species collected during this survey and in subsequent intensive sampling by Bennelongia Environmental Consultants in the Fortescue Lower and Fortescue Middle sub-regions. Species recorded only during this survey are given as Not recollected, species recorded only by BEC are given as Extra to BEC. See Figure 14 for distribution of sampling in the two survey programs.

| Region          | Survey | BEC  | Not recollected | Extra to BEC | Total |
|-----------------|--------|------|-----------------|--------------|-------|
| Fortescue Lower | 45     | 37   | 16              | 19           | 64    |
| Fortescue Middle| 78     | 58   | 34              | 24           | 102   |

**Table 11**  Numbers of stygofauna species in different regions of the world (modified from Gibert and Culver 2009).

| Region        | No. of species | Area (km²) |
|---------------|----------------|------------|
| Pilbara       | 550            | 261,144    |
| Europe        | 2000           | 10,180,000 |
| Dinaric karst | 396            | 60,000     |
| France        | 380            | 674,843    |
| Italy         | 265            | 294,140    |
| Romania       | 193            | 233,391    |
| Asia          | 561            | 44,579,000 |
| Japan         | 210            | 377,944    |
| Africa        | 335            | 30,221,532 |
| North America | 500            | 24,709,000 |
| USA           | 300            | 9,826,675  |
| South America | 100            | 17,840,000 |
Pilbara stygofauna

established for paramelitid taxonomy, the evidence from taxa such as Paramelitidae sp. 2 (PSS) is that an order of magnitude more species may exist than were recognised. The degree of taxonomic lumping was also high for syncarids; impact assessment reports have shown that numerous species of Bathynelliidae occur in the Pilbara (e.g. Bennelongia 2012b), and the geographic distribution of survey records suggests that at least seven species were collected although only three were recognised. Among the Parabathynellidae, single species within the genera Billibathynella and Atopobathynella were recognised, although the geographic distribution of records suggests at least nine and five species, respectively, were collected. Recent work by Guzik et al. (2008) and Cook et al. (2012) suggests that most areas across Australia support multiple species of parabathynellid.

**Global significance of Pilbara fauna**

As explained above, we regard Eberhard et al.’s estimate of 500–550 stygofauna species in the Pilbara as too low but, even using the estimate of 550 species, the Pilbara is a globally important region for stygofauna in terms of the number of species present and, on the basis of current knowledge, it supports higher stygofauna species density than any region of the world other than the Dinaric karst in the Balkan Peninsula (6.6 species per 1,000 km² in the Dinaric karst v. 2.1 in the Pilbara, Table 11). While global comparisons must be treated with some caution because much of Africa, Asia and South America is unstudied, Europe and North America are moderately well surveyed and provide a baseline against which the Pilbara, Table 11. While global comparisons may, in fact, have similar richness to the Pilbara. Most species in the Yilgarn occur in the calcrete aquifers of palaeodrainage valleys (Humphreys et al. 2009).

Stygofauna occur north of the Pilbara in the Kimberley region (e.g. Wilson and Ponder 1992; Karanovic, I. and Marmonier 2002; Cho et al. 2005; Susac et al. 2010) but, while there has been relatively little survey in the Kimberley and publications are mostly taxonomic, it appears to have a less well-developed fauna than the Pilbara and Yilgarn (Hancock and Bennison 2005; Bennelongia 2012c). Poorly developed stygofauna communities also occur in south-western Australia and the Nullarbor region. Groundwater of the Swan Coastal Plain and adjacent Darling Range in the south-west supports communities consisting mostly of syncarids and stygophilic copepods (e.g. Schmidt et al. 2007; Tang and Knott 2009; GHD 2010; Bennelongia 2009a). Various relatively small cave systems support limited stygofauna communities (e.g. Jasinska et al. 1996; Karanovic, I. 2003; Eberhard et al. 2005c; Moulds 2007). The cave systems of the Nullarbor support few stygofauna species despite their large size (Holmes et al. 2001) but some of the species present are scientifically interesting (Karanovic, T. and Eberhard 2009).

Less is known about stygofauna communities elsewhere in mainland Australia, although a significant amount of work by researchers, especially at the South and Western Australian Museums, is currently underway and inventory surveys are being undertaken by industry as part of the environmental approval process (e.g. ALS 2012; Subterranean Ecology 2012a). Based on existing work, it appears that cave stygofauna communities in the Northern Territory are relatively depauperate (Moulds and Bannink 2012). Richer communities occur in calcrete aquifers of the Ngalia basin (Humphreys 2008) but there is little published evidence of widespread occurrence of stygofauna in other aquifers of the Northern Territory. Eastern parts of New South Wales and Queensland appear to support stygofauna communities of only moderate richness in alluvial aquifers (Hancock and Boulton 2008; Asmyhr and Cooper 2012; Cook et al. 2012), with about 40 species known from Queensland to date (ALS 2012) and slightly higher richness in New South Wales. Caves in New South Wales appear to have moderately diverse stygofauna communities, although in many cases identifications have not been to species level (Thurgate et al. 2001a, b).

In South Australia stygofauna communities have been found along the coast (Leij et al. 2011) and in the Flinders Ranges (Leys et al. 2010; Abrams et al. 2013). There are also rich communities of groundwater-associated invertebrates in the springs of the Great Artesian Basin and some of these species are related to stygofauna in the Yilgarn (Murphy et al. 2009).

In their review of Australian stygofauna, Guzik
et al. (2010) emphasised the importance of aridity as a factor leading to regional richness of stygofauna. Mostly on the basis of unpublished information, they suggested that arid parts of the Northern Territory and central Queensland may support numbers of stygofauna species similar to those in the Yilgarn region of Western Australia. Existing data on numbers of species, however, clearly differentiate the stygofauna communities of the Pilbara and Yilgarn from the rest of Australia, and most of the world, in terms of their species richness.

Uniqueness of Pilbara fauna

One of the major differences between stygofauna studies in the Pilbara and those in most other regions has been the way that Pilbara studies have highlighted the occurrence of stygofauna in groundwater throughout the landscape in shallow and deeper subterranean habitats (see Pipan and Culver 2012 for discussion of shallow subterranean habitats). Stygofauna have been recorded in nearly all parts of the Pilbara and the areas from which they appear to be absent are likely mostly to be sampling errors (Figure 20), whereas a recent European survey found stygofauna in only 26% of 4668 sampling units, each measuring 300–400 km² (Deharveng et al. 2009).

Nearly all stygofauna species occurring in the Pilbara are endemic in the region. Of the 290 species for which range at the regional level could be determined, 98% of the stygobites and 83% of the other groundwater species (stygophiles and perhaps a few stygoxenes) are known only from the Pilbara, with no more than three stygobitic species having ranges extending outside the Pilbara. The calanoid copepod Stygorigedgewayia trispinosa occurs on the Cape Range peninsula (Tang et al. 2008), the cyclopoid copepod Diacyclops humphreysi humphreysi occurs on Barrow Island and in south-western Australia (Karanovic, T. 2006) and the species of Stygiocaris on the Pilbara coast may be S. lancifera, which occurs on the Cape Range peninsula (a related species, S. stylifera, also occurs at Exmouth as well as on Barrow Island; Page et al. 2008).

In addition to the high level of endemism, the taxonomic composition of stygofauna in the Pilbara differs from most regions of the world because of the dominance of ostracods, which represent 41% of species in the Pilbara compared with an average 3% in the rest of the world (Figure 7; Eberhard et al. 2005a). Copepods or a combination of copepods, isopods and amphipods dominate the stygofauna communities of most regions of the world (Deharveng et al. 2009). The ostracod richness of the Pilbara principally reflects an extraordinary radiation of the family Candonidae, with 86 described and many undescribed species (Karanovic, I. 2007; Karanovic, I. and McKay 2010). Other regions of Australia have low proportions of ostracod species and very few candonids (Karanovic, I. 2003, 2005), although the genus Candonopsis has also radiated in the Yilgarn region (Karanovic, I. and Marmonier 2002).

The other notable feature of the Pilbara is the near absence of stygofaunal beetles, which form an important component of the fauna of the Yilgarn, where more than 80 species are known (Watts and Humphreys 2009). Beetles also occur in the Ngalia Basin in the Northern Territory well to the east of the Pilbara and at lower numbers in New South Wales (Watts et al. 2007, 2008) and South Australia (Leys et al. 2010). No beetles were recorded during the survey and, despite a considerable amount of other sampling for environmental impact assessment, only one stygofaunal beetle species of the family Dytiscidae has been recorded in the Pilbara (Watts and McRae 2013).

Further unusual features of Pilbara stygofauna are the low proportions of mollusc and syncarid species (1% and 5%, respectively, compared with world averages of 10% and 16%). The low number of mites in the Pilbara is markedly different from the situation in alluvial communities of eastern Australia where mites account for 23% of species (Hancock and Boulton 2008) but molluscs appear to scarce in most Australian stygofauna communities (e.g. 3% of the fauna, Hancock and Boulton 2008).

As already pointed out, the currently recognised proportions of amphipod and syncarid species (9% and 4%, respectively) in the Pilbara are likely to be considerable underestimates because of the relatively cryptic morphological differentiation amongst species. The Pilbara probably supports at least the world average proportion of amphipods (19%) and above the average proportion of syncarids (4%). The proportion of syncarids in the surveyed areas of eastern Australia is relatively high (22%, Hancock and Boulton 2008) and the group appears to be speciose throughout Australia, including in the Yilgarn (Guzik et al. 2008; Camacho and Hancock 2011; Abrams et al. 2012, 2013), especially in alluvial and colluvial aquifers.

The Pilbara supports close to the world average proportion of copepod species (18%, world average 17%; Eberhard et al. 2005a). Sampling collected relatively large numbers of copepods per species, so that the group comprised nearly half of all stygofaunal animals (Figure 7B), which is the situation in most regions of the world (Galassi et al. 2009). The proportion of copepod species in eastern Australia appears to be similar (14%) to the Pilbara, in contrast to the much richer copepod fauna of the Yilgarn that may comprise more than half the stygofauna species present (Bennellongia 2007; Karanovic, T. and Cooper 2011, 2012).
Factors affecting stygofauna yields

In line with the notion that sampling efficiency is an important determinant of how much stygofauna is collected, characteristics of the wells sampled appeared to influence stygofauna yield more than factors related to the community in the surrounding groundwater. Well characteristics include age and diameter of the well, with yields being low in recently installed wells (Bennelonga 2009b). Large diameter wells (≥2750 mm) had the highest yields, while numbers of animals caught appeared to be uniform in wells between 50 mm and 400 mm diameter (Figure 15). Sampling of a small number of 25 mm wells in the eastern Pilbara after completion of the Pilbara Biodiversity Survey suggests that yields decline in very small wells for logistical reasons: nets are difficult to lower in small wells and a substantial pressure wave is created in front of the net on retrieval.

Analyses indicate that well casing has relatively little effect on yields of stygofauna (Figure 16), although cased wells are easier to sample because there are few obstructions to snag sampling equipment. The higher yields from wells cased with PVC rather than steel matches anecdotal observations reported by Hose and Lategan (2012).

It is often remarked that stygofauna yields show considerable temporal variation and that recharge events may affect yield (e.g. Hancock and Boulton 2008, 2009). Yields frequently varied between sampling events but there was no obvious pattern and it is considered that the temporal variability was largely a result of the low detectability of species occurring at low abundance (Eberhard et al. 2009), combined with particular events at a well, such as small vertebrates falling down the well shaft.

Factors affecting stygofauna occurrence

The factors exerting most effect on the number of stygofauna animals and species in the groundwater around the well were depth to groundwater, DO in the top layer of groundwater, distance from the coast and geology, although none of these variables had strong explanatory power. While stygofauna have been found at depths of greater than 2,190 m from the surface in caves (Sendra and Reboleira 2012) and probably occur at depths greater than 100 m in the Pilbara, depth to groundwater is likely to be a major constraint on the complexity and abundance of stygofauna communities in the Pilbara. Numbers of animals collected were low where groundwater was more than 32 mbgl (Figure 17) and a similar pattern was observed by Hancock and Boulton (2008). Hahn (2006) argued that it is hydrological connectivity between the surface and groundwater that controls the negative relationship between stygofauna occurrence and depth. The depth to which tree roots extend may be another important parameter because the roots appear to provide a food source for many species (Jasinska et al. 1996; see also Hancock and Boulton 2008). Groundwater recharge in the Pilbara, together with an influx of carbon, occurs after heavy monsoonal rainfall events (Dogramaci et al. 2012).

Using almost the same dataset as presented here, Reeves et al. (2007) examined the occurrence of stygobitic ostracods and found that less than 6% of the variance in ostracod occurrence was explained by topographic and water chemistry variables. While this was considered to be largely a result of low sampling efficiency resulting in ostracods not being collected from many sites where they actually occurred (see Eberhard et al. 2009), several other studies have also found that water chemistry has little effect on stygofauna occurrence (e.g. Hahn 2006; Dole-Olivier et al. 2009), despite the expectation that divalent cation and bicarbonate/carbonate levels should be important for ostracods (Forester and Brouwers 1985; Gouramanis and De Deckker 2010) and perhaps other crustaceans (Glazier 1991). Little effect of water chemistry (other than salinity) was observed in the current analyses of the survey data.

Dissolved oxygen is usually considered to be a limiting factor for stygofauna because DO levels in groundwater can be very low (see Figure 5; Malard and Hervant 1999). Both abundance and richness of stygofauna would be expected to decline at low DO, despite stygofauna being more tolerant of low DO than most animals (Malard and Hervant 1999; Mosslacher 2000). However, the relationship between stygofauna occurrence and DO in the survey data was not strong and groundwater with almost no DO sometimes contained multiple species (Figure 18). One possible reason for the lack of relationship is erroneous meter readings, especially when values of 0% were recorded. A second reason is that DO measured 1 m bswl probably did not reflect DO throughout the well profile in a significant proportion of wells (Figure 5D, G) and, therefore, did not adequately reflect DO in the surrounding aquifer (Figure 5A). Given the uncertainty about how reliably measurements reflected DO in the aquifer, and the small amount of variation in stygofauna occurrence explained by this variable, the impact of DO is not considered further other than to emphasise that it is a difficult variable to measure accurately during stygofauna investigations in deep groundwater. It is more amenable to study, and better understood, in shallow groundwater (Datry et al. 2005).

Perhaps the most surprising result of the survey was the apparently very minor role of geology in determining stygofauna sampling yields and, by inference, stygofauna occurrence.
in the surrounding aquifer (Figure 17). The most productive geology appeared to be mafic volcanics, which was unexpected and at variance with other studies that have suggested that porous and karstic aquifers (alluvium and calcrete in the Pilbara) are the more productive (Maurice and Bloomfield 2012). There are two reasons why the sampling results from the survey may be misleading in relation to geology. First, wells were assigned to the single geology in which they were located in Geological Survey maps. In reality, there was often a vertical succession of several geologies in a well, so that wells may have sampled a more prospective geology than maps indicated. Second, wells were usually deliberately located in places with relatively high transmissivity, which may have resulted in atypically high yields from some geologies.

In contrast to the overall sampling results of the survey, eight of the nine areas of high stygofauna richness identified by kriging were located in Quaternary alluvials. Mafic volcanics were present in part of just one area of high richness around the Sherlock River (Table 9). Similarly, calcrete was present in just part of just one area of high richness. Early inventory work in the Pilbara focused on calcrete (e.g. Humphreys 1999) and subsequent work in the Yilgarn has shown calcrete to be the most important habitat for stygofauna in that region (Humphreys 2001b; Cooper et al. 2002). Projects such as PASCALIS in Europe have shown karst and alluvium to be the main stygofauna geologies (Dole-Olivier et al. 2009) and, despite the survey results, it is likely that these are also the most productive geologies in most of the Pilbara.

Stygofauna management in the Pilbara

The principal aims of the survey were related to setting priorities for nature conservation, improving the understanding of factors affecting invertebrate stygofauna distribution and sampling yields (discussed above), and providing a framework for assessing stygofauna species and community significance in relation to environmental impact assessment. Given that little is known about the life histories of stygofaunal invertebrates in the Pilbara, environmental impact assessments have focused on the threat to species persistence rather than population impacts (EPA 2003, 2013). Consequently, the species of interest have been those with small ranges that may potentially be threatened by groundwater drawdown or, less frequently, groundwater pollution.

A recent European study by Michel et al. (2009) to compare methods of selecting reserves for the conservation of stygofauna showed that use of a complementarity algorithm led to more efficient protection of species than selecting reserved areas on the basis of high richness or endemism. However, the area of occupancy criterion chosen by Michel et al. (2009) required that 46% of cells known to contain stygofauna in Europe were conserved. Given that stygofauna occur over most of the Pilbara, such an extensive network of groundwater reserves for stygofauna is unlikely to be achievable, especially considering the spatial extent of groundwater drawdown associated with some large mines (Johnson and Wright 2001).

No areas for stygofauna conservation have been proposed here. However, mapping of sampling yields and fitting a species richness surface for stygofauna across the Pilbara (Figure 21) resulted in identification of nine areas of high stygofauna richness where some protection of stygofauna values may be warranted (Table 9, Figure 21). One of the smaller areas of richness identified by the survey, Ethel Gorge aquifer near Newman, has already been recognised as important for stygofauna and is listed as a Threatened Ecological Community by the Minister for the Environment. Listing reflects high conservation value of a community and the existence of a potential threat – in this case mine dewatering – to these values (English and Blyth 1999). The Ethel Gorge community was listed in 2001 prior to commencement of the survey because a large number of species of Chydaekata amphipods had been described in the area (Bradbury 2000). Subsequent genetic work suggested that a single species of Chydaekata occurred there (Finston et al. 2004, 2007) but results of the current survey and monitoring in the area (e.g. Subterranean Ecology 2012b) have shown that about 80 species occur in the vicinity of Ethel Gorge.

The largest of the nine areas of high richness identified in the survey extended from the lower Robe River valley, where well PSS016 on the coastal plain yielded 54 species of stygofauna from 11 samples (see Eberhard et al. 2009), southwards along the boundary of the Ashburton Lower and Middle sub-regions. The boundaries of this and other areas of high stygofauna richness identified by kriging are not exact and further work should be undertaken to define the spatial extent of this area of species richness before implementing any management actions to protect stygofauna. Nevertheless, there is sufficient information to identify this area as one of the most important areas for stygofauna in a region that is a global hotspot.

Another large area important for stygofauna is on the Sherlock River in the Port Hedland Coast Lower and Upper sub-regions. The western Fortescue Plain, east of Millstream-Chichester National Park, is another important area for stygofauna. The borefield to the west of Millstream was not
identified here as important for stygofauna but its value as a water supply area, together with its proximity to Millstream, make it an area where stygofauna values may deserve more attention in water management planning than they are currently receiving (DoW 2010).

Framework for environmental impact assessment

The survey has provided a broad framework for assessing the environmental impacts of projects affecting groundwater in the Pilbara by facilitating taxonomic description of a large number of species, providing information about the typical size of species’ ranges and about areas where abundant stygofauna occur, and refining the protocols for sampling (see Eberhard et al. 2005b, 2009).

Based on existing information, most species of copepod and oligochaete collected at mine sites or wellfields in the Pilbara have ranges extending into other sub-regions and are widespread at the likely scale of project impacts (see Table 7). However, most isopod and stygobitic ostracod species will be restricted to the sub-region in which a project occurs. Amphipod and syncarid species are also likely be restricted to single sub-regions (Finston et al. 2007; Guzik et al. 2008), although poorly developed taxonomy prevented the survey dataset from showing this clearly.

The average range of species collected from multiple sites but known from a single sub-region was estimated to be 682 km². This estimate makes several assumptions and is probably more realistic in flat landscapes near the coast than in the elevated and dissected landscapes of some headwaters. However, it suggests that persistence of about half of the stygofauna species known only from vicinity of a mine may potentially be threatened, or at least species populations may be substantially reduced, where mine dewatering drawdown extends over a radius of 10 km or more (e.g. Shepard et al. 2009). Large-scale water supply borefields may also affect stygofauna, although there is usually a requirement in such borefields for production rates to be sustainable, which prevents loss of large amounts of habitat (though see Kalf and Woolley 2005; Humphreys 2009).

The simple calculation above, and the conclusions about the proportion of species under threat, may have poor biological underpinning but it highlights the potential for deleterious conservation outcomes of groundwater abstraction (and perhaps reinjection) in a region that is globally important for stygofauna. The actual threat to stygofauna species can be determined only by two processes. First, species’ ranges must be defined through adequate sampling or the effective use of surrogate information to define ranges. Second, the actual loss of habitat within the area of groundwater drawdown must be quantified. Shallow drawdowns will usually not affect persistence of stygofauna in an area unless there is stratification and the species are restricted to the upper layer of an aquifer.

Defining the range of a species requires accurate taxonomic delimitation of the species, whether or not it is formally described. There is increasing use of genetics to assist in delimiting species but it might be expected that the low dispersal capacity of stygofauna will often lead to significant genetic structuring within a species (e.g. Guzik et al. 2009, Asmyhr et al. 2013). Thus, if stygofauna species are to be recognised genetically, it is even more important than for surface species that large numbers of samples are collected to document the extent of intraspecific variation and detect interspecific divergences (Bergsten et al. 2012). Given the logistical difficulties associated with collecting large numbers of stygofauna species, morphology is likely to remain an important element of species delimitation even when genetics is used (see Carstens et al. 2013).

It is frequently impossible to determine from additional sampling whether the ranges of low abundance species, often collected as single animals or from single wells, extend beyond project impact areas (Guisan et al. 2006), especially when sampling is limited to whatever wells are available. Accordingly, the assessment guidance released by the EPA (2013) suggests that two types of surrogate information may be used to infer wider distribution of the low abundance species: namely, the distributions of other species; and the continuity of stygofauna habitat into undisturbed areas, as shown by contemporary gene flow within widespread species collected inside and outside the area of drawdown. In reality, the poor dispersal capacity of many stygofauna species means contemporary gene flow is unlikely to occur across the ranges of many species (e.g. Asmyhr et al. 2013). Maximising the available information on distributions of all stygofauna species in the local ‘community’ will probably provide more information about ranges of low abundance species than can be obtained from gene flow. Obtaining such distributional information requires sampling outside the area of groundwater drawdown and highlights the importance of the sampling program in impact assessment.

The sampling guidance released by the EPA (2007) recommended that 40 samples should be collected from the impact area of a development project (usually the area of groundwater drawdown) to characterise the potentially threatened fauna. The recommendation was based on, first, 12 samples being needed to collect nearly all the fauna from a single well (Eberhard et al. 2009) and, second, information from the current
survey suggesting that half to a third of wells in any area yield substantially better than the surrounding wells (EPA 2007). This information was taken to indicate that perhaps only a third of wells were likely to reflect fully what was in the surrounding aquifer, with factors such as the exact position of the well, its construction, history of use and contamination contributing to variation in stygofauna yield. Thus, often 36 (rounded up to 40) samples are required to collect nearly all species if the impact area is in a single homogeneous aquifer type.

Subsequent sampling programs for impact assessment have shown that in most circumstances 40 samples collect only 50–80% of the fauna (e.g. Subterranean Ecology 2011; see also Figure 22B). The likely cause of this lower than predicted collection efficiency is that most projects do not cover only one type of aquifer. Instead, there may be several aquifers and fine-scale heterogeneity of habitat and water chemistry within these aquifers (Larned 2012), leading to impact areas containing multiple assemblages of stygofauna species, each of which reflects different habitat and water chemistry conditions (Hahn and Fuchs 2009; Maurice and Bloomfield 2012). Further studies of stygofauna ecology are likely to improve sampling efficiency and the capacity to predict stygofauna occurrence in different aquifer types. Until this occurs, significant sampling programs will continue to be required to assess the threat to stygofauna species from projects with large-scale impacts on groundwater (Karanovic, T. et al. 2013).

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**APPENDIX 1 [ELECTRONIC]**

Locations of wells sampled.

**APPENDIX 2 [ELECTRONIC]**

Water chemistry at each well.

**APPENDIX 3 [ELECTRONIC]**

List of stygofauna species collected with occurrence at each well.

See CD inside the back cover or visit

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