Effects of increasing salinity on freshwater ecosystems in Australia

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Abstract. Salt is a natural component of the Australian landscape to which a number of biota inhabiting rivers and wetlands are adapted. Under natural flow conditions periods of low flow have resulted in the concentration of salts in wetlands and riverine pools. The organisms of these systems survive these salinities by tolerance or avoidance. Freshwater ecosystems in Australia are now becoming increasingly threatened by salinity because of rising saline groundwater and modification of the water regime reducing the frequency of high-flow (flushing) events, resulting in an accumulation of salt. Available data suggest that aquatic biota will be adversely affected as salinity exceeds 1000 mg L\textsuperscript{-1} (1500 EC) but there is limited information on how increasing salinity will affect the various life stages of the biota. Salinisation can lead to changes in the physical environment that will affect ecosystem processes. However, we know little about how salinity interacts with the way nutrients and carbon are processed within an ecosystem. This paper updates the knowledge base on how salinity affects the physical and biotic components of aquatic ecosystems and explores the needs for information on how structure and function of aquatic ecosystems change with increasing salinity.

Introduction

Salt is a natural component of the Australian landscape and has been deposited from a variety of sources over millions of years. Salt enters aquatic systems from groundwater, terrestrial material via the weathering of rocks or from the atmosphere, transported by wind and rain (Baldwin 1996a; Williams 1987). The relative contributions of these sources depend on factors such as distance inland, climate and geology (Williams 1987).

Under natural flow conditions in many wetlands and rivers, periods of low flow resulted in the concentration of salts in wetlands and riverine pools. Evaporation, combined with intrusions of groundwater often caused natural salinity levels to be high for periods of time (Close 1990; Williams 1999; Kay \textit{et al.} 2001) (Fig. 1A). During these periods of low flow/high salinity, biota that could not readily disperse managed to survive either with little or no reproduction and recruitment (Mills and Geddes 1980; Williams and Williams 1991) or as dormant propagules (Williams 1985; Brock \textit{et al.} 2003). Biota that are unable to tolerate these periods either perish or disperse to recolonise when more favourable conditions occur (Williams 1985).

In many river system such as the River Murray, alteration of flows, through modification of temporal and spatial patterns, has reduced the periods of high flow/low salinity and low flow/high salinity. The periods in which salt concentrations exceed the critical thresholds of biota now rarely, if ever, occur, but secondary salinisation, caused by run-off from the terrestrial landscape, has increased the amount of salt entering rivers. The reduction in the frequency of high-flow (flushing) events is causing an accumulation of salt in these river systems and a gradual increase in the mean concentration over time (Close 1990; MDBC 1999; DLWC 2000). While the salinity threshold levels for mature biota may no longer be exceeded, the mean salinity thresholds for more sensitive life stages may eventually be surpassed (Fig. 1B).

A similar pattern of salt accumulation occurs in wetlands. Prior to the removal of the terrestrial vegetation, most of the water carrying dissolved salts from the surrounding catchments was trapped by vegetation and transpired or evaporated. The salt that did wash into wetlands became concentrated by evaporation, often exceeding tolerance levels of sensitive biota. Once these wetlands dried the salt accumulated in the sediments and was removed by flushing during the next high-flow events. Removal of vegetation has increased the amount of water entering the groundwater and the amount of water and salt that enters wetlands. Many
wetlands are no longer flushed, so the continual input of salt increases concentration of salt in the sediments and this will influence biota such as aquatic plants and benthic animals (Bailey and James 2000). The increase in salt may also affect the long-term viability of dormant eggs of micro-invertebrates and seed of aquatic plants. When sediments with raised salt concentration are wetted during subsequent wetland refilling, salt concentrations in the water column again can exceed the tolerance of wetland biota (Fig. 2).

A taskforce on salinity and biodiversity established by the Australian and New Zealand Environment and Conservation Council has predicted that by the year 2050 more than 40 000 km of waterways and associated wetlands will have significantly elevated salt concentrations (ANZECC 2001). Management organisations such as the Department of Infrastructure, Planning and Natural Resources (DIPNR) in New South Wales (NSW) and the Murray–Darling Basin Commission (MDBC), have set interim end-of-catchment (or valley) targets for salinity on the basis of existing knowledge. Available data suggest that aquatic biota will be adversely affected as salinity exceeds 1000 mg L–1 (1500 EC) (Hart et al. 1991). For many plants and animals there is information available on the threshold levels of salinity on mature life stages. For most of these, there is very little information on threshold levels for earlier developmental stages (Fig. 3). The early stages of development for some biota (i.e. fish) have been shown to be more sensitive to salt than mature stages. For example, Macquarie perch (Macquaria australasica) has been shown to have a salinity tolerance of more than 30 000 mg L–1 but if eggs are exposed to salinity of only 4000 mg L –1 egg survivorship is reduced by 100% (O’Brien and Ryan 1997). While some native aquatic biota appear to be tolerant of increase in salinity above 10 000 mg L–1 (Williams and Williams 1991), early life forms may be potentially most at risk from gradual increases in salinity. There is even less information about how salinity interacts with processes in aquatic ecosystems, such as carbon and nutrient cycling.

The large spatial and temporal scales of salinity mean that if our current best land-management practices were fully implemented, salinisation would continue to increase in aquatic ecosystems throughout Australia. Although the
Effect of salinity on freshwater ecosystems in Australia

Australian Journal of Botany 657

The effect of increasing salinity on aquatic biota has been extensively reviewed. We do not understand the ecological consequences of salinisation in Australian freshwaters (Hart et al. 1991; Bailey and James 2000; Nielsen and Hillman 2000; Clunie et al. 2002). The aim of this paper is to review and update the knowledge base on how salinity affects not only aquatic biota but also the physical component of aquatic ecosystems.

Physical and chemical environment

Salinisation of a freshwater body can potentially change both the light climate and the mixing properties, which in turn have an impact on the cycling of energy and nutrients. Salt-induced aggregation and flocculation of suspended matter is recognised as a major factor in the removal of particles from the water column, resulting in an increase in light penetration, and may increase photosynthesis. The rate of clarification is enhanced by the presence of divalent cations (particularly Ca$^{2+}$ and Mg$^{2+}$) common in saline ground water (Grace et al. 1997). Increased water clarity, as a consequence of saline groundwater intrusions has been implicated in the formation of significant blooms of cyanobacteria (Geddes 1988; Donnelly et al. 1997). Alternatively, flocculation of colloids may also remove trace elements and nutrients from the water column making them less readily available to pelagic organisms (Donnelly et al. 1997).

Salinisation can alter the relative proportions of cations and anions in water that can change chemical equilibria and solubility of some minerals. The major cationic (Na$^+$, K$^+$, Mg$^{2+}$, and Ca$^{2+}$) and anionic species (Cl$^-$, SO$_4^{2-}$, HCO$_3^-$/CO$_3^{2-}$) vary between locations in both abundance and concentration. Freshwater biota are influenced as much by the ionic composition and pH of water as by the total concentration of dissolved substances (Frey 1993). The relative proportions of the main cations and anions modify the way biota respond to high salinities (Bayly 1969; Bailey and James 2000; Radke et al. 2002). Bayly (1969) suggested that the ratio of (Na$^+$ + K$^+$)/(Mg$^{2+}$ + Ca$^{2+}$) is important in determining toxicity and suggested that the monovalent ions are more toxic than divalent ions. This offers an explanation as to why many species of copepods have been found across a broad range of salinities in Australia (Hammer 1986). For example, Boekella triarticulata is a freshwater species that has been shown to survive in highly saline waters (Bayly 1969). Bayly (1969) hypothesised that the upper limit of salinity tolerance of freshwater animals is determined by the chloride content in the blood and that the suppression of this by a regime of bicarbonate might permit survival in higher than usual salinities.

Salt-dependent stratification can occur in freshwater systems following groundwater incursions. Establishment of a salt gradient can reduce mixing and solute transport within aquatic ecosystems. The halocline is a barrier for transport of materials between the surface and bottom strata and has important implications for nutrient and carbon cycling. In particular, it may become a barrier for the movement of oxygen from the surface water to the bottom, causing the rate of oxygen consumption in the bottom waters to exceed the rate of replenishment from the surface, which ultimately leads to anoxia and the death of benthic organisms (Legovic et al. 1991). Anoxia can also alter the microbially mediated cycling of nutrients. Anoxia of bottom waters has been reported in rivers where intrusions of saline water occur (Anderson and Morison 1989; McGuckin 1990; Donnelly et al. 1997; Ryan et al. 1999). Salinity of the groundwater intrusion does not need to be substantially higher than the salinity of the surface water to induce stratification and anoxia. Stratification in the Wimmera River has been observed at a salinity gradient between 300 and 700 mg L$^{-1}$ (Anderson and Morison 1989). Gribben et al. (2003) reported the formation of a seasonal salinity gradient in a

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**Fig. 3.** Changes in life-history traits as a consequence of modifying the delivery of salt. (A) Natural delivery. (B) Increased delivery. Solid line indicates increasing salinity over time. Dotted lines indicate tolerance levels for each life history phase. The time available for completion of each stage is decreased as the rate of delivery of salt is increased.
shallow freshwater wetland of only about 70 mg L\(^{-1}\), which they attribute to a ground-water intrusion during the drier summer months. This gradient, coupled with a corresponding thermal gradient, is sufficient to prevent mixing between the surface and bottom waters, with resultant anoxia in the bottom waters.

Saline ground waters can also lead to elevated levels of sulfate, dissolved iron and nitrate (Nines et al. 1992). Sulfate has been implicated in the cycling of phosphorus (Carraco et al. 1989). Sulfate-reducing bacteria use the sulfate ion for anaerobic respiration. The respiratory end product from sulfate reduction is hydrogen sulfide, which is a reducing agent that can facilitate the dissolving of iron minerals with a release of phosphorus (Boström et al. 1988). It has also been suggested that sulfide can displace P from insoluble Fe\(^{2+}\) phases (Rodent and Edmonds 1997). On the other hand, if a saline groundwater intrusion has a high level of dissolved iron, oxidation and subsequent precipitation of the iron can lead to the removal of phosphorus from solution (Baldwin 1996b). Similarly, increases in the concentration of calcium can also lead to the loss of phosphorus from solution through precipitation (House 1999).

The increase in ionic strength as a consequence of salinisation can also disrupt chemical equilibria between dissolved and particulate phases, either through changes to ion activity co-efficients or through salt ions blocking mineral surface adsorption sites (Chang 1977; Stumm and Morgan 1996). The activity co-efficient of phosphate decreases with increasing salinity, suggesting that phosphate should be more soluble in saline systems than in freshwater systems. Surface chemistry may also be disrupted with increasing salinity as cations present in salt compete with other ions for adsorption sites on particle surfaces. For example, Seitzinger et al. (1991) have shown that the concentration of exchangeable ammonium in freshwater sediments is significantly greater than in marine sediments. They attributed this difference to cations out-competing ammonium ions for adsorption sites on the sediment.

**Biological communities**

Some organisms are adapted for living in freshwater, others for living in salt water. In general, freshwater biota do not extend into saline or slightly saline water. Consequently, as salinity increases, the species richness and growth of freshwater biota is reduced (Hart et al. 1991). Freshwater is generally defined as water in which salinity is less than 3000 mg L\(^{-1}\) and sea water as 35000 mg L\(^{-1}\). These are the world average values for those systems (Boulton and Brock 1999); 3000 mg L\(^{-1}\) is often considered the lower limit for saline waters (Hart et al. 1991). Water between 3000 mg L\(^{-1}\) and 10000 mg L\(^{-1}\) can be defined as saline as biotic effects are well known within this range. Animals are divided on the basis of their ability to regulate their internal osmotic concentrations against the external environment: those that regulate internal salt concentrations well can adapt to a wide range of salinities (euryhaline regulators), whereas those that are poor regulators cannot and are restricted only to a narrow range of salinities (steno-haline regulators). Salt-tolerant plants (halophytes) tend to prefer brackish or saline conditions rather than freshwater, whereas most freshwater plants (non-halophytes) do not tolerate increasing salt concentration.

Changes in salinity can affect biota in freshwater directly or indirectly. Toxic effects as a consequence of increasing salinity cause physiological changes, resulting in a loss (or gain) of species. Indirect changes can occur where increasing salinity modifies community structure and function by removing (or adding) taxa that provide refuge, food or modify predation pressure. Other factors such as water-logging or loss of habitat may interact with salinity or have a more immediate impact on species richness (Savage 1979; Froend et al. 1987; Bailey and James 2000; Clunie et al. 2002).

Over the past 12 years several reviews on the effects of salinity in freshwater ecosystems have highlighted the paucity of suitable information for making informed predictions on what future aquatic communities will look like as salinity increases (Hart et al. 1991; Metzeling et al. 1995; Gutteridge, Haskins and Davey Pty Ltd 1999; Bailey and James 2000; Nielsen and Hillman 2000; Clunie et al. 2002). In this review, increases in salinity from less that 500 mg L\(^{-1}\) up to above 10000 mg L\(^{-1}\) are considered. This is the most likely range of salinities that Australian freshwater rivers and wetlands may experience in the next 50 years. Pulses of higher salinity are also likely to be encountered in some rivers and for some wetlands higher levels of salinity may be experienced as they evaporate and dry out. Ecological effects of salinity are likely to be observed within these ranges (Hart et al. 1991).

**Microbial function and community structure**

Bacteria have a major role in carbon and nutrient cycling. Our understanding of how microbially mediated processes change with changing salinity has come from cross-ecosystem comparisons, in which rates of various processes have been measured in freshwater, estuarine, marine and hypersaline environments. A less common approach has been to examine bacterial populations along a salinity gradient within rivers as they undergo transition from freshwater to brackish at estuaries. Understanding of the function, structure and diversity of microbial community has recently been advanced with the availability of molecular DNA methods to identify the presence and diversity of microbes, and techniques to estimate *in situ* bacterial production (growth) or the metabolic capacity of microbes. In general, aerobic bacterial heterotrophic production in different aquatic ecosystems has been found to be broadly predictable, with no consistent differences existing between
maritime and freshwater systems (Cole et al. 1988). Where
differences occur, factors such as carbon and nutrient input
and temperature are more important in regulating production
concluded that although marine and freshwater microbes
have different physiological methods for tolerating high salt
concentrations, the ecology of marine and freshwater
microbes is virtually identical. As such, it has been assumed
that a process of species replacement will occur in salinised
freshwater systems, that is, increased salinisation of
freshwater ecosystems will simply select for new
physiological types that are able to tolerate given salt levels,
but possessing the same metabolic capabilities (Hart et al.

Molecular DNA techniques have established that distinct
differences occur in the phylogenetic make up of microbial
populations in freshwater and marine ecosystems (Nold and
Zwart 1998; Crump et al. 1999). Recently it was also shown
that shifts in microbial composition occur along fresh to
brackish gradients in riverine/estuarine systems (Bouvier and
del Giorgio 2002). Metabolic activities of planktonic bacteria
also are known to vary in space and time along a riverine/
estuary gradient (Schultz and Ducklow 2000; del Giorgio and
Bouvier 2002). In the latter study, salinity was an important
determinant in separating bacterial communities.

The sparse information on the response of cyanobacteria
to salinity indicates that some members of this group occur
at salinities greater than that of seawater (\(>35000\ \text{mg}\ \text{L}^{-1}\)).
Freshwater cyanobacteria appear to be inhibited by variations in salinity (Hart et al. 1991) but may adapt to
gradual increases. Species of Anabaena have been found to
acclimatise to salinities of 7000 mg L\(^{-1}\) after several days’
exposure (Hart et al. 1991; Winder and Cheng 1995).

The relationship between salinity and specific bacterial
processes has been examined, although not extensively.
Nitrogen fixation and nitrification are known to occur in
environments with widely differing salt levels. Nitrogen
fixation by planktonic organisms generally is greater in
freshwater than in marine systems; however, within given
ecosystems, the rate of nitrogen fixation generally is
regulated by nutrient status and not salinity (Howarth et al.
1988). There appears to be no difference in the rate of
nitrogen fixation by benthic communities with respect to
different salinities. However, nitrifying and nitrogen-fixing
communities are known to vary significantly across such
systems (Affourtit et al. 2001; de Bie et al. 2001). Specific
linkages between structure and function of nitrifying and
nitrogen-fixing organisms have not been made.

A major difference between freshwater and marine
systems is the processes in anaerobic degradation of
Carbon. In marine and estuarine systems sulfate-reduction is
the major step, whereas methanogenesis dominates in fresh
systems. (Capone and Kiene 1988). This difference is driven
by the presence of sulfate ions in sea water stimulating
sulfate-reducing bacteria, which in turn are able to
out-compete methanogens for substrates (Widdel 1988).
Marine and freshwater species of sulfate-reducing bacteria
and methanogens are known to exist (Postgate 1984;
Oremland 1988).

Denitrification occurs in all aquatic ecosystems; however,
they have been suggested that in general terms the range of rates
denitrification in marine systems is greater than in
freshwater systems (Seitzinger 1988). Denitrification rates
tend to be limited by nitrate concentration, and salinity by
itself may not be the underlying regulating factor in nitrate
reduction. Molecular studies have been carried out on
denitrifying bacteria from freshwater and marine ecosystems
(Braker et al. 1998; Bothe et al. 2000; Scala and Kerkhoff
2000). However, no extensive cross-system comparisons of
denitrifying populations have been made.

Studies on rivers and estuaries continue to provide useful
insights as to how bacterial populations change across salt
grades. Whether such comparisons can readily be
transferred to freshwater ecosystems that undergo long-term
increases in salinity remains to be tested.

**Algae**

There is only sparse information on the sensitivity and
tolerance of freshwater algae; however, the majority of taxa
do not appear to be tolerant of increasing salinity (Hart et al.
1991; Bailey and James 2000; Nielsen and Hillman 2000;
Clunie et al. 2002).

The majority of algae do not appear to tolerate salinities
in excess of 10000 mg L\(^{-1}\) (Bailey and James 2000). Field
observations indicate that as salinity increases, diatoms
decrease in both abundance and richness (Blinn 1993; Blinn
and Bailey 2001). Experimental flooding of sediments has
suggested that some phytoplankton emerge in substantial
numbers when exposed to saline water but diversity is
reduced (Skinner et al. 2001; L. Bowling, unpubl. data).
Some unicellular algae such as Dunaliella salina produce
resting cysts that allow them to survive high salinities.
Species such as D. salina also undergo morphological and
physiological changes that allow them to survive across a
broad range of salinities (Borowitska 1981; Brock 1986).

**Aquatic plants**

In general, freshwater aquatic plants are not tolerant of
increasing salinity. The majority of data on the response of
aquatic plants to increasing salinity come from field
observations. The upper limit of salinity tolerated by most
freshwater aquatic plants appears to be 4000 mg L\(^{-1}\). Above
this, non-halophytes such as Myriophyllum are replaced by
more tolerant halophytic species such as Ruppia spp. and
Lepilaena spp. which have been recorded in salinities several
times that of seawater (Brock 1981, 1985, 1986).

At salinities above 1000 mg L\(^{-1}\), adverse effects on
aquatic plants appear, with reduced growth rates and reduced
development of roots and leaves. Both sexual and asexual reproduction become suppressed (James and Hart 1993; Warwick and Bailey 1997, 1998). The development of below-ground tubers, necessary for growth in the following year, and the development of flowers are also prevented (Warwick and Bailey 1996).

Information on sublethal effects of increasing salinity on germination, growth or development of aquatic plants is limited. Salt sensitivity may differ among various life stages of a species, which may reflect exposure to different environmental conditions (Bailey and James 2000). High salinity is usually inhibitory or toxic to seed germination of most freshwater plants (Ungar 1962; Williams and Ungar 1972; Baskin and Baskin 1998). For example, germination of seeds from both Sagittaria latifolia and Ruppia megacarpa decreases as salinity increases (Brock 1982; Delesalle and Blum 1994). However, there are isolated cases in which germination of a halophytic species has increased under higher salinities (e.g. Ruppia tuberosa) (Brock 1982).

Results from the experimental inundation of sediments from seven wetlands across inland New South Wales under five salinities (300, 1000, 2000, 3000 and 5000 mg L\(^{-1}\)) indicated that salinity has a significant impact on the germination of seeds of aquatic plants when it exceeds 1000 mg L\(^{-1}\). The greatest impact was on species richness and abundance in communities developing from sediment subjected to shallow flooding. Communities developing from sediment subjected to deeper flooding showed a lesser effect of salinity. This suggests that submerged aquatic plant communities may be buffered from elevated concentrations of salt, whereas those plants that live in the margins of wetlands may be more susceptible to increases in salinity (D. L. Nielsen and M. A. Brock, unpubl. data).

**Invertebrates**

It has been predicted that salinity exceeding 1000 mg L\(^{-1}\) will have adverse affects on invertebrates (Hart et al. 1991). Results from field studies examining salinity gradients in rivers or across wetlands indicate that as salinity increases there is a loss of diversity. Diversity decreases rapidly as salinity increases up to 10000 mg L\(^{-1}\), but less rapidly above 10000 mg L\(^{-1}\) (Williams et al. 1990).

Invertebrates can be divided into the following two groups: (1) microinvertebrates, comprising protozoa, rotifers and micro-crustaceans (particularly copepods, cladocerans and ostracods) (Shiel 1990) and (2) macroinvertebrates in which the major taxonomic groups are insects, worms, snails and macro-crustaceans (shrimp, yabbies) (Bennison and Suter 1990).

**Microinvertebrates**

Microinvertebrates are generally considered to be of non-marine origin (De Deckker 1983; Hammer 1986) and as a group they appear not to be tolerant of increasing salinity.

As salinity increases, there is a general decrease in abundance and richness of rotifers and microcrustaceans (Brock and Shiel 1983, Campbell 1994). There is little information on salt tolerance in protozoa, although they have been recorded from Lake Gregory, Western Australia, when the lake contains freshwater but not when it is saline (Halse et al. 1998).

Field studies have shown that there is a decrease in the number of rotifer species occurring in lakes at salinities above 2000 mg L\(^{-1}\) (Brock and Shiel 1983; Green and Mengestou 1991). In freshwater wetlands in Australia, over 200 taxa have been recorded from individual sites (Boon et al. 1990), but at high salinities, taxon richness is substantially reduced, often to as little as one or two taxa (Timms 1981; Brock and Shiel 1983; Halse et al. 1998). The rotifers and *Brachionus plicatilis*, *Hexarthra fennica* and *Trichocerca* spp. have been recorded in saline lakes (Timms 1981, 1987, 1998; Brock and Shiel 1983) and many ostracods also appear to tolerate a broad range of salinities (De Deckker 1983).

Few studies have examined the effect of increasing salinity on the emergence of microfauna from dormant eggs. It has been shown that increases in salinity may inhibit emergence from resting eggs (Skinner et al. 2001). High salinity has been linked to blocking hatching of the rotifer *Brachionus plicatilis* (Pourriot and Snell 1983), and the microcrustacean *Daphniopsis pusilla* (Geddes 1976). Ostracods have also been noted as emerging only in saline lakes when salinities are low (De Deckker 1983). In the experimental inundation of sediments from seven wetlands across inland New South Wales under five salinities (300, 1000, 2000, 3000 and 5000 mg L\(^{-1}\)), the majority of microinvertebrate taxa had significantly reduced emergence at salinities of 2000 mg L\(^{-1}\) and above. For some taxa there was a significant reduction in emergence below 1000 mg L\(^{-1}\) (Nielsen et al. 2003; D. L. Nielsen and M. A. Brock, unpubl. data). Increasing salinity may be reducing the viability of the eggs or it may be blocking the required cues to trigger emergence.

Food availability may also influence the ability of animals to tolerate increased salinity. The estuarine copepod *Sulcanus conflictus* has been shown to have lower survival at high salinities when the available food is of poor quality (Rippingale and Hodgkin 1977). However, in the case of rotifers, decreases in numbers have been linked more to specific physiological tolerances rather than food availability (Green and Mengestou 1991). The effects of salinity may also be sex-dependent. Females of the copepods *Boekella hamata* and *Acartia tonsa* are larger than males and exhibit higher survival at increased salinities (Hart et al. 1991; Cervetto et al. 1999; Hall and Burns 2001).

**Macroinvertebrates**

A large proportion of Australian macroinvertebrates has a marine ancestry (Hart et al. 1991) and as a group they appear
to be more tolerant of increasing salinity than the microinvertebrate group.

There is more information on salinity effects on macroinvertebrates than other biotic groups, as they have been widely used in the monitoring of the health of aquatic systems. Data are generally from field surveys comparing taxon presence with conductivity (salinity) collected as an environmental parameter. In some cases, there is limited monitoring of community changes at a site over time so ranges of tolerance for some taxa can be inferred from within-site as well as between-site data. Much of these data have been collated into a database (Boon et al. 2002).

In river ecosystems, macroinvertebrate diversity and salinity are not closely correlated. While salinity may cause the loss of some taxa and facilitate the intrusion of estuarine taxa upstream, increasing salinity may not be a catastrophic event. The macroinvertebrate fauna of rivers appear to be tolerant and relatively resilient to increasing salinity (Williams et al. 1991; Metzeling 1993; Metzeling et al. 1995). Data from wetlands confirm this view. Substantial changes in diversity of wetland macroinvertebrates are not likely to occur until salinities exceed 10000 mg L$^{-1}$, after which substantial loss of diversity and changes in community composition may occur (Suter et al. 1993; Halse et al. 2000). The groups most sensitive to increasing salinity are the structurally simple, often soft-bodied animals such as hydra, insect larvae and molluscs (Hart et al. 1991). Data from acute 72-h toxicity tests (LC$_{50}$) of 59 macroinvertebrate taxa indicate that the salinity tolerance ranged from 5000 up to 500000 mg L$^{-1}$, with baetid mayflies the least tolerant (LC$_{50}$ = 5500 mg L$^{-1}$) and macrocrustaceans the most tolerant (LC$_{50}$ = 38000 mg L$^{-1}$) (Kefford et al. 2003).

Although the adults and larvae of many macroinvertebrates appear to be tolerant of elevated salinity, there is little information on modifications to egg development or early instar and juvenile development.

**Fish**

Most adult native and introduced fish are tolerant of increasing salinity, but juveniles and eggs of some species are susceptible (Clunie et al. 2002).

The majority of native Australian fish are derived from relatively recent marine ancestors. Only the lung fish (Neoceratodus forsteri), spotted barramundi (Scleropages leichardti) and the Western Australian salamanderfish (Lepidogalaxias) have long evolutionary histories in freshwater (Merrick and Schmida 1984). Studies have shown that the majority of native and introduced fish in Australia appear to be tolerant of salinities exceeding 3000 mg L$^{-1}$ (Chessman and Williams 1974; Hart et al. 1991; Williams and Williams 1991; O’Brien and Ryan 1997; Whiterod 2001).

There has been only limited examination of the effect of salinity on juveniles and eggs, although evidence suggests some are susceptible to increased salinity. Eggs of the native Macquarie perch have only 50% survival when exposed to 3000 mg L$^{-1}$ and juveniles that hatched in this salinity were smaller than the controls. In a similar experiment, trout cod egg survival was reduced by 50% at 4500 mg L$^{-1}$ (O’Brien 1995; O’Brien and Ryan 1997). Eggs of silver perch are not affected until salinity exceeds 9000 mg L$^{-1}$; however, juveniles hatched at 6000 mg L$^{-1}$ had better survival than those hatched in freshwater, possibly resulting from decreased mortality as a consequence of salt-inhibiting diseases that commonly affect larvae (Guo et al. 1993). The Australian grayling, which is found in coastal rivers of south-eastern Australia and spends part of its life cycle in estuaries, produces eggs that are tolerant of salinities up to 5000 mg L$^{-1}$ (Bacher and O’Brien 1989).

**Discussion**

There is a general acceptance that freshwater ecosystems undergo little ecological stress when subjected to salinities up to 1000 mg L$^{-1}$. However, much of our understanding of the effects of salinity on freshwater ecosystems comes from lowland rivers where exposure to significant salt concentrations already occurs; other systems may be more sensitive. Hence, this view could lead to the misinterpretation that freshwater ecosystems below 1000 mg L$^{-1}$ are ‘healthy’, and there will be no adverse effects on biota or ecosystems. For many taxa, sublethal effects may not be apparent at the community level for many generations. Much of our knowledge of the impacts of salinity on aquatic ecosystems comes from field sampling along a gradient of salinity, from which it is difficult to attribute cause of ecological change. Other underlying factors such as habitat modification, loss of food resources or modification of predation pressure may also be causing changes within these systems (Blinn and Bailey 2001).

Some biotic groups are more tolerant of salinity than others. Communities of adult fish and macroinvertebrates appear to tolerate increasing salinity because they either comprise salt-tolerant remnants left after salt-sensitive species have been eliminated or reflect an evolution from marine ancestors (Williams et al. 1991; Bunn and Davies 1990; Mitchell and Richards 1992; Metzeling 1993; Kay et al. 2001). The freshwater algae, aquatic plants and microinvertebrates, appear to be less tolerant of increased salt. For these groups the general trend is to reduction of species richness as salinity increases with either a loss (or gain) in abundance. The freshwater taxa in these groups appear restricted to below 3000 mg L$^{-1}$, which may reflect a non-marine recent ancestry.

Life cycles of aquatic organisms generally are controlled by the presence of water, in association with other triggers (e.g., temperature) that cue the onset of processes such as germination of seeds, hatching of invertebrates from diapausing eggs or spawning of fish. Although specific
information on impacts of increasing salinity is limited, we
do know that life-history traits related to fitness, such as
survival, growth and reproduction, can be reduced by stress
(Hoffman and Parsons 1991). Hence, stresses such as
long-term exposure to salinity may lead to reduction in
reproduction, recruitment and ultimately depletion of biotic
reservoirs, reducing the sustainability of communities and
their ability to respond when a flush of freshwater occurs.

The current rate of change of salinity in freshwater
ecosystems may be much faster than freshwater biota can
evolve or adapt. Although lowland river biota may have
mechanisms that allow survival during periods of extreme
salt concentrations, upland rivers potentially have
experienced lower natural variation in salinity and therefore
biota in these systems may be less salt-tolerant. Induced
changes in salinity in upland systems may be too rapid for
taxa to adapt, suggesting that freshwater taxa may be lost and
communities will become dominated by salt-tolerant taxa.
Pulses of salt into freshwater ecosystems will influence
survival of a range of biota and although such increases in
salt may be rapid and short-lived, the consequences to the
survival of a range of biota and although such increases in
salt may be rapid and short-lived, the consequences to the
freshwater biota are unknown.

We need to know how salinity changes ecosystem
functioning through alteration of biotic and abiotic processes: do
changes to ecological processes change community
composition? Managers need to know more about the
relationship between flow patterns, salt concentrations and
environmental damage to predict consequences of management
actions. How a combination of changes in flow and salt affect
river and wetland communities is also relevant to management
predictions. We have many systems that are naturally variable
in both salinity and hydrology, yet we do not know how increasing
salinity will affect the biota or ecosystem integrity. Linking
salinity levels directly to mortality or recruitment potential of
aquatic biota is not sufficient to predict the outcome of
increasing salinity on freshwater ecosystems. Second- and
third-order effects must also be taken into account in describing
the full effect of salinity on aquatic ecosystems. Of particular
interest are the effects of increasing salinity on primary and
secondary production, nutrient dynamics and food-web
structure. Once we understand these interactions, links and the
flow on consequences, managers and researchers will be in a
better position to predict the condition of aquatic ecosystems
under modified salinity and move towards focusing on effective
rehabilitation. For example, the use of environmental water
allocations (environmental flows) could be considered as a tool
in managing salinity in aquatic ecosystems, once the
relationships between hydrology, salinity and environmental
damage are further delineated. Use of this relationship could
enhance effective disposal of salt-contaminated water, with
minimal damage to the environment.

If ecosystem health and salt can be related, then tools such
as water allocations, river operation, engineering
intervention and catchment management programs can be
designed to manipulate salt loads to increase the health of
aquatic ecosystems. Innovative experimental science,
together with imaginative predictive management can work
together to underpin salinity management issues on both
broad and local scales.

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References

microorganisms along the Neuse estuary, North Carolina. *Microbial
Ecology* 41, 114–123.

Anderson JR, Morison AK (1989) Environmental flow studies for the
Rylah Institute for Environmental Research, Melbourne.

ANZECC (2001) Implications of Salinity for Biodiversity,
Conservation and Management. Report prepared by a Taskforce on
Salinity and Biodiversity. Australian and New Zealand Environment
and Conservation Council, Adelaide.

Bacher GL, O’Brien TA (1989) Salinity tolerance of the eggs and larvae
of the Australian grayling, *Prototroctes maraena* (Salmoniformes:
Prototroctidae). *Australian Journal of Marine and Freshwater
Research* 40, 227–230.


LWRRDC, Canberra.


Baldwin DS (1996b) Effects of exposure to air and subsequent drying on
the phosphate sorption characteristics of sediments from a

Baskin CC, Baskin JM (1998) ‘Seeds: ecology, biogeography and
evolution of dormancy and germination.’ (Academic Press: CA)

Bayly IAE (1969) The occurrence of calanoid copepods in athalassic
saline waters in relation to salinity and ionic proportions.
*Internationale Vereinigung für Theoretische und Angewandte
Limnologie, Verhandlungen* 17, 449–455.

Commission: Canberra)

de Bie MMJ, Speksnijder AGCL, Kowalchuck G, Schuurman T,
the dominant populations of ammonia-oxidizing β-sub class
Proteobacteria along the eutrophic Schelde estuary. *Aquatic
Microbial Ecology* 23, 225–236.

Blinn DW (1993) Diatom community structure along physicochemical

Blinn DW, Bailey PCE (2001) Land-use influence on stream water
quality and diatom communities in Victoria, Australia: a response to
secondary salinisation. *Hydrobiologia* 466, 231–244.

doi:10.1023/A:1014541029984
phosphorous across sediment–water interface. Hydrobiologia 170,
229–244.
analysis of ammonia oxidation and denitrification in natural
environments. FEMS Microbiology Reviews 24, 673–690.
doi:10.1016/S0168-6445(00)00053-X
cesses and management.’ (Glenheg Publishing: Glen Osmond, SA)
Bouvier TC, del Giorgio PA (2002) Compositional changes in free-
living bacterial communities along a salinity gradient in two
systems for the amplification of nitrite reductase genes (nirK and
nirS) to detect denitrifying bacteria in environmental samples.
Applied and Environmental Microbiology 64, 3769–3775.
Brock MA (1981) The ecology of halophytes in salt lakes in the
south-east of South Australia. Hydrobiologia 81, 23–32.
Brock MA (1982) Biology of the salinity tolerant genus Ruppia L. in
saline lakes in South Australia. 1. Morphological variation within
and between species and ecophysiology. Aquatic Botany 13,
219–248. doi:10.1016/0304-3770(82)90062-6
Brock MA (1985) Are Australian salt lake ecosystems different?
Evidence from the submerged aquatic plant communities.
Brock MA (1986) Adaptions to fluctuations rather than to extremes of
environmental parameters. In ‘Limnology in Australia’. (Eds P De
Deckker, WD Williams) pp. 131–140. (Melbourne/Dordecht:
CSIRO/Junk)
Brock MA, Shiel RJ (1983) The composition of aquatic communities in
saline wetlands in Western Australia. Hydrobiologia 105, 77–84.
Drought and aquatic community resilience: the role of eggs and
seeds in sediments of temporary wetlands. Freshwater Biology 48,
1207–1218.
Bunn SE, Davies PM (1990) Why is the stream fauna of south-western
Campbell CE (1994) Seasonal zooplankton fauna of salt evaporation
basins in South Australia. Australian Journal of Marine and
Freshwater Research 45, 199–208.
Capone DG, Kiene RP (1988) Comparison of microbial dynamics in
marine and freshwater sediments. Contrasts in anaerobic carbon
metabolism. Limnology and Oceanography 33, 725–749.
phosphorous release from sediments of aquatic systems. Nature
341, 316–318. doi:10.1038/341316a0
distribution of Acartia tonsa (Copepoda, Calanoida). Journal of
doi:10.1016/S0022-0981(99)00023-4
Chang R (1977) ‘Physical chemistry with application to biological
systems.’ (Macmillian Publishing: New York)
saline waters in Victoria, Australia. Australian Journal of Marine
Close A (1990) River salinity. In ‘The Murray’. (Eds N Mackay,
D Eastburn) pp. 127–146. (Murray–Darling Basin Commission:
Canberra)
Clunie P, Ryan T, James K, Cant B (2002) Implications for rivers from
salinity hazards: scoping study. Report produced for the
Murray–Darling Basin Commission, Strategic Investigations and
Riverine Program—Project R203. Department of Natural
Resources and Environment, Vic.
Cole JJ, Findlay S, Pace ML (1988) Bacterial production in fresh and
salt water ecosystems: a cross-system overview. Marine Ecology
Progress Series 43, 1–10.
Crump BC, Armbrust EV, Barross JA (1999) Phylogenetic analysis of
particle-attached and free-living bacterial communities in the
Columbia River, its estuary and the adjacent coastal ocean. Applied
and Environmental Microbiology 65, 3192–3204.
De Deckker P (1983) Notes on the ecology and distribution of
Delesalle VA, Blum S (1994) Variation in germination and survival
among families of Sagittaria latifolia in response to salinity and
doi:10.1086/297158
(Department of Land and Water Conservation: Sydney)
Donnelly TH, Grace MR, Hart BT (1997) Algal blooms in the
Darling–Barwon River, Australia. Water, Air, and Soil Pollution 99,
487–496. doi:10.1023/A:1018351709174
coupling of bacterial and algal production in a heterotrophic
ecosystem: the Hudson River estuary. Limnology and Oceanography
36, 268–278.
Hydrobiologia 267, 233–248.
Froend RH, Hedde EM, Bell DT, McComb AJ (1987) Effects of
salinity and waterlogging on the vegetation of Lake Toolin,
Geddes MC (1976) Seasonal fauna of some ephemeral saline waters in
Western Victoria with particular reference to Parartemia zietziana
Sayce (Crustacea: Anostraca). Australian Journal of Marine and
Freshwater Research 27, 1–22.
Geddes MC (1988) The role of turbidity in the limnology of Lake
Alexandra, River Murray, South Australia; comparison between
clear and turbid phases. Australian Journal of Marine and
Freshwater Research 39, 201–209.
del Giorgio PA, Bouvier TC (2002) Linking the physiologic and phylo-
genetic successions in free-living bacterial communities along an
estuarine salinity gradient. Limnology and Oceanography 47,
471–486.
groundwater on the aggregation and settling on suspended particles
in a turbid Australian River. Colloid Surface A 120, 123–141.
doi:10.1016/S0927-7757(96)03863-0
Green J, Mengestu S (1991) Specific diversity and community
structure of Rotifera in a salinity series of Ethiopian inland waters.
Hydrobiologia 209, 95–106.
Gribben D, Rees GN, Croome RJ (2003) Anoxicogenic phototrophic
bacteria and aerobic phototrophs in Normans Lagoon, a billabong
adjacent to the Murray River, south-eastern Australia. Lakes
and Reservoirs: Research and Management 8, 95–104.
development of silver perch (Bidyanus bidentus) eggs and larvae.
Comparative Biochemistry and Physiology 104A, 531–535.
Final report to the Murray–Darling Basin Commission. Reference
Number 311/1048/06/00.
Hall CJ, Burns CW (2001) Effects of salinity and temperature on
survival and reproduction of Bocchella hamata (Copepoda:
Calanoida) from a periodically brackish lake. Journal of Plankton
Research 23, 97–103. doi:10.1093/PLANKT/23.1.97


Hammer UT (1986) ‘Saline lake ecosystems of the world.’ (Dr. W. Junk: Dordrecht)


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