

Alma Mater Studiorum Università di Bologna
Archivio istituzionale della ricerca

Sediment metal enrichment and ecological risk assessment of ten ports and estuaries in the World Harbours Project

This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Sediment metal enrichment and ecological risk assessment of ten ports and estuaries in the World Harbours Project / Birch, G.F.; Lee, J.-H.; Tanner, E.; Fortune, J.; Munksgaard, N.; Whitehead, J.; Coughanowr, C.; Agius, J.; Chrispijn, J.; Taylor, U.; Wells, F.; Bellas, J.; Besada, V.; Viñas, L.; Soares-Gomes, A.; Cordeiro, R.C.; Machado, W.; Santelli, R.E.; Vaughan, M.; Cameron, M.; Brooks, P.; Crowe, T.; Ponti, M.; Airolidi, L.; Guerra, R.; Puente, A.; Gómez, A.G.; Zhou, G.J.; Leung, K.M.Y.; Steinberg, P.. - In: MARINE POLLUTION BULLETIN. - ISSN 0025-326X. - STAMPA. - 155:(2020), pp. 111129.1-111129.23.
DOI: 10.1016/j.marpolbul.2020.111129

This version is available at: <https://hdl.handle.net/11585/758497> since: 2024-02-24

Published:

DOI: <http://doi.org/10.1016/j.marpolbul.2020.111129>

Terms of use:

Some rights reserved. The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. For all terms of use and more information see the publisher's website.

This item was downloaded from IRIS Università di Bologna (<https://cris.unibo.it/>).
When citing, please refer to the published version.

(Article begins on next page)

This is the final peer-reviewed accepted manuscript of:

Birch, G. F. *et al.* Sediment metal enrichment and ecological risk assessment of ten ports and estuaries in the World Harbours Project. *Marine Pollution Bulletin* 155, 111129 (2020).

The final published version is available online at:

<https://doi.org/10.1016/j.marpolbul.2020.111129>

Terms of use:

Some rights reserved. The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. For all terms of use and more information see the publisher's website.

This item was downloaded from IRIS Università di Bologna (<https://cris.unibo.it/>)

When citing, please refer to the published version.



Highlights

Ten global harbours were assessed for anthropogenic change (AC) and ecological risk (ER)

AC was high for Derwent River, Santander and Sydney estuaries

AC was moderate for Rio de Janeiro and Dublin Port, slight for Hong Kong, minimal for Darwin.

Derwent River sediment was rated at high ER, Sydney and Santander estuaries with moderate risk.

An improved technical framework for sediment quality assessment is provided.

Sediment metal enrichment and ecological risk assessment of ten ports and estuaries in the World Harbours Project

Birch, G. F.¹, Lee, J.-H.^{1, 2}, Tanner, E.³, Fortune, J.⁴, Munksgaard, N.⁵, Whitehead, J.⁶, Coughanowr, C.⁶, Agius, J.⁶, Chrispijn, J.⁶, Taylor, U.⁶, Wells, F.⁶, Bellas, J.⁷, Besada, V.⁷, Viñas, L.⁷, Soares-Gomes, A.⁸, Cordeiro, R. C.⁹, Machado, W.⁹, Santelli, R.E.¹⁰, Vaughan, M.¹¹, Cameron, M.¹¹, Brooks, P.¹², Crowe, T.¹², Ponti, M.¹³, Airolidi, L.¹³, Guerra, R.¹⁴, Puente, A.¹⁵, Gómez, A. G.¹⁵⁺, Zhou, G.J.¹⁶, Leung, K.M.Y.^{16,17} and Steinberg, P.³.

¹ School of Geosciences, The University of Sydney, NSW, Australia; ² Lotsearch, 3/68 Alfred Street, Milsons Point, NSW, 2061; ³ Sydney Institute of Marine Science, Mosman, NSW, Australia; ⁴ Aquatic Health Unit, Department of Environment and Natural Resources, Northern Territory, Australia; ⁵ Research Institute for the Environment and Livelihoods (RIEL) Charles Darwin University, Darwin, Northern Territory, Australia; ⁶ Derwent River Program. DPIPWE, Tasmania, Australia; ⁷ Centro Oceanográfico de Vigo, 36390, Vigo, Pontevedra, Spain; ⁸ Federal Fluminense University, Marine Biology Department, Valonguinho Campus, Niterói, RJ, Brazil; ⁹ Federal Fluminense University, Geochemistry Department, Valonguinho Campus, Niterói, RJ, Brazil; ¹⁰ Rio de Janeiro Federal University, Chemistry Institute, Rio de Janeiro, RJ, Brazil; ¹¹ Research and evaluation unit, Auckland Council, New Zealand; ¹² UCD Earth Institute and School of Biology and Environmental Science, University College Dublin, Dublin, Ireland; ¹³ Department of Biological, Geological and Environmental Sciences and Interdepartmental Research Centre for Environmental Sciences, University of Bologna, 48123, Ravenna, Italy; ¹⁴ Department of Physics and Astronomy and Interdepartmental Research Centre of Environmental Sciences, University of Bologna, 48123, Ravenna, Italy; ¹⁵ Environmental Hydraulics Institute, Universidad de Cantabria, Avda. Isabel Torres, 15, 39011, Santander, Spain; ¹⁶ The Swire Institute of Marine Science and School of Biological Sciences, The University of Hong Kong, Pokfulam, Hong Kong, China; ¹⁷ State Key Laboratory of Marine Pollution, City University of Hong Kong, Kowloon, Hong Kong, China.

+ Present address: Balearic Islands Coastal Observing and Forecasting System, SOCIB, 07122, Edificio Naorte, Bloque A, Parc Bit, Palma de Mallorca, Spain.

ABSTRACT

Ten global harbours were assessed for sediment quality by quantifying the magnitude of anthropogenic change and ecological risk. Anthropogenic change (enrichment) was high for Derwent River and Sydney estuary, moderate for Santander Harbour, Rio de Janeiro and Dublin Port, slight for Hong Kong, minimal for Darwin. All 10 enrichment indices used showed similar results. Derwent River sediment was rated at high ecological risk, followed by Sydney and Santander estuaries with moderate risk. Auckland and Darwin sediments exhibited minimal ecological risk and sediment in the remaining harbours (Dublin, Hong Kong, Ravenna, Ria de Vigo and Rio de Janeiro) were assessed at slight ecological risk.

The extraordinary variety of environments and types/quantities/qualities of data investigated resulted in as much a critique and development of methodology, as an assessment of human impact, including unique techniques for elemental normalisation and contaminant classification. Recommendations for an improved technical framework for sediment quality assessment are provided.

Keywords: Sediment Quality Guidelines, Environmental Indices, Anthropogenic Change, normalisation, Classification scheme

1. INTRODUCTION

Substantial anthropogenic stress on coastal marine environments has reduced sediment and water quality in many global harbours and estuaries resulting in threatened benthic and pelagic populations (Costanza et al., 1997; Chapman and Wang, 2001; Costanza et al., 2014). It is important that the extent of contamination of an estuarine environment is assessed as accurately as possible and that the level of threat to the health of these sensitive environments is determined with care. Choosing an appropriate methodology to assess environmental condition is complex and requires an integrated strategy (Rees et al., 2008). Ecosystem indicators used to assess anthropogenic stress are commonly compromised by natural spatial and temporal variability. This confounding of natural and anthropogenic-induced stress results in inappropriate and often erroneous assessment (Hogg and Norris, 1991). Increasingly, sediments are being used to assess the status of aquatic environments due to the advantage sediments have in faithfully recording and time integrating environmental events, both temporally and spatially (Rodrigues et al., 2006; Birch et al., 2008). Moreover, sediments greatly influence the quality of overlying and interstitial water and are an extensive habitat to a large number of faunal and floral species (Simpson et al., 2015).

The condition of estuarine environments may be described using a wide range of approaches, e. g. ecological risk indicators (Singh et al., 2005), concentration factors/indicators (Guo et al., 2010) and enrichment factors/indicators (Caeiro et al., 2005). However, from a management perspective, it is important to know by how much the system has deviated from the pristine condition, i. e. the magnitude of anthropogenic change, and the degree of risk of potential harm posed by sedimentary contaminants to biological communities (Birch, 2016). These two types of information differ fundamentally and are based on different types of data. A full and comprehensive assessment of sediment health is complex and requires a raft of chemical, sedimentological and ecological approaches (Ponti et al., 2009; Birch, 2016; Birch, 2018), however information on the degree of anthropogenic change and level of ecological risk posed by sedimentary chemicals provides a useful initial screening assessment of environmental quality in marine environments.

The objectives of the present work are to determine the magnitude of anthropogenic change and the risk posed by sedimentary contaminants to the biotic ecosystem for the ten harbour estuaries involved in the World Harbour Project (WHP) from a wide range of environments in multiple locations across the globe (Fig. 1) in an effective and regionally consistent manner using traditional, as well as innovative methodologies.

1.1 The WHP Concept

The WHP was initiated by the Sydney Institute of Marine Science (SIMS) and aims to develop resilient urban harbours through a global network of collaborating researchers (Steinberg et al.,

2016). The project brings together international research institutions and agencies concerned with the health of these heavily urbanised waterways and the increasing challenges these environments face. Like Sydney, many of the great cities of the world, such as Auckland, Rio de Janeiro and Hong Kong, are located on the coast. These working harbours are part of the fundamental fabric of those communities and the relationship between the residential, industrial and marine environments require ongoing study and management. The WHP tackles issues surrounding the multiple uses of harbours through global collaborative projects and targeted workshops. The Project works to facilitate and link programs across major international harbours, with a proactive focus on investigating and restoring ecosystem functioning and the consequent development of management best-practices that can be applied by all partner cities

1.2 Harbours of the WHP

The morphodynamic characteristics of estuaries influence benthic contaminant distributions and concentrations through tidal patterns, flushing and structural control (Roy et al., 2001; Birch et al., 2016). Numerous classification schemes have been developed in an attempt to combine diverse physical, biological and ecological characteristics of coastal waterways (Dalrymple et al., 1992; Edgar et al., 2000; Roy et al., 2001). The ten harbour estuaries assessed in the current work (Fig. 1) have been loosely separated into five morphodynamic groups, namely (1) funnel-shaped, wave-dominated, drowned-valley estuaries: Derwent River (Tasmania, Australia), Sydney estuary (New South Wales, Australia), Ría de Vigo (Spain) and Auckland Harbour (Waitemata Bay, New Zealand); (2) tide-dominated, drowned-valley estuaries: Darwin Harbour (Australia) and Santander Harbour (Spain); (3) open ocean embayments: Hong Kong; (4) partially enclosed embayments: Rio de Janeiro, Guanabara Bay, (Brazil); and (5) river mouths/canals: Dublin Harbour (Ireland) and Ravenna Harbour (Italy).

Auckland Harbour, Waitemata Bay, New Zealand

The Waitemata Harbour (80 km²) is the largest east coast estuary in the Auckland region and is comprised of tidal creeks, embayments and a central basin (Aguirre et al., 2016). Sediment studies have been mainly confined to urban and rural tidal creeks, local harbours and embayments, which receive the major contaminant loads, leaving the central basin largely unsurveyed. These peripheral environments are predominantly muddy and fringed by mangrove mudflats. The catchment (185 km²) comprises mainly urban (21%), rural (25%), forest (21%) and minor industrial landuses. Urbanised catchments have been identified as major sources of fine sediment and metals to the harbour (Aherns, 2008; Mills et al., 2012; Mills and Williamson, 2014). Henderson Creek, which drains the largest urban subcatchment, as well as a substantial area of rural land, contributes the largest loads of sediment and metals to Central Waitemata Harbour. Present-day surface sediments show spatially-variable concentrations of metals with maximum concentrations occurring on intertidal flats near tidal creek outlets and stormwater drains in the south western embayment of Central Waitemata Harbour and upper Shoal Bay. Marked increases of metal discharge was initiated around 1950 coinciding with the beginning of

149 rapid urbanisation and reached a maximum more than 20 years ago, while declining slightly to
150 the present time.

Darwin Harbour, Northern Territory, Australia

151 Darwin Harbour (~1220 km²) is approximately 35 km long and 5 km wide at the mouth. The
152 waterway is macro-tidal (range up to 8m) and comprises extensive mudflats (containing
153 substantial calcium carbonate from shell material) fringed by large stands of mangroves
154 separated by deep tidal channels. Parts of the Harbour are relatively poorly flushed, especially in
155 the dry season when the residence time in the upper arms is ~20 days. Most (85%) of the
156 catchment (~2010 km²) is open space, while urban and light industry occupy 7% (no heavy
157 industry), horticulture makes up about 1-2 % of the area and rural/residential comprises 6 %.
158 Although sedimentary metal concentrations are generally low, moderate environmental concern
159 is for elevated metals along the developed eastern side of Darwin Harbour and in the vicinity of
160 the sewage treatment plant outfall north of the city (Munksgaard et al., 2012; 2013; 2015; 2018).
161 Metal loads have declined as a consequence of treatment system improvements, however
162 emerging sources of metals are from removal of marine sediments associated with coastal
163 development and dredging activities.

164 Derwent River, Tasmania, Australia

165 The Derwent River is approximately 45 km long, occupies 200 km² and is relatively deep (av. 25
166 m) (Whitehead et al., 2010), The waterbody is stratified in the narrow upper reaches and well
167 mixed in the lower, broad seaward area. The tidal range is 1 m and average flushing time is 12
168 days. The catchment (8,900 km²) is mainly agriculture and forest (76%) and urban/industry (8%)
169 and supports 40% of the state population. The main environmental issues are metal
170 contamination of water and sediment from a zinc smelter and nutrients from a paper mill and
171 waste water treatment plants. Nutrient loads have declined, but Zn, mainly from groundwater,
172 remains elevated.

173 Dublin Harbour, Ireland

174 Dublin Bay hosts Dublin, the capital city of Ireland (population >1m) (Brooks et al., 2016) and
175 Dublin Port, busiest shipping port in Ireland, which accounts for half of the nation's annual
176 imports and exports (DPC, 2014). Dublin Port (3.267 km²) has developed on both sides of the
177 Liffey channel, but most of the shipping docks are on the northern side, while the southern port
178 encompasses major infrastructure. Technical development and regulation backed by
179 environmental monitoring has considerably reduced most adverse effects of deliberate
180 contamination. Despite such controls, input of contaminants to the port from discharge of sewage
181 and industrial waste, spillage of cargo, or ship lubricants, stormwater runoff and warm water
182 from power stations still occurs. Riverine inputs (mainly from the Liffey River), which drain the
183 highly urbanized and rural areas around Dublin city, are a major source of contaminants into
184 Dublin Bay (EPA, 2006; 2015; Brooks et al., 2016; Murphy et al., 2016; Cunningham, 2018).

Resuspension of contaminated sediments may occur due to strong tidal forces, storm surges, or increased water movement from boats (Davoren et al., 2005; Macken et al., 2008; Bedri et al., 2011; Briciu-Burghina et al., 2014).

Hong Kong Harbours

Hong Kong is situated in the Pearl River Estuary (PRE), and covers about 8000 km² (Chen et al., 2013). The estuary (1649 km²) is up to 30 m deep (av. <15 m) and is microtidal with a <2 m average tidal range (Mao et al., 2004). Bottom sediment is mainly mud with subordinate sand and gravel (Tanner et al., 1993). Sedimentary metals are significantly higher in harbours, e. g. Tolo Harbour and Victoria Harbour, than in offshore areas (Chan et al., 2016; Blackmore, 1998; Zhou et al., 2007a, b; Tang et al., 2008; Chen and Jiao, 2010; Liu et al., 2015; Chen et al., 2013). The territory (1108 km²) supports >7.4 million people and comprises Hong Kong Island, Lantau Island, Kowloon Peninsula, the New Territories and 261 islands with a complex and long coastline (1200 km) (EPD, 2017; SMO, 2018). Hong Kong is a highly urbanized (81 %), coastal city receiving substantial metal loads from industrial and municipal waste waters, especially during the 1950s-1980s (Morton, 1989). Metal contamination of marine sediment is mainly attributed to: (1) historical discharge of untreated industrial wastewater and partially treated sewage (e. g., Victoria Harbour and Tolo Harbor); (2) surface runoff from Pearl River and local rivers (e. g. Deep Bay), and (3) other sources, including infiltration from septic tanks and leachate of antifouling compounds from shipping (Chen and Jiao, 2010; Liu et al., 2015). Improved conditions since the late 1980s are due to industry moving to the New Territories, Southern China and other areas of Asia (Morton, 1996; Blackmore, 1998) and improved treatment of domestic sewage (Lai et al., 2016). However, metal contamination persists due to release from legacy contaminated sediments.

Ravenna Harbour, Italy

Ravenna is the largest harbour (3.62 km²) in the western Adriatic and is one of the most extensive commercial seaports in Italy. The harbour was established in lagoonal systems surrounding the city and is structured as a major 'canal' port extending for 11 km from the centre of Ravenna to the tourist seacoast. The canal is directly connected to surrounding lagoons (Pialassa Baiona and Pialassa Piomboni), which are included in the southern part of the Po River Delta Park, inscribed in the World Heritage List. Construction of two large converging jetties (2400 m long each) to protect the harbour from siltation has altered sediment transport and has re-shaped nearby tourist beaches. Environmental concerns include degradation of natural habitats, contamination of sediments and management of highly urbanised areas (Airoldi et al., 2016). Ravenna Harbour and coastal lagoons receive civil and industrial wastewater carrying nutrients, pollutants and cooling water from two power stations and industrial plants. Although discharges now comply with current laws, lax legal constraints between 1958 and 1976 resulted in sediment of Pialassa Baiona being heavily impacted by industrial metals, including Hg (Fabbri et al., 1998; 2000; 2001; McRae et al., 2000; Guerra, 2012; Guerra et al., 2014). However,

sediment resuspension due to frequent maintenance dredging and to deepen the port have had minimal effects on macrobenthic assemblages inhabiting the lagoons (Guerra et al., 2007; 2009; Ponti et al., 2009; 2011).

Ría de Vigo, Spain

Ría de Vigo (156 km²) is the most populated (411,363 inhabitants) and developed (approximately 7% industrial, 12% construction) ria in Galicia and is home to Vigo city (population 292,986) (Galician Institute of Statistics, 2017; <http://www.ige.eu>). The ria is approximately 30 km long and 12 km wide at the mouth. The waterway is relatively deep (av. 16 m, max. 45 m) and the water column is well mixed with a tidal range of 4 m and an average flushing time of 3-4 days (Barton et al., 2015). Sediments are predominantly organic-rich and fine grained. The catchment (578.2 km²) is heavily urbanised and industrialised (>21%) (Fernández et al., 2016) and activities include shipbuilding, canning, automobile and steel manufacturing. Metal pollution is restricted to the inner estuary as a result of urban and industrial discharges and to intense activity of the Port of Vigo with chronic Pb pollution due to discharge from a ceramic factory located at the head of the estuary (Rubio et al., 2000; Prego and Cobelo-García, 2003; Alvarez Iglesias et al., 2006; 2007; Quelle et al., 2011). Other sources are natural, related to catchment and upwelling processes (Quelle et al., 2011). Mariculture rafts in the northern estuary have influenced the distribution of metals by increasing the carbon content and decreasing grain size producing metal sinks. Tidal currents act to redistribute metals from accumulation zones.

Rio de Janeiro, Guanabara Bay, Brazil

Guanabara Bay (449 km²) is located in the southeastern Brazil Marine Ecoregion in the most economically developed region of the country. The bay comprises a large, shallow (av. 5.1 m) inner semi-circular water body with a maximum length of 48.2 km and a narrow inlet with a main central channel depth of 58 m. Mean spring tidal range is 1.05 m. The catchment is the largest of this ecoregion (~3700 km²) (Kjerfve et al., 1997; Kjerfve et al., 2001) and supports Rio de Janeiro, the second largest Brazilian city (6.5 million people). The drainage basin of the Guanabara Bay (4180 km²) is drained by approximately 45 rivers (JICA, 1994) and the main rivers are the Macacu, Iguaçu, Estrela and Sarapui. The Guanabara Bay catchment includes, partially or totally, 12 municipalities with a population of almost 10 million inhabitants, equivalent to 80% of the population of the State of Rio de Janeiro. As a consequence of urban, agricultural, and industrial development, the bay is one of the most altered and polluted in the country (Carreira et al., 2002; Xavier de Brito et al., 2002; Silva et al., 2013; Camargo et al., 2017; Cordeiro et al., 2015; Baptista Neto et al., 2017). Anthropogenic metal sources to the bay include direct discharges of untreated and treated industrial waste, domestic sewage, inputs from rivers, atmospheric fallout, dockyards and agricultural activities, landfill and road runoff (Rebello et al., 1986; Abuchacra et al., 2015; Aguiar et al., 2018). Metal loads to the bay have increased substantially over the last 70 years caused by population growth in the metropolitan region (from ~2.5 to 12 million) accompanied by extensive urbanization, deforestation and

industrial and agricultural growth (Moraes, 2012; Covelli et al., 2012; Figueiredo Jr. et al., 2014). Higher concentrations of metals located in the inner bay from river discharge and in sediments adjacent to harbours (Baptista Neto et al., 2006; Cordeiro et al., 2015; Aguiar et al., 2018) exhibit significant ecotoxicological effects on aquatic organisms (Moraes et al., 2000; Maranhão et al., 2009; 2010; Campos et al., 2019).

Santander Bay, Spain

Santander Bay (22.5 km²) is one of the most important and largest estuaries in northern Spain (Biscay Gulf) and includes a Special Protection Area (SPA) (Gómez et al., 2014). The bay is a natural harbour and hosts a major commercial port. The bay is characterized by a semidiurnal tidal regime with a medium tidal range of 2.9 m and interacts with freshwater discharges from the Cubas River (Puente et al., 2002). The estuary is dominated by extensive, shallow water (max. depths 10–12 m) intertidal areas (67%), which have been greatly modified by urban development and port activity (Ondiviela et al., 2013). Anthropogenic activities began in Roman times (Vigurí et al., 2007) reaching a maximum impact during the 1970s (Vigurí et al., 2007). Since marsh reclamation in 1903, the inner bay has undergone intensive industrial expansion of mainly metallurgical and chemical industries. During the last 150 years between 37% and 50% of the original intertidal zone has been reclaimed (Vigurí et al., 2007; Remoundou et al., 2015; Calleja et al., 2017) and used as grasslands, to expand the Port, and to create new industrial and residential areas, including the city of Santander. Continuous, untreated industrial discharges ceased in 2001 when a new sewer system came into operation (López et al., 2013; Echávarri et al., 2007) and all direct discharges to the bay were eliminated in 2010 when a new wastewater treatment plant was commissioned. Industrial contaminant sources are mainly located in the inner bay and on the western shore, where the port is located. Maximum metal concentrations occur in the subtidal and inner estuary related to industrial sources and are minimal in the intertidal flats (Puente et al., 2002).

Sydney estuary, New South Wales (NSW), Australia

Sydney estuary is approximately 30 km long and up to 3 km wide with an area of 50 km², while the catchment (500 km²) is highly industrialized and urbanized (76%) (Birch et al., 2015; 2016; Birch, 2016) and supports the City of Sydney (population 5.5m). Estuarine water is generally well-mixed marine, but becomes stratified after prolonged heavy rain (Lee et al., 2011; Birch and McCready, 2009). Typical flushing times are 5 to 10 days, however in the upper reaches of the waterway it may be up to 130 days. Sediments in the estuary are mainly muddy in the upper reaches and sandy in the mid- and lower estuary. Sediments are significantly contaminated by metals and organic compounds (Birch et al., 2000a; 2008; 2013; McCready et al., 2004; 2006; Birch, 2017) and the waterway is classified as “severely modified” (NLWRA, 2002; Birch and Taylor, 2000a). Until recently, the harbour was a busy commercial and naval port and the shoreline was lined by factories, however, industries have moved away from Sydney and the area has converted into a mainly tourist and recreational hub (Birch and Taylor, 2000b, c).

2. METHODS AND MATERIALS

2.1 Analytical Methods

Auckland Harbour

Total recoverable metals were determined on the <500 µm fraction by hot acid digestion (HNO₃/HCl) (USEPA Method 200.2) (Table 1).

Darwin Harbour

Sediment samples were wet sieved to <2 mm grain size and digested with in a strong HNO₃/HClO₄ (perchloric acid) mix using open digestion tubes in a heating block. Elemental analysis was by inductively coupled plasma mass spectrometry (ICP-MS) using a marine sediment Certified Reference Material (CRM) MESS-3.

Derwent River

Shallow cores were taken using a triplicate multi-corer, with the upper 5 cm extruded and mixed to provide a total sediment integrated surface sample. Subsamples were also size normalised to 62.5 µm and analysed using inductively coupled plasma atomic adsorption spectrometry (ICP-AES).

Dublin Harbour

Samples were dried at <30°C, crushed and sieved at 2mm. Samples were digested in HNO₃ using open digestion tubes in a heating block. Analysis was via inductively coupled plasma optical emission spectrometry (ICP-OES).

Hong Kong

Marine sediments were collected using a Van Veen grab sampler and samples were digested by microwave-assisted acid extraction (HNO₃) (ISO, 1995) and analysed by ICP-MS (USEPA, 1994).

Ravenna Harbour

Samples were digested in closed Teflon vessels by a mixture of HNO₃ + HCl (3:1) in a microwave system and measured by graphite furnace atomic adsorption spectroscopy (GFAAS) using a Certified Reference Material PACS-2 (Marine Sediment, NCR-CNRC, Canada).

TABLE 1

Ría de Vigo

Total sediment was digested with a strong acid mix of HNO₃/HCl/HF in Teflon digestion bombs using a microwave oven and analysed by spectrophotometer. Quality assurance systems using Certified Reference Materials (BCSS-1 and BEST-1, National Research Council of Canada).

Rio de Janeiro

Sediments were digested using a microwave-assisted procedure with concentrated HNO₃ solution (USEPA, 2007a) and analysed by ICP-OES. Analytical quality control was assessed by analysing a certified reference material (NIST 2782 Industrial Mud).

Santander Harbour

Sediment samples were collected with a Van Veen grab and grain size was determined by dry sieving. Metals analysis was for the <63 µm fraction and followed the U.S. Standard method (US EPA 6020, 2007b) using a HNO₃/HCl/HF mix and analysed by ICP-MS and a Certified Reference Material (Loamy Clay, CRM 052, Resource Technology Corporation, US).

Sydney estuary

The fine (<63 µm) fraction and total sediment were analysed by aqua regia digestion (HCl/HNO₃) (modified US EPA 200.8 Rev 4.4 method) (US EPA 1994) and analysed by ICP-OES using a reference material (AGAL-10), procedural blanks and blind replicates (Siaka et al., 1998; Birch and Taylor, 2000b).

2.2 Data availability and sample distribution

Data used for Auckland Harbour assessment were for five metals (no Al, Fe, Ni) for 41 sites sampled in 2008 (Table 2). Sampling (Cu, Pb and Zn n=1221 and for As, Cd, Cr n=37) targeted areas of interest located primarily within embayments, tributaries and coastal zones. The 298 samples collected from Darwin Harbour in 2012 were distributed on inter-tidal flats and were absent in strong-flowing tidal channels. Metals data (n=123) sampled in 2000 in the Derwent River provided an excellent coverage for contaminant mapping. Sampling (n=42) undertaken in Dublin Port in 2006 in the main channel of the docks and in two berthing basins provided sufficient coverage to map the distribution of contaminants.

TABLE 2

Duplicate samples from 45 sites for the years 1995 to 2015 (n=2676) were available for the Hong Kong coastal region, however only the most recent vintage (2015/16) of samples (n=9) confined to the harbour areas (63.0 km²) were used for mapping in the present assessment. Several of the canals of Ravenna Harbour have been sampled, but lacked Al, Fe, or size data and only sediments (n=52) of Canale Candiano (3.52 km²) have been assessed. The 39 samples taken in 2011 in Ria de Vigo estuary were well distributed and provided a good spatial spread of data throughout the embayment. Although sampling was undertaken for multiple years between 2005 and 2015 in Santander Harbour, sites were sparsely distributed and the data for 2015 (n=10) mapped in the present study were located close to the shoreline with an absence of samples in the central harbour (22.5 km²). Approximately decadal sampling has been undertaken in Sydney Harbour between the years 1975 to 2015. The 2000-2015 dataset (n=1175) used in the current assessment was well distributed throughout the estuary and provided an excellent spatial platform for mapping and assessing contaminants.

As an adequate sampling density is required for GIS mapping, metal distributions of only six harbours (Darwin, Derwent, Hong Kong, Ria de Vigo, Rio de Janeiro and Sydney) could be plotted as filled contour maps (Figs. 2-8), while contaminant distributions of the remaining four harbours (Auckland, Dublin, Ravenna and Santander) were depicted as points in the Supplementary Material file (Figs, S1-S7). The amount of data available was not consistent across all 10 harbours and thus assessment could not be completed for all techniques for all harbours.

2.3 Differentiation between ‘anthropogenic enrichment’ and ‘sediment quality’

In a management perspective, two important attributes define the ‘environmental health’ of sediment in an aquatic ecosystem. Anthropogenic ‘enrichment’ is the magnitude of human-induced change in the aquatic environment and measures the departure of the system from the pristine condition. ‘Sediment quality’ is the ability of sediment to maintain a healthy benthic community and is measured by ecological risk assessment. Enrichment does not imply ecological effect, or toxicity. These two metrics are based on different methodologies and criteria, require different types of data and are entirely unrelated, however these attributes are frequently confounded and often aggregated into a single value, or index (Caeiro et al., 2005; Wilson and Jeffery, 1987; Kabir et al., 2011). To measure human-induced change requires pre-anthropogenic contaminant concentrations (commonly referred to as ‘background’) to be known and that data are normalised (either size-, or elemental normalisation) to reduce the confounding of variable grain size, while ecological risk is based on total sediment using sediment quality guidelines (SQGs). In the present work, only sedimentary metals (Cd, Cr, Cu, Ni, Pb and Zn) were considered, as a full suite of organic and inorganic pollutants were not available for all harbours.

2.4 Techniques to measure anthropogenic change (enrichment) in the WHP

Two complications with the WHP dataset, i. e. that background concentrations for metals were not available for all locations and that size-normalised data were accessible for only three locations (Derwent River, Ria de Vigo and Sydney) and Post-Extraction Normalisation (PEN) was undertaken for Sydney and Darwin. In the PEN technique, the >63 µm fraction is removed by sieving after digestion thereby capturing metals associated with the total sediment, providing an advantage over the usual size-normalisation procedure, which only measures contaminants in the <63 µm fraction (Birch and Taylor, 2000; Birch, 2003). The lack of these data restricted the ability to determine the magnitude of anthropogenic change, i. e. sedimentary metals enrichment.

Early researchers used global average upper crust and marine shale metal concentrations as ‘standard reference materials’ (SRMs), (Taylor, 1964; Bowen, 1979; Turekian and Wedepohl 1961; Wedepohl, 1995). SRM values are not site, or material, specific and the preferred methods of determining background are identification of a nearby ‘pristine environment’, or the use of sedimentary cores to obtain sediment from below the level of anthropogenic influence. Nevertheless, SRMs are still being used extensively and are entrenched in the literature (Pekey, 2006; Karbassi et al., 2008; Kabir et al., 2011).

The European Union has been concerned with the problem of establishing background metal values for their community for some time and the OSPAR Coordinated Environmental Monitoring Programme (CEMP) working group on monitoring (MON) have declared that core data will form the basis by which background is determined for the OSPAR region using only fine-grained and/or sieved samples (size normalised) to reduce the confounding of variable grain size and, if possible, samples should be taken from sediment older than 1850 AD (OSPAR, 2008). Using data from multiple studies, OSPAR (2005; 2008) established a single background metals dataset for the entire northeast Atlantic region. An investigation of background metal concentrations for 52 harbours and estuaries globally showed a remarkably narrow range of concentrations (Birch, 2016) with means of 27 µg/g, 33 µg/g and 97 µg/g for Cu, Pb and Zn, respectively. A similar study conducted as part of the current work, but exclusively for <63 µm data, gave values of 17 µg/g, 26 µg/g and 79 µg/g, respectively, which compared closely with the OSPAR concentrations of 20 µg/g, 25 µg/g and 90 µg/g, respectively for fine sediment. Background metal concentrations are strongly influenced by geology, climate and soil processes (Birch, 2018) and will be different for each of the WHP harbours. However, in the absence of local information on background concentrations, the OSPAR pre-anthropogenic values have been adopted in the present study in order that results be consistent between the 10 harbours being assessed and that WHP outcomes can be related to other global studies.

The second difficulty in assessing the magnitude of human-induced change in ports of the WHP, i. e. the absence of normalised metals data, was addressed using a unique elemental-normalisation procedure. Aluminium has been used as a normalising element (denominator) for some time (Rubia et al., 2000; Caeiro et al., 2005; Brady et al., 2015), however an increasing number of studies are using an assumption that sediment containing 100% clay mineral material has an Al concentration of 50,000 µg/g Al and that total metal concentrations can be normalised to this concentration of Al (OSPAR, 2005; 2008; Munksgaard et al., 2012; 2013; 2015; 2018). Further assumptions are that all metals are associated with the clay mineral phase and that the Al concentration used for normalisation (50,000 µg/g Al) was the same for all locations. These

assumptions were tested by comparing normalisation and enrichment determined by size-normalised core data with that determined using 50 000 µg/g Al for the Sydney estuary dataset. Pearson (product moment) correlation was used to determine the linear association between size-fractionated metals and metals normalized to various concentrations of Al. The relationship between Al and sediment size for the Sydney estuary dataset showed that a considerably lower concentration of Al corresponded to the composition of 100% clay mineral phase, i. e. 35,000 µg/g Al (and 35,000 µg/g for Fe, another normalising element). A unique opportunity to comprehensively study the relationship between Al (and Fe) and sediment size was made available by a large body of data obtained from 41 central NSW estuaries, which included sediment size, total sediment and size-normalised metals (Birch et al., 2016). Results from this study showed that the relationship between Al (and Fe) concentrations in total sediment and the fine fraction content was unique for each estuary. The procedure of choosing the appropriate Al concentration for normalisation is described in detail in Birch (2020). This understanding has led to a break-through in an ability to normalise metals data using unique Al normalising concentrations for each individual WHP harbour (the Canale Candiano of Ravenna and Auckland had neither Al and/or Fe) The use of Al and Fe in normalisation procedures should be used cautiously as Al enrichment in glacial sediments (Loring, 1991) and elevation of Fe by diagenetic processes, is well documented (Grant and Middleton, 1990; Whalley et al.,1999).

2.5 Indicators used to estimate the magnitude of anthropogenic change (MAC) (sedimentary metal enrichment)

A raft of environmental indicators is being used to assess human influence on marine ecosystems. However, terms, e. g. ‘contamination factor’, ‘concentration factor’ and ‘enrichment factor’ used synonymously in the literature need first to be defined for clarity. The terminology used in the present work combines terms promulgated by Caeiro et al. (2005) and Brady et al. (2015). ‘Concentration factors’ do not employ background, or normalisation procedures, ‘contamination factors’ involve pre-anthropogenic values, but do not incorporate normalised data, whereas ‘enrichment factors’ apply both background and some form of normalisation. Also important is whether enrichment indices are for multiple elements and whether the index is linked to a classification scheme. Assessment schemes used in the present study have been ranked based on these attributes in Table 3 and the formulae used to calculate the indices are presented in Supplementary Material.

TABLE 3

2.6 Ecological Risk Assessment (ERA) Risk of adverse effects to benthic populations posed by anthropogenic sedimentary chemicals

Chemical concentrations *per se* do not provide an effective means for determining potential adverse effects on benthic resources. To assess the ecological significance of contaminants bound

to sediments information on toxicity, bioaccumulation and effect on the structure of biological communities are needed. These measurements require a high degree of expertise, are time consuming and expensive and, as in the case of the WHP, are frequently not available. Instead, sediment quality guidelines (SQG) are commonly used to make preliminary assessments of sediment toxicity when direct biological effects information is unavailable. Empirical methods, employing matching sediment chemistry and biological effects data, have been used in development of SQGs for determining adverse outcomes of contaminants on ecological populations using total sediment chemistry. Numerical-effects based SQGs are now in common use globally as a screening management tool to identify and prioritise contaminants and regions of concern.

There are several sediment quality effects-based guidelines commonly in use, e. g. the apparent-effects threshold (AET) scheme and screening level concentrations (SLC) (Birch, 2018). However, the most commonly used SQG for estuarine and marine environments is the US National Oceanic and Atmospheric Administration (NOAA) scheme based on concurrent sediment chemical and ecological data from the laboratory and field for a variety of techniques and benthic end points (Long and Morgan, 1990; Long et al., 1995; MacDonald et al., 1996). The scheme comprises two observed ecological effects concentrations, i. e. the effects range low (ERL) and the effects range median (ERM). The former level identifies the concentration below which adverse ecological effects are seldom observed and the latter level distinguishes concentrations above which adverse ecological effects occur frequently. Concentrations between the two levels exhibit irregular ecological response. A wide range of chemicals, including organic and metallic contaminants, has been incorporated into these SQGs and the approach is now well established in North America and in many countries in Europe, Asia, South America and Africa.

Contaminants do not occur as single chemicals within marine sediments and a number of schemes have been developed to assess the effects of chemical mixtures for aquatic sediments. The mean ERM quotient (MERMQ) scheme has been used to estimate adverse ecological effects of chemical mixtures in this WHP (Long and MacDonald, 1998). The MERMQ method requires normalising the concentration of each chemical with respect to its ERM value, summing the quotients for each substance and dividing by the number of chemicals for which guidelines are being used. MERMQ ranges of >1.5; 1.5-0.5; 0.5-0.1 and <0.1 have been related to the probability of toxicity (76%, 49%, 21% and 9%, respectively) in amphipod assemblages. The number of ERL and ERM exceedances has also been related to toxicity through whole sediment bioassays. However, these toxicity relationships should be used with caution due to area-specific nature of benthic populations and sedimentary chemicals and instead the MERMQ is considered in the current assessment as a level of risk of adverse effects to sediment-dwelling animals, rather than as a probability of toxicity.

3. RESULTS

For harbours with adequate sample density (Darwin, Derwent, Hong Kong, Ria de Vigo, Rio de Janeiro and Sydney) figures are presented as filled contour maps (Figs. 2-8) and for the remaining locations (Auckland, Dublin, Ravenna and Santander) distributions are given as points

in Supplementary Material (Figs. S1-S7). Because distributions of most elements being considered in this work co-occur, only Zn is depicted in figures to limit the number of diagrams. Total metal concentrations (Table 4, Fig. S8), enrichment (Table 5, Fig. S9), contamination and enrichment indices/factors (Table 6) and ecological risk (Tables 7 and 8, Fig. 9) are described for each of the WHP locations below.

Auckland Harbour

Mean and maximum total sediment metal concentrations were low and were highest in tributaries in the south-west and west.

Concentration (CF) and contamination factors were low indicating no to slight contamination, however metal enrichment could not be estimated as no sediment size data, or total Al/Fe data were available.

No sampled areas exceeded ERL concentrations for Cu and Pb, while samples only exceeded this guideline for Zn in the upper reaches of tributaries in a small part of the southern estuary. Ecological risk is minimal (MERMQ=0.09).

TABLE 4

Darwin Harbour

Total mean and maximum metal concentrations were low for sediments mantling Darwin Harbour and were greatest in tributaries of Palmerston and Darwin, as well as along the coastal fringe towards the southeast. Copper total sediment concentrations were particularly low with mean and maximum levels of 5.4 µg/g and 23 µg/g, respectively.

Post-extraction normalised (PEN) data for Darwin Harbour showed low mean concentrations for Cu, Pb and Zn (16 µg/g, 29 µg/g and 79 µg/g, respectively) and a large number of samples were at, or close to background levels. Low PEN metal concentrations may be due to high and variable carbonate content in surficial sediments (Munksgaard et al., 3012).

Concentration and contamination factors were low, suggesting no to slight contamination. Enrichment values indicated sediments were generally uncontaminated with mean values of 0.9, 1.4 and 1.0 for Cu, Pb and Zn, respectively, except for the tidal flat sediment near Darwin City and port where enrichment was generally >2.5.

Maximum total sediment concentrations of Cr (95 µg/g), Cu (23 µg/g), Ni (27 µg/g), Pb (50 µg/g) and Zn (190 µg/g) exceeded ERL values by small margins. Metals showed a similar spatial distribution and no metals in any area of the harbour exceeded ERM concentrations. Overall ecological risk (MERMQ=0.06) was minimal.

TABLE 5

Derwent River

Mean and maximum total sediment Cd (14 µg/g and 128 µg/g, respectively), Pb (450 µg/g and 1880 µg/g, respectively) and Zn (2130 µg/g and 14600 µg/g, respectively) concentrations were extremely high and the maximum concentration for Zn is possibly the highest recorded. Total sediment, mean and maximum Cu concentrations were moderately high and Cr and Ni concentrations were slightly elevated. A very strong down-stream gradient was apparent for Cd, Cu, Pb and Zn with maximum concentrations centered on the Glenorchy area.

In the absence of total Al, normalisation was accomplished using 40K Fe as a normalising agent. Normalised metal concentrations were only moderately higher (~25%) than total concentrations due to the mainly muddy nature of bottom sediments. All environmental indicators were highest for Derwent River sediments in the WHP, indicating extreme to severe enrichment. Mean enrichment quotient, especially for Cd (77), Pb (19) and Zn (23) was also extremely high, resulting in a MEQ of 21.

Sediment in most of the estuary exceeded ERL concentrations for Pb and Zn and for large areas for Cu. Sediment in the Glenorchy area exceeded ERM concentrations for Cu and over large parts of the estuary sediment exceeded ERM values for Cd, Pb and Zn. Similar distributions of risk were presented by Pb and Zn distributions, except that Zn ERM concentrations extended further seawards than did Pb. High risk for Cd (1.40), Pb (2.04) and Zn (5.2) were moderated by low risk levels for Cr (0.10) and Ni (0.32) to give an overall risk of MERMQ=1.58.

Dublin Port

The total sediment metal concentrations reported for three vintages (2006, 2008 and 2013) of data for Dublin Port varied greatly. Vintages 2006 and 2008 had similar spatial distributions covering most of the harbour area, while 2013 data were mainly confined to the Alexandra Basin loading dock. Metal total sediment concentrations in 2006 (Cu, Pb and Zn concentrations were 49 µg/g, 81 µg/g and 217 µg/g, respectively) were substantially higher than in 2008 (mean 27 µg/g, 44 µg/g and 152 µg/g, respectively), but lower than in Alexandra dock in 2013 (mean 60 µg/g, 138 µg/g and 663 µg/g, respectively). In particular, the 2006 vintage data were normalised using 25K Al (mean Al 15,900) resulting in increased enrichment and a MEQ of 4.6. However, for the 2008 survey which covered the majority of the harbour, sediment metal concentrations exceeded ERL values only in a minor part of the inner harbour, while one site exceeded ERM values for Pb and Zn. As a result, the overall risk for this vintage was low and the MERMQ was 0.31. Concentrations of Cu, Pb and Zn displayed similar distribution patterns, i. e. increasing towards the inner harbour with a moderate elevation in a dock in the central harbour area (based on 2008 data).

These apparent high temporal fluctuations in contaminant levels for the 2006 and 2008 data from the same harbour area may have been due to changes in sediment size (ship turbulence, fluvial- or tidal-derived currents) as indicated by different mean Al concentrations. Or may be due in part to more stricter controls on inputs both within the Port/Harbour environ and upstream along the River Liffey catchment area (Brooks et al., 2016). Substantial spatial variance and high concentrations in the 2013 data may have been related to small-scale variability (debris from loading vessels), or the variance may have been analytical. The Alexandra basin has a long history of ship-building and vessel cleaning and maintenance and it is deemed the most likely source for the high concentrations found in sediments within this area (Brooks et al., 2016). Indeed, due to proposed developmental plans within the Port (most of which is to occur within the basin area – DP, 2014), it may be that this sample was selected to ascertain how contaminated sediments were so that appropriate precautions for handling could be developed prior to commencement of Port developments (DPC, 2014).

Hong Kong Harbour

Total sediment metal concentrations for Hong Kong coastline and Hong Kong Harbours were low, while maximum concentrations were also moderately low. Copper, Pb and Zn displayed similar spatial distributions and were more elevated in the harbour than for the adjacent coastline. Metal concentrations decreased rapidly seawards and were very low in blue water regions.

Normalisation was conducted using 35K Al, which increased total concentrations moderately. Total metal/total Al plots suggested some sites were slightly contaminated and total Al and total Fe were closely related ($r=0.865$). Environmental indicators showed Hong Kong Harbour sediments to be slightly enriched. Mean enrichment quotient, especially for Cu was moderate for coastal samples (mean 5.1 and maximum 8.8) and mean enrichment for Pb and Zn in the harbour and coastal region was <3.0 .

Ecological risk in all areas was low and only in the Hong Kong Harbour area do any metal exceed ERL values, while no ERM concentrations were exceeded. Overall ecological risk is low (MERMQ=0.20) and most elevated for Zn (0.36) followed by Cu (0.25).

Ravenna

Sediments in the Canale Candiano were moderately rich in total mean Cd and Pb concentrations (1.6 $\mu\text{g/g}$ and 49 $\mu\text{g/g}$, respectively) and metal concentrations increased towards the city of Ravenna. Concentration and contamination indicators were slightly elevated, however no enrichment factors could be determined due to an absence of sediment size data and normalising elements for the Canale Candiano. Ecological risk was low (MERMQ=0.26) with Zn posing the highest concern (MERM=0.43).

Rio de Janeiro

Rio de Janeiro Bay was almost entirely mantled in muddy sediments ($>80\%$) and total sediment metals were moderately high for Cu and Pb (means 62 $\mu\text{g/g}$ and 66 $\mu\text{g/g}$, respectively) and high

for Zn (mean 318 µg/g). All metals increased towards the SW and were greatest adjacent to Rio de Janeiro city and the Rio Pauvna and lowest in the sandy sediment off San Francisco Beach in the SE. Maximum concentration for Zn was third highest (2039 µg/g) for the 10 harbours studied.

Total sediment was normalised to 20K Al increasing total concentrations by approximately 30% for all metals. Total Al/total metal plots indicated a substantial number of contaminated samples. Environmental factors suggested slight to moderate enrichment. Mean enrichment was moderate for Cu, Pb and Zn, i. e. 4.0, 3.1 and 4.6, respectively resulting in a MEQ of 3.5 (moderate enrichment).

Sediments in the NE and SE of Rio de Janeiro Bay posed no risk to benthic resources with total metal concentrations <ERL. For Cu and Pb sediments mantling the remainder of the bay had an intermediate risk of adverse effects with total concentrations >ERL<ERM. Sediments posed a high risk due to total Zn concentrations (>ERM) adjacent to the city and off Rio Pavuna. Overall ecological risk was slight (MERMQ=0.33).

Ria de Vigo estuary

Mean Cr (70 µg/g), Cu (82 µg/g) and Ni (29 µg/g) concentrations were moderate, while maximum concentrations (198 µg/g, 479 µg/g and 43 µg/g, respectively) for these metals were reasonably high. Chromium and Zn outliers increased maximum concentrations for these metals. Sedimentary Zn concentrations displayed a distinctive decreasing gradient away from the main harbour. Copper showed a similar pattern to Zn, but trends were less strong, nevertheless concentrations were clearly elevated at the harbour, while Pb concentrations increased regularly up estuary due to discharges from a ceramic factory located at the head of the estuary (in operation since the late 1960s and closed in 2001). Secondary Pb inputs are attributed to industrial and port activities. The harbour was a significant source of Zn and Cu, and possibly Pb to the estuary.

Normalised metal concentrations were only marginally higher than total values due to a consistent muddy substrate. Contamination factors showed slight to moderate elevation and enrichment was highest for Cu (mean 4.9) and Pb (mean 5.2) with a MEQ of 4.1.

Mean sediment Cu, Ni and Pb concentrations were greater than ERL values and maximum Cu, and Zn concentrations exceeded ERM values. Ecological risk for sediments was high for Zn in the vicinity of the harbour (concentrations >ERM), while risk was moderate for Cu and Pb for most of the estuary with samples exceeding ERL values. Overall ecological risk was slight (MERMQ=0.38).

Santander Harbour

Sediments in Santander Harbour contained the second highest total Cr (83 µg/g) and Ni (48 µg/g) concentrations in the WHP. Lead and Zn sediment metal concentrations were high in the

dockland area west of the harbour and Pb and Cu concentrations were also elevated in the embayment to the south. An apparent decreasing metals gradient from the upper embayment in the south towards the harbor mouth in the northeast requires verification with additional sampling.

Normalisation was undertaken using 27K Al. Although almost all samples in the 2015 vintage database were enriched (>2.5), contamination was not apparent in the total metals/total Al plots due probably the lack of uncontaminated samples. Environmental factors suggested moderate to high modification and enrichment was greatest for Cd and Zn (means 9.9 and 9.5, respectively), high for Pb (mean 5.2) and moderate for Cu (mean 2.5) and mean enrichment was $MEQ > 5$.

Mean total Cr, Ni and Pb concentrations exceeded ERL values and mean Zn concentrations were $>ERM$. With a spatially limited dataset it was difficult to determine accurately the areas exposed to ecological risk by sediments, however it appeared that large parts of the port may be at risk for Cu, Pb and Zn. Overall ecological risk was moderate ($MERMQ = 0.51$). Additional sampling and mapping is required to verify the risk distribution.

Sydney estuary

Total sediment mean (133 $\mu\text{g/g}$) and maximum (1060 $\mu\text{g/g}$) Cu concentrations were the highest recorded in the current study and maximum concentrations were highest for Pb (1932 $\mu\text{g/g}$) and second highest for Cr (298 $\mu\text{g/g}$) and Zn (11300 $\mu\text{g/g}$), while minimum values were commonly below detection due to sandy substrate in parts of the harbour. Sedimentary metal concentrations declined markedly from the upper reaches of the estuary towards the mouth and with distance from stormwater discharge points at the headwaters of offchannel embayments and tributaries. Sediments of the four south, central embayments of Blackwattle/Rozelle Bay, Iron Cove, Hen and Chicken Bay and Homebush Bay consistently contained the highest concentration of metals. Moderate metal concentrations were located in the western embayments of Middle Harbour.

Individual embayments had distinctive metal distributions in Sydney estuary (Birch et al., 2015a, b). Sediments in Homebush Bay generally had high Pb levels related to paint manufacturing, whereas surficial sediments in Hen and Chicken Bay had high Cu concentrations originating from a bronze processing plant. Sediments of Iron Cove were elevated in Pb and Cd due to historical industrial discharge and Blackwattle/Rozelle Bay sediments were highly enriched in Cu, Pb and Zn from shoreline heavy industry. Chromium was anomalously high in sediments of some bays in northwest Middle Harbour and Lane Cove related to tanning industries.

Size-normalised data were available for Sydney estuary, which allowed enrichment to be determined directly using surficial sediment concentrations and size-normalised OSPAR background metal concentrations without the use of elemental normalisation. Enrichment factors indicated moderate to high modification and enrichment was highest in the WHP for Cu (9.1) and third highest for Cd (5.1), however MEQs (8.6) were reduced by low enrichment for Cr (2.0) and Ni (0.7), especially when six elements are considered (5.7).

Copper, Pb and Zn are the contaminants of most concern in sediments of Sydney estuary and areas of the waterway with sediment exceeding ERM concentrations for these metals represented approximately 2%, 50%, and 36% of the estuary, respectively (Birch and Taylor, 2002a, b, c). Sediment in the entire estuary, except a small area near the entrance, exceeded ERL concentrations for at least one metal. Overall ecological risk was moderate (MERMQ=0.53).

4. Discussion

4.1 Sampling and Analytical methods

Sediment samples used by WHP institutions were recovered by grab (van Veen), corer or box corer. When using these different sampling devices, it is important to remove only the uppermost sediment layer so as to sample only the most recently deposited material and not to mix this surficial material with underlying, pre-anthropogenic substrate. Sampling design, density and distribution needs to be consistent. However, sampling was frequently focused on perceived source locations and points of interest with low density cover over the remaining (sometimes majority) waterway preventing a full spatial assessment. Sample density should be relative to small-scale spatial variance and proximity to discharge locations to provide optimal coverage for source identification and dispersion tracking. Sample density, which provided satisfactory regional coverage for reasonably consistent abundances was approximately 0.5-1.0 samples/km², but increased to 5-10 samples/km² in areas of interest, or in places of high variability based on data provided in the current study.

Metals in the sedimentary environment are present in the matrix of minerals and as the absorb phase of mainly fine-grained particles. The method chosen for chemical analysis of anthropogenic chemicals requires that metals from the mineral matrix be excluded from the analysis, especially as some sedimentary minerals contain high concentrations of metals incorporated in the structure. This is especially important in the coastal environment where marine and terrestrial sediments are immature and commonly contain metal-rich matrix minerals. Analytical schemes that result in assessment of both the absorbed and matrix phases confound interpretation and identification of anthropogenic contribution to the sediment. The approach used to analyse metals in sediments is therefore fundamentally important in assessment of sediment condition (Table 1). Weak acids have the advantage of providing an estimate of the trace metal bioavailable fraction (1M HCl) (Ying et al., 1982) and may be used in assessing potential toxicity (6 mol/L HCl solution for acid-volatile sulphides - simultaneously extracted metals, AVS-SEM analysis) (Di Toro et al., 1990; 1992). Strong acid digestions (HF) break down minerals and releases both matrix and adsorbed components resulting in a 4- to 9-fold elevation of metal concentrations compared to the more frequently used aqua regia (Katz and Kaplan, 1981), whereas the dilute HCl solutions only recover approximately 60% of metals relative to aqua regia. Digestion procedures used in the WHP varied from weak to strong acids, which may have resulted in a mixed proportion of matrix and adsorbed metals in the analyses.

Most analyses undertaken in the WHP were by ICP, either OES, or MS, which would result in a high level of accuracy and precision, especially as most laboratories used International Reference Materials and appropriate QA/QA procedures. Not all studies incorporated sediment size, Al and Fe in the analytical stream, limiting an ability to normalise data needed for enrichment determinations.

Preferably, unconsolidated sediments should be chemically characterised using a wide variety of analytes, including metals and a range of organic contaminants, including organochlorine pesticides (OCs), polycyclic aromatic hydrocarbons (PAHs) and polychlorinated biphenyls (PCBs). However, inconsistent organic chemical data and gaps in metal analyses across the WHP dataset prohibited such a holistic approach in the current investigation. Instead, metals that were ubiquitous in the WHP dataset were used to determine the magnitude of human induced change and possible ecological stress resulting from contamination. Although the absence of organic contaminant data in the present work is a disadvantage, the extent to which metals reflect the distribution of other priority contaminants of concern (OCs, PAHs, PCBs) in estuarine environments is also surprisingly consistent. The correlation between three metals (Cu, Pb and Zn) and OCs, PCBs and PAHs in Sydney estuary was significant ($r=0.793$, $p<0.05$) and with the same metals and nine OCs and hexachlorobenzene from Hawkesbury River (NSW) $r=0.866$ ($p<0.05$) and the same metals with PCBs from New York Harbour $r=0.878$ ($p<0.05$) and with sediments from San Diego Bay $r=0.655$ ($p<0.05$) (Birch et al., 2008). In the current study, Cu, Pb and Zn correlated with each other and with six metals (Cd, Cr, Cu, Ni, Pb and Zn) ($r=0.997$, $p<0.05$) and with PAHs in sediments of the Ria de Vigo Harbour ($r=0.769$, $p<0.05$). Estuarine sedimentary contaminants will more likely to covary in regions dominated by stormwater discharge because the chemical mix of urban stormwater is reasonably consistent (NURP, 1983; US EPA, 1983; Fletcher et al, 2004), however in areas receiving point-source, chemical-specific industrial discharge, contamination may be less likely to covary spatially.

4.2 Magnitude of anthropogenic change

The magnitude of anthropogenic change followed total metal concentrations with Derwent River having the highest Cd, Pb and Zn enrichment and Sydney the highest Cu enrichment, while Santander sediments were most enriched by Cr and Ni. The overall magnitude of anthropogenic change (MEQ) was high for Derwent River, Santander and Sydney estuary (>5.0), moderate for Rio de Janeiro (3.0-5.0), slight for Dublin, Hong Kong and Ria de Vigo (1.5-3.0) and minimal for Darwin (<1.5) (no data were available for Canale Candiano in Ravenna and Auckland) (Table 5). This enrichment appeared to be more related to location and magnitude of source than the morphodynamic characteristics of the WHP harbour estuaries.

Enrichment (MEQ) was calculated for six metals ($MEQ/6=Cd, Cr, Cu, Ni, Pb$ and Zn) and for three metals ($MEQ/3=Cu, Pb$ and Zn) to determine whether the extra number of elements effected enrichment calculations (Table 5). A close correlation between the two data sets ($r=0.953$, $p<0.05$) supports similar results, which show metals are invariably closely correlated

with each other, e. g. the three metals (Cu, Pb, and Zn) were closely correlated with a suite of six metals for Sydney estuary ($r = 0.959$, $p < 0.05$), Hawkesbury River ($r = 0.932$, $p < 0.05$), New York Harbour ($r = 0.843$, $p < 0.05$) and for Tampa Bay ($r = 0.825$, $p < 0.05$) (Birch et al., 2008) and in many other locations (Zhang et al., 2009).

Enrichment has been calculated in the present work based on a novel elemental normalisation procedure employing variable concentrations of Al to accommodate for regional changes in local clay mineral chemical assemblages. This approach has been investigated recently in great detail (Birch, 2020) and is discussed here only in its application to the WHP. The concentration of the Al normaliser varied across the 10 WHP datasets from 20K Al (Rio de Janeiro) to 70 K Al (Darwin) and was most commonly between 32K and 35K Al. The normaliser value used was estimated from the relationship between sediment size and total Al concentration, which was available for six (Darwin, Derwent, Hong Kong, Rio de Janeiro, Ria de Vigo and Sydney) of the 10 harbours. For harbours without sediment size information, normaliser values were estimated from total Al alone, which introduced speculation and no normalisation could be undertaken for harbours without sediment size, Al, or Fe data (Auckland and Canale Candiano in Ravenna).

An opportunity to test the validity of normalisation and enrichment determined by elemental normalisation (Al and Fe) was afforded by the availability of size-fractionated metals data from Sydney estuary. Enrichment based on size-normalised data and 35K Al normalisation (employing SOPAR (2008) background values for both data sets) was closely related. i. e. 9.1 and 9.8, respectively for Cu, 11 and 13, respectively for Pb and 6.5 and 7.6, respectively for Zn (Table 5). Moreover, enrichment determined by sediment size- and Al normalisation was also closely correlated. i. e. $r = 0.921$ ($p < 0.05$) for Cu and $r = 0.854$ ($p < 0.05$) for Zn and less so for Pb ($r = 0.516$, $p < 0.05$), which showed moderate scatter due to analytical difficulties. These results generate confidence in the approach being used in the current work based on variable Al elemental normalisation.

The availability of PEN data for Darwin Harbour allowed a comparison between another size-normalisation process and elemental normalisation. Normalisation of Darwin Harbour data to 70K Al produced similar results to PEN normalisation, i. e. mean concentrations of 15 $\mu\text{g/g}$, 29 $\mu\text{g/g}$ and 79 $\mu\text{g/g}$ compared to 18 $\mu\text{g/g}$, 36 $\mu\text{g/g}$ and 87 $\mu\text{g/g}$ for Cu, Pb and Zn, respectively. Normalisation using Fe produced inconsistent results due to anomalously high concentrations of this element for some samples related to the presence of Fe oxyhydroxides (Munksgaard et al., 2013). Mean enrichment, also using 70K Al as a normaliser, was closely correlated, i. e. 0.9, 1.4 and 1.0 for Cu, Pb and Zn, respectively compared to 0.7, 1.0 and 0.9, respectively for PEN data (Table 5). Enrichment using PEN and 70K Al data was moderately correlated ($r = 0.641$, $p < 0.05$), however PEN data were less consistent than using 70K Al possibly due to variable carbonate content.

Although Li is the normaliser of choice for Ria de Vigo sediments, these data afforded the opportunity to test the appropriateness of Al- and Fe-normalisation on the same data set. Normalised concentrations produced by 90K Al and 35K Fe were remarkably similar, i. e. 98 $\mu\text{g/g}$ and 97 $\mu\text{g/g}$, respectively for Cu, 131 $\mu\text{g/g}$ and 125 $\mu\text{g/g}$, respectively for Pb, and 244 and 235, respectively for Zn. Similarities in enrichment determined by these two techniques were

also close, i. e. 4.9 and 4.9 for Cu; 5.2 and 5.0 for Pb and 2.9 and 2.7 for Zn. MEQ for the two approaches were both 4.1 and correlation between the two techniques for Cu was $r=0.964$ ($p<0.05$). These results indicated both Al and Fe may be used as elemental normalisers in the absence of diagenetic modification.

Determining the most appropriate pre-anthropogenic metal concentrations was the second difficulty in assessing enrichment in the WHP. The lack of these data for individual harbours reduced the ability to determine the magnitude of anthropogenic change. In the absence of data derived from sedimentary cores, or from local pristine environments, OSPAR (2008) background metal values were adopted in the present WHP study. Various Al concentrations were used in the normalisation process to accommodate for changes in local clay mineral characteristics however, a single suite of background metal concentrations was applied to all WHP data. Background values will be different for each of the world harbours and should be estimated from local fine-grained down-hole core data, or pristine fine sediment. The absence of size-normalised sediment metal concentrations and valid local background data will introduce errors in enrichment determinations for the WHP.

4.3 Use of concentration, contamination and enrichment factors in assessment of anthropogenic change

Derwent River and Sydney estuary registered high values for all 10 enrichment indices used in the current investigation (Table 6), while Auckland and Darwin resulted in low assessments for most of these tools. Other harbours had mixed values across the spectrum of indices with Ria de Vigo, Rio de Janeiro and Santander having slightly elevated outcomes. An attempt has been made to combine the outcomes of assessment factors calculated in the current work to give an overall quantification for the 10 estuaries in the WHP (Table 6). The classification schemes for the EF (Rubio et al., 2000), mDC (Abraham and Parker, 2008), mPI (Brady et al., 2015) and MEQ (Birch and Olmos 2008) techniques are substantially similar providing a unique opportunity to produce an overall classification scheme for these enrichment factors. Enrichment factors of EF <1.5; 1.5-3; 3 to 5; 5-10; 10 -25; > 25 were classified as not enriched; slightly enriched; moderately enriched; highly enriched; extremely enriched; and severely enriched, respectively. Based on this approach, the overall score for the Derwent River was 20, for Sydney estuary 14, for Santander 12, for Ria de Vigo and Dublin 11, for Rio de Janeiro 9, for Hong Kong 5 and for Darwin 4. Insufficient normalised data were available to rank Canale Candiano of Ravenna and Auckland, however these harbours probably have scores of <4 considering the low total metal concentrations.

A generalised observation of the 10 metrics used in the current study (Table 6) showed that the Nemerow Pollution Index (*PI*) and the Metal Pollution Index (*MPI*) values were high, while the Geo-accumulation index (*I_{geo}*) for both total sediment and for Al-normalised data were low with respect to results from other indices. PI values were higher than reported in the literature (Brady et al., 2015) possibly due to unusually elevated metal concentration in some of the WHP

estuaries, whereas *Igeo* figures are commonly cited low, or negative (Abraham and Parker, 2008). The Surface Enrichment Factor (*SEF*), Enrichment Factor (*EF*), the Modified Degree of Contamination (*mDC*) and the Mean Enrichment Quotient (*MEQ*) results are similar. An outcome of these assessments was that, although the factors are based on different attributes, the ranking produced was similar for all assessment tools. A further outcome was that the Concentration Factor (*CF*) (un-normalised data), *mDC*, *EF*, *mPI* and the *MEQ* are closely correlated (*r* values between 0.942 and 0.959, $p < 0.05$), while the *Igeo* and the *PI* are poorly correlated with the previous group of factors, probably due to the different basis on which these indicators are calculated.

TABLE 6

Estimation of metal background concentrations has a substantial influence in the determination of anthropogenic change established by contamination and enrichment factors. Four types of background value were used in the study of Ria de Vigo (Northwest Spain) (Rubio et al., 2000). One background value was derived from local geology, a second from global shale and two further background values were based on regional studies. An assessment of contamination varied substantially when each of these background concentrations were used to assess contamination. Abraham and Parker (2008) found that using continental shale Fe values as a normalising element resulted in a significant increase in *EF* compared to using Fe concentrations from the base of local cores and warned that using global material as background to calculate enrichment should be undertaken with caution. Background metal concentrations used in calculating the Degree of Contamination (*DC*) were derived from pristine sediments by Hakanson (1980) based on sediment from 50 lakes in Europe and North America already influenced to varying degrees by anthropogenic activity and also possibly by variable grain sizes. Background values were determined by adding one standard deviation to the mean, which resulted in doubling of the final background concentrations due to considerable variation in the data. Resulting background concentrations are high relative to other reported pre-anthropogenic values, nevertheless this technique remains in use globally and is frequently cited in the literature.

The original Geo-accumulation Index (*Igeo*) (Müller 1969, 1979; 1986) was based on fine sediment samples and background metal concentrations, however recently different backgrounds have been used (global shale) (Ghani et al., 2013) and various sediment sizes (total sediment) have been included in the computation (Buruaem et al., 2012; Pang et al., 2015). resulting in confusion and incompatible outcomes. More recently confounding due to variable size when using total sediment has been taken into account by normalising to Al and adjusting the data to 100 % mud (Kim et al., 2018; 2019). Geo-accumulation index values were low and often negative in the current work, similar to other studies where results for most elements were also negative, e. g. Cevik et al. (2009); Kaushik et al. (2009); Thuong et al. (2013), Abraham and Parker, (2008) and Kim et al. (2018).

The original Degree of Contamination (DC) included seven specific metals (As, Cd, Cu, Cr, Hg, Pb, and Zn) and an organic pollutant (PCB) and required a minimum of five samples. The numeric sum of the eight specific contamination factors expressed overall degree of contamination and all eight contaminants had to be included in the calculation. The limited and specific number of pollutants led Abraham and Parker (2008) to modify the factor to include any number of metallic contaminants and the analysis of at least three samples of impacted sediments. Background values were determined from the lower sections of cores and the six-division classification scheme was modified accordingly. Abraham and Parker (2008) restricted examination to fine-grained samples for both contemporary and background materials.

Single-element pollution indicators present a number of limitations and do not take into account the complex interaction of metal contamination in mixed urban and industrial environments. These limitations have led to the development of multi-element indices, e. g. *mPI*, *DC* and *mDC*, which include a suite of metals to make a more integrated assessment of contamination.

The skewed nature of some contaminant data has also led to modification of the *PI* index. In the case where one metal is highly enriched and the calculation is averaged over a suite of metals, the impact of the enriched element is subdued. This problem has been addressed by including the maximum concentration of the elevated element as a separate factor in the weighted-average value. The *mPI* also uses enrichment factors, which accounts for the non-conservative nature of contaminated sediments. However, no guidance is provided of how to identify an ‘enriched’ element, or how to conduct the calculation if there is not one. In the WHP datasets, Zn was always the highest concentration, however it not clear whether this element should be considered ‘anomalously’ elevated and included as a separate factor, or not in the computation. Inclusion of Zn as a separate factor in the current study resulted in an over estimation of *mPI* values, e. g. in the Derwent River and Sydney estuary, and production of values that are far in excess of other assessment types in the current study and of *mPI* results in others work (Brady et al., 2015).

4.4 Ecological risk posed by sedimentary metals

Only sediment from Derwent River was rated high risk ($MERMQ > 1.5$), followed by Sydney and Santander estuaries at moderate risk ($MERMQ > 0.50$). Auckland and Darwin sediments exhibited minimal risk and sediment in the remaining harbours (Dublin, Hong Kong, Ravenna, Ria de Vigo and Rio de Janeiro) was assessed at slight risk to benthic communities (Table 7). The statistic that separates Derwent River from other harbours is the number of samples with at least one element with concentrations > 3 times ERM values (51 samples, 45%) and > 5 times ERM levels (40 samples, 36%) (Table 8). Ecological risk for sediments of the Derwent River are driven by high Cd, Pb and Zn concentrations, Sydney by high Pb and Zn concentrations and Santander by Ni and Zn concentrations (Table 8).

TABLE 7

A more detailed examination of harbours exposed to slight and minimal risk shows Auckland and Darwin with only a few samples >ERL for any one metal (Table 8). Harbours mantled with sediment assessed at slight risk of adverse effects (Dublin, Hong Kong, Ravenna, Ria de Vigo and Rio de Janeiro) have a high proportion (54 % - 88%) of samples with at least one metal >ERL values, suggesting possible risk.

TABLE 8

The second popular effects-based sediment quality guideline for single contaminants, i. e. the Probable Effects Level (PEL) (MacDonald et al., 2000) and for chemical mixtures, i. e. the mean PEL quotient (MPELQ) provide the highest values the Derwent River (1.34), followed by Sydney 0.97) and Santander (0.77), similar to results produced by the MERMQ. The MERMQ for three metals (Cu, Pb and Zn) and for six metals are consistent, except for Sydney where the MERMQ for six metals has been reduced by low Cr and Ni values.

A more extensive evaluation of the highly impacted harbours was made by assessing areas and proportions of harbours adversely affected by sedimentary metals. Greater than 80% of the Derwent River and Sydney estuary are mantled in sediment enriched >5 times over pre-anthropogenic times, while Rio de Janeiro sediments exhibit a large range of enrichments. Sediments in Hong Kong and Ria de Vigo Harbours are mainly enriched between 1.5 and 5 times. Sediments in more than 25% of Derwent River are at high ecological risk (MERMQ>1.5). while only 2% of Sydney estuary was at this risk level. Over 90% of the area of the remaining harbours (Hong Kong, Ria de Vigo and Rio de Janeiro) had a slight to moderate ecological risk (MERMQ = 0.1 - 0,5)

4.5 Overall magnitude of anthropogenic change and ecological risk

An overall assessment of anthropogenic change and ecological risk has been undertaken by ranking enrichment (Table 5), environmental indices (Table 6) and MERMQ (Table 7) for eight harbours (enrichment data were not available for Auckland and Canale Candiano of Ravenna) (Table 9). Ranking was remarkably consistent across the three schemes, i. e. the Derwent River, Sydney and Santander estuaries were placed first, second and third most impacted environments, respectively for all assessments, while Darwin Harbour was the least influenced by human activities. The remaining harbours changed only one or two places between schemes.

TABLE 9

991

992 A new categorisation scheme (Birch, 2018) has been applied to results of enrichment and
993 ecological risk obtained in the current study to assess overall anthropogenic change and
994 ecological risk (Table 10). Derwent River, Sydney and Santander estuaries are highly enriched,
995 while Rio de Janeiro is moderately enriched. Dublin, Hong Kong and Ria de Vigo are slightly
996 enriched, whereas Darwin is not enriched. Only the Derwent River is at high ecological risk,
997 while Sydney and Santander estuaries are at moderate risk. Auckland and Darwin are at minimal
998 risk and Dublin, Hong Kong, Ravenna Ria de Vigo and Rio de Janeiro are at slight ecological
999 risk.

1000

1001 TABLE 10

1002

1003

1004 5 Conclusions

1005 Sediments of Derwent River are distinctive with very high total mean concentrations of Cd, Pb
1006 and Zn. Sydney estuary sediments contain the highest mean Cu concentrations and Santander
1007 Harbour sediments are characterized by the highest mean concentrations of Cr and Ni of the 10
1008 WHP ports.

1009 An innovative technique for Al and Fe normalisation was applied and tested against accepted
1010 size-normalisation and post-extraction (PEN) methods, which satisfactorily allowed
1011 measurement of anthropogenic change. Metals enrichment followed total metal concentration
1012 and the mean enrichment for multiple sedimentary metals (MEQ) was high for Derwent River,
1013 Santander Harbour and Sydney estuary (>5.0), moderate for Rio de Janeiro and Dublin Port (3.0-
1014 5.0), slight for Hong Kong (1.5-3.0) and minimal for Darwin (<0.15) (no sediment size. Al, or Fe
1015 data were available for Canale Candiano of Ravenna and Auckland.

1016 Only sediment from Derwent River was rated at high ecological risk ($MERMQ > 1.5$), followed
1017 by Sydney and Santander estuaries with moderate risk ($MERMQ > 0.50$). Dublin, Hong Kong,
1018 Ravenna, Ria de Vigo and Rio de Janeiro were assessed at slight risk to benthic communities and
1019 Auckland and Darwin sediments exhibited minimal ecological risk.

1020 All 10 enrichment indices used in the current investigation showed elevated values for Derwent
1021 River and Sydney estuary, while Auckland and Darwin registered low values for most of these
1022 tools. Other harbours had mixed values across the spectrum of indices. A unique, multi-index
1023 classification scheme provided an overall score of 20, 14, 12, 11, 9, 5 and 4 for the Derwent
1024 River, Sydney estuary, Santander, Ria de Vigo and Dublin, Rio de Janeiro, Hong Kong and
1025 Darwin, respectively. Insufficient data were available to rank Canale Candiano of Ravenna and
1026 Auckland, however these harbours are considered to score <4 considering total metal
1027 concentrations.

A new categorisation scheme applied to results of the current study assessed overall anthropogenic change and ecological risk. Derwent River, Sydney and Santander estuaries were rated highly enriched, while Rio de Janeiro was moderately enriched. Dublin, Hong Kong and Ria de Vigo were slightly enriched, whereas Darwin was not enriched. Only the Derwent River was at high ecological risk, while Sydney and Santander estuaries were at moderate risk. Dublin, Hong Kong, Ravenna Ria de Vigo and Rio de Janeiro were at slight ecological risk and Auckland and Darwin were at minimal risk. The similarity in the ranking of the harbours in assessment of enrichment and ecological risk is reassuring and the minor difference for Rio de Janeiro is due to the different criteria used in the two assessment techniques.

The wide range in environments and a large variety in types of data provided by partner organisations resulted in a useful critique and development of methodologies used in assessment of sediment quality in maritime regions. It is important to restate that these assessments are the result of a screening procedure to identify and prioritise contaminants and region of concern and that further evaluation of other risk factors, e. g. bioaccumulation, bioavailability and toxicity, are required to determine potential impact.

Recommendations

Sample coverage was inconsistent amongst WHP partners and tended to be focused on perceived point sources and nearshore environments, often leaving large central areas un-surveyed, which prevented a full spatial assessment for 40% of the harbours. Sufficient sample density for regional coverage was estimated at $\sim 0.5\text{-}1.0$ samples/km², while for areas of concern, or high variability, a density of 5-10 samples/km² is recommended.

The suite of analytes also varied within the WHP and no consistent combination of chemicals other than metals (and even these were not consistent) were available for assessment. Ideally, a full set of metallic and organic contaminants would be required to conduct a satisfactory environmental assessment, however evidence is available to show that metals are strongly correlated to other organic pollutants.

Most analyses were undertaken using weak (HCl) to moderately strong acid (aqua regia) mixtures, which are suitable for assessing the adsorbed phases required to assess the magnitude of anthropogenic change and to establish ecological risk. The use of stronger acids, e. g. HNO₃/HClO₄ (nitric plus perchloric acid) would extract inert mineral forms and may make assessment of ecological risk problematic.

A global inadequacy exists in the availability of suitable data for setting pre-anthropogenic sedimentary metal concentrations. Background values should be estimated from local, fine-grained down-hole, core data and will be different for each of the WHP locations. Cores used for background estimation should be recovered from undisturbed areas of deposition and subsurface bioturbation and post-depositional physical and chemical remobilisation should be avoided. Instead, a single suite of background metal concentrations had to be applied to all WHP data, introducing possible error in enrichment assessments of the 10 harbour estuaries.

Some form of normalisation is essential for enrichment assessment to moderate confounding by varying grain size. WHP data frequently lacked data necessary for assessing metal enrichment, i.

e. sediment size, Al and Fe. Size-fractionated metal data are the preferred data for enrichment estimation, however tests conducted in the current work confirm the use of both Al and Fe as normalisation elements and the use of PEN data for determining human-induced change in the absence of elevated carbonate content.

The individual schemes comprising the plethora of indices now available for estimating metals enrichment have not yet been thoroughly tested for validity and some are based on uncertain assumptions. Indices supporting multiple elements, include a classification scheme and based on normalised data and background information are recommended and results from these schemes are closely correlated ($r > 0.95$), i. e. Enrichment Factor (EF), modified Degree on Contamination (*mDC*) and the Mean Enrichment Quotient (MEQ). The original Geo-accumulation Index (*Igeo*) and the modified version (*mIgeo*) exhibits reduced sensitivity and the Nemerow Pollution Index (*PI*) and the modified version (*mPI*) over emphasises elevated metals resulting in loss of discretionary power. Despite the different formulations on which the 10 indices are based, ranking of index results was similar for all assessment tools and as many as possible indices should be tested in assessing anthropogenic change.

Acknowledgements

We are grateful to Alberto Righetti who provided the data on sediments from the Ravenna 'canal' port, collected as part of his doctoral thesis.

REFERENCES

- Abraham, G.M.S., Parker, R.J. 2008. Assessment of heavy metal enrichment factors and the degree of contamination in marine sediments from Tamaki Estuary, Auckland, New Zealand. *Environmental Monitoring and Assessment*, 136, 227-238.
- Abuchacra, P.F.F., Aguiar, V.M.C., Abuchacra, R.C., Baptista Neto, J.Á., Oliveira, A.S. 2015. Assessment of bioavailability and potential toxicity of Cu, Zn and Pb, a case study in Jurujuba Sound, Rio de Janeiro, Brazil. *Marine Pollution Bulletin*, 100, 414-425.
- Aguiar, V.M.C., Abuchacra, P.F.F., Baptista, Neto J.A., Oliveira, A.S. 2018. Environmental assessment concerning trace metals and ecological risks at Guanabara Bay, RJ, Brazil. *Environmental Monitoring Assessment*, 190, 448.
- Aguirre D., Bollard-Breen, B., Cameron, M., Constantine, R., Duffy, C., Dunphy, B., Hart K., Hewitt, J., Jarvis, R., Jeffs, A., R. Kahui-McConnell, Kawharu, M., Liggins, L., Lohrer, A., Middleton, I., Oldman, J., Sewell, M., Smith, A... Wilson, R. 2016. Loved to pieces: Toward the sustainable management of the Waitematā Harbour and Hauraki Gulf, *Regional Studies in Marine Science*, 8/2, 220-233.

1104 Aherns, M., Swales, A., Wadhwa, S., Lewis, M., Hart, C. 2008. Central Waitemata Harbour
 1105 Contaminant Study. Trace Metal Concentrations in Harbour Sediments. Prepared by NIWA Ltd
 1106 for Auckland Regional Council. Auckland Regional Council Technical Report 2008/035

1107 Airoidi, L., Ponti, M., Abbiati, M. 2016. Conservation challenges in human dominated
 1108 seascapes: The harbour and coast of Ravenna. *Regional Studies Marine Science*, 8, 308-318.

1109 Álvarez-Iglesias, P., Quintana, B., Rubio, B., Pérez-Arlucea, M. 2007. Sedimentation rates and
 1110 trace metal input history in intertidal sediments derived from ²¹⁰Pb and ¹³⁷Cs chronology.
 1111 *Journal of Environmental Radioactivity*, 98, 229-250.

1112 Álvarez-Iglesias, P., Rubio, B., Pérez-Arlucea, M. 2006. Reliability of subtidal sediments as
 1113 “geochemical recorders” of pollution input: San Simón Bay (Ría de Vigo, NW Spain). *Estuarine,
 1114 Coastal and Shelf Science*, 70, 507-521.

1115 Balls, P.W., Hull, S., Miler, B.S., Pirie, J.M., Procter, W. 1997. Trace metal Scottish estuarine
 1116 and coastal sediments. *Marine Pollution Bulletin*, 34/1, 42 – 50.

1117 Baptista, Neto J.A., Gingele, F.X., Leipe, T., Brehme, I. 2006. Spatial distribution of heavy
 1118 metals in surficial sediments from Guanabara Bay: Rio de Janeiro, Brazil. *Environmental
 1119 Geology*, 49, 1051-1063.

1120 Baptista, Neto, J.A., Barreto, C.F., Vilela, C.G., Fonseca, E.M., Melo, G.V., Barth, O.M. 2017.
 1121 Environmental change in Guanabara Bay, SE Brazil, based in microfaunal, pollen and
 1122 geochemical proxies in sedimentary cores. *Ocean Coastal Management*, 143, 4-15.

1123 Barton, E.D. Largier, J., Torres, R., Sheridan, M., Trasviña, A., Souza, A.J., Pazos, Y., Valle-
 1124 Levinson, A. 2015. Coastal upwelling and downwelling forcing of circulation in a semi-enclosed
 1125 bay: Ria de Vigo. *Progress in Oceanography*, 134, 173-189.
 1126 <https://doi.org/10.1016/j.pocean.2015.01.014>

1127 Buat-Menard, P. 1979a. Influence de la retombée atmosphérique sur la chimie des métaux en
 1128 trace dans la matière en suspension de l’Atlantique Nord. Thèse Doctorat d’Etat, Paris VII, pp.
 1129 434.

1130 Buat-Menard. P.. Chesselet. R. 1979b. Variable influence of the atmospheric flux on the trace
 1131 metal chemistry of oceanic suspended matter. *Earth Planetary Science Letters*, 42, 399 – 411.

1132 Bedri, Z., Bruen, M., Dowley, A., Masterson, B. 2011. A three-dimensional hydro-
 1133 environmental model of Dublin Bay. *Environmental Modeling and Assessment*, 16(4), 369-384.

1134 Birch, G.F., Robertson, S.E., Taylor, S.E., McConchie, D. 2000. The use of sediments to detect
 1135 human impacts on the fluvial system, *Environmental Geology*, 39, 1015-1028.

1136 Birch, G.F., Taylor, S.E., 2000. Use of sediment quality guidelines in the environmental
 1137 assessment of Port Jackson estuary, Sydney, Australia. AMSA 2000 National Conference,
 1138 University of New South Wales, Sydney, September, 2000.

1139 Birch, G.F., Taylor, S.E., 2000. The use of size-normalisation procedures in the analysis of
1140 organic contaminants in estuarine environments. *Hydrobiologia*, 431, 129-133.

1141 Birch, G.F., Taylor, S.E., 2000. Possible ecological significance of contaminated sediments in
1142 Port Jackson, Sydney. Geological Society of Australia, Abstracts No. 59. 15th Australian
1143 Geological Convention, Sydney, July, 2000.

1144 Birch, G.F., Taylor, S.E. 2002. Possible biological significance of contaminated sediments in
1145 Port Jackson, Sydney, Australia. *Environmental Monitoring and Assessment*, 77, 179-190.

1146 Birch, G. F., 2003. A test of normalisation methods for marine sediments, including a new post-extraction
1147 normalisation (PEN) technique. *Hydrobiologia*, 492, 5-13.

1148 Birch, G.F., Olmos, M. 2008. Sediment-bound heavy metals as indicators of human influence
1149 and biological risk in coastal water bodies. *Journal of Marine Science*, 65, 1407-1413

1150 Birch, G.F., McCready, S., Long, E.R., Taylor, S.E., Spyarakis, G. 2008. Contaminant chemistry
1151 and toxicity of sediments in Sydney Harbour, Australia: spatial extent and chemistry-toxicity
1152 relationships. *Marine Ecology Progress Series*, 363, 71-87.

1153 Birch, G.F., McCready, S. 2009. Catchment sources of heavy metal contamination and influence
1154 on the quality of receiving basin sediments in Port Jackson, Australia. *Science of the Total*
1155 *Environment*, 187,314. DOI: 10.1007/s10661-015-4481-y

1156 Birch, G.F., Gunns, T., Olmos, M. 2016. Sediment-bound metals as indicators of anthropogenic
1157 change in estuarine environments. *Marine Pollution Bulletin*. 101/1, 23-257.
1158 [doi:10.1016/j.marpolbul.2015.09.056](https://doi.org/10.1016/j.marpolbul.2015.09.056).

1159 Birch, G.F. 2016. Determination of sediment metal background concentrations and enrichment in
1160 marine environments – A critical review. *Science of the Total Environment*, 580, 513-531.

1161 Birch, G.F. 2017. Assessment of human-induced change and biological risk posed by
1162 contaminants in estuarine/harbour sediments: Sydney Harbour/estuary (Australia). *Marine*
1163 *Pollution Bulletin*, 116, 234-248. DOI: 10.1016/j.scitotenv.2016.12.028.

1164 Birch, G.F. 2018. A review of chemical-based sediment quality assessment methodologies for
1165 the marine environment. *Marine Pollution Bulletin*, 133, 218-232.
1166 doi.org/10.1016/j.marpolbul.2018.05.039

1167 Birch, G.F. 2020. An assessment of aluminium and iron in normalisation and enrichment
1168 procedures for environmental assessment of marine sediment, *Science of the Total Environment*,
1169 in review.

1170 Blackmore, G. 1998. An overview of trace metal pollution in the coastal waters of Hong Kong.
1171 *The Science of the Total Environment*, 214: 21–48.

1172 Bowen, H.J.M. 1979. *Environmental Chemistry of the Elements*. Academic Press, London.

1173 Brady, J.P., Ayoko, G. A., Martens, W.N., Goonetilleke, A. 2015. Development of a hybrid
 1174 pollution index for heavy metals in marine and estuarine sediments. *Environmental Monitoring*
 1175 *and Assessment*, 187, 306 – 319.

1176 Briciu-Burghina, C., Sullivan, T., Chapman, J., Regan, F. 2014. Continuous high-frequency
 1177 monitoring of estuarine water quality as a decision support tool: a Dublin Port case study.
 1178 *Environmental monitoring and assessment*, 186(9), 5561-5580.

1179 Brooks, P.R., Narin, R., Harris, M., Jeffrey, D., Crowe, T.P. 2016. Dublin Port and Dublin Bay:
 1180 Reconnecting with nature and people. *Regional Studies in Marine Science*.
 1181 <http://dx.doi.org/10.1016/j.rsma.2016.03.007>

1182 Buruaem, L.M., Castro, L.B., Hortellani, M.A., Taniguchi, S., Fillnmann, G., Sasaki, S.T., Petti,
 1183 M.A.V., Sarkis, J.F.S., Bicigeo, M.C., Maranoho, L.A., Davanso, M.B, Nonato, E.F., Cesar, A.,
 1184 Costa-Lotufu, L.V, Abessa, D.M.S., 2013. Integrated quality assessment of sediments from
 1185 harbour areas in Santos-Sao Vicente estuarine systems, Southern Brazil. *Estuarine and Coastal*
 1186 *Shelf Science*, 130, 179-189

1187 Caeiro, S., Cost, M.H., Ramo,s T.B., Ferandes, F., Silveira, N., Coimbra, A., Medeiros, G.,
 1188 Painho, M. 2005. Assessing heavy metal contamination in Sado Estuary sediment: an index
 1189 analysis approach. *Ecological Indicators*, 5, 151 – 169.

1190 Calleja, F., Galván, C., Silió, A. Juanes, J.A., Ondiviela, B. 2017. Long-term analysis of *Zostera*
 1191 *noletii*: a retrospective approach for understanding seagrasses' dynamics. *Marine Environmental*
 1192 *Research*, 130: 93-105. DOI: 10.1016/j.marenvres.2017.07.017

1193 Camargo, M.Z., Sandrini-Net, L., Carreira, R.S., Camargo, M.G. 2017. Effects of hydrocarbon
 1194 pollution in the structure of microbenthic assemblages from two large estuaries in Brazil. *Marine*
 1195 *Pollution Bulletin*, 125, 66-76.

1196 Campos, B.G., Moreira, L.B., Pauly, G.F.E., Cruz, A.C.F., Monte, C.N., Dias da Silva, L.I.,
 1197 Rodrigues, A.P.C., Machado, W., Abessa, D.M.S. 2019. Integrating multiple lines of evidence of
 1198 sediment quality in a tropical bay (Guanabara Bay, Brazil). *Marine Pollution Bulletin*, 146, 925-
 1199 934.

1200 Carreira, R.S., Wagener, A.L, Readman, J.W., Fileman, T.W., Macko, S.A., Veiga, A. 2002.
 1201 Changes in the sedimentary organic carbon pool of a fertilized tropical estuary, Guanabara Bay,
 1202 Brazil: an elemental, isotopic and molecular marker approach. *Mar Chem.* 79, 207–227.

1203 Cevik, F., Goksu, M., Derici, O., Findik, O. 2009. An assessment of metal pollution in surface
 1204 sediments of Seyhan dam by using enrichment factor, geoaccumulation index and statistical
 1205 analysis. *Environmental Monitoring and assessment*, 152/1-4, 309-317.

1206 Chapman, P.M., Wang F. 2001. Assessing sediment contamination in estuaries. *Environmental*
 1207 *Toxicology and Chemistry*, 20/1, 3-22.

1208 Chan, A.K.Y., Xu, W.Z., Liu, X.S., Cheung, S.G., Shin, P.K.S. 2016. Sediment characteristics
1209 and benthic ecological status in contrasting marine environments of subtropical Hong Kong.
1210 *Marine Pollution Bulletin*, 103: 360–370.

1211 Chen, K., Jiao, J.J. 2010. Spatio-temporal trends of heavy metals and source apportionment in
1212 Tolo Harbour, Hong Kong. *Environmental Earth Sciences*, 60, 1439–1445.

1213 Chen, X., Gao, H., Yao, X., Chen, Z., Fang, H., Ye S. 2013. Ecosystem health assessment in the
1214 Pearl River Estuary of China by considering ecosystem coordination. *Plos one*, 8, e70547.

1215 Cordeiro, R.C., Machado, W., Santelli, R.E., Figueiredo, Jr. A.G., Seoane, F.C.S., Oliveira, E.P.,
1216 Freire, A.S., Bidone, E.D., Monteiro, F.F., Silva, F.T., Meniconi, M.F.G. 2015. Geochemical
1217 fractionation of metals and semimetals in surface sediments from tropical impacted estuary
1218 (Guanabara Bay, Brazil). *Environmental Earth Science*, 74, 1363–1378.

1219 Costanza, R., d'Arge, R., deGroot, R., Farber, S., Grasso, M., Hannon, B., Limburg, K., Naeem,
1220 S., Oneill, R.V., Paruelo, J., Raskin, R.G., Sutton, P. vandenBelt, M. 1997. The value of the
1221 world's ecosystem services and natural capital. *Nature*, 387 (6630), 253-260.

1222 Costanza, R., de Groot, R., Sutton, P., van der Ploeg, S., Anderson, S.J., Kubiszewski, I., Farber,
1223 S., Turner R.K. 2014. Changes in the global value of ecosystem services. *Global Environmental*
1224 *Change*, 26, 152-158.

1225 Coughanowr, C., Whitehead, S., Whitehead, J., Einoder, L., Taylor, U., Weeding, B. 2015. State
1226 of the Derwent Estuary 2015: a review of environmental data from 2009 to 2014. Derwent River
1227 Program. DPIPW, Tasmania

1228 Covelli, S., Protopsalti, I., Acquavita, A., Sperle, M., Bonardi, M., Emili, A. 2012. Spatial
1229 variation, speciation and sedimentary records of mercury in the Guanabara Bay(Rio de Janeiro,
1230 Brazil). *Continental Shelf Research*, 35, 29-42.

1231 Cunningham, A. 2018. Distribution and sources of PAHs and trace metals in Bull Island, Dublin
1232 Bay (Doctoral dissertation, Dublin City University).

1233 Dalrymple, R.W., Zaitlin, B.A., Boyd, R., 1992. Estuarine facies models, conceptual basis and
1234 stratigraphic implications. *Journal of Sedimentary Petrology*, 62, 1130-1146.

1235 Davoren, M., Shúilleabháin, S.N., O'Halloran, J., Hartl, M.G.J., Sheehan, D., O'Brien, N.M.,
1236 Van Pelt, F.N.A.M., Mothersill, C. 2005. A test battery approach for the ecotoxicological
1237 evaluation of estuarine sediments. *Ecotoxicology*, 14(7), 741-755.

1238 Di Toro, D.M., Mahony, J.D., Hansen, D.J., Scott, K.J., Hicks, M.B., Mayr, S.M., Redmond,
1239 M.S. 1990. Toxicity of cadmium in sediments: the role of acid volatile sulfide. *Environmental*
1240 *Toxicology and Chemistry*, 9, 1487-1502.

1241 Di Toro, D.M., Mahony, J.D., Hansen, D., Scott, K.J., Carlson, A.R., Ankley, G.T. 1992. Acid
 1242 volatile sulfide predicts the acute toxicity of cadmium and nickel in sediments. *Environmental*
 1243 *Science and Technology*, 26, 96-101.

1244 DPC. 2014. Alexandra Basin Redevelopment Project. Environmental Impact Statement. Dublin
 1245 Port Company, Dublin.

1246 Echavarri, B., Juanes, J.A., García-Castrillo, G., Revilla, J.A. 2007. Medium-term responses of
 1247 rocky bottoms to sewage discharges through a deepwater outfall in the NE Atlantic. *Marine*
 1248 *Pollution Bulletin*, 54, 941-954.

1249 Edgar, G.J., Barrett, N.S., Graddon, D.J., Last, P.R. 2000. The conservation significance of
 1250 estuaries: a classification of Tasmanian estuaries using ecological, physical and demographic
 1251 attributes as a case study. *Biological Conservation*, 92/3, 383-397.

1252 EPA. 2006, Environmental Protection Agency. Water Framework Directive monitoring
 1253 programme. EPA Ireland:
 1254 [http://www.epa.ie/pubs/reports/water/other/wfd/EPA_water_WFD_monitoring_programme_mai](http://www.epa.ie/pubs/reports/water/other/wfd/EPA_water_WFD_monitoring_programme_main_report.pdf)
 1255 [n_report.pdf](http://www.epa.ie/pubs/reports/water/other/wfd/EPA_water_WFD_monitoring_programme_main_report.pdf)

1256 EPA. 2015. Environmental Protection Agency. Water quality in Ireland 2010- 2012, EPA
 1257 Ireland: <https://www.epa.ie/pubs/reports/water/waterqua/wqr20102012/WaterQualityReport.pdf>

1258 EPD. 2017. Environmental Protection Department. Marine water quality in Hong Kong in 2016.
 1259 Retrieved June 19, 2018 from The Government of Hong Kong Special Administrative Region,
 1260 Environmental Protection Department Web site: [http://wqrc.epd.gov.hk/pdf/water-](http://wqrc.epd.gov.hk/pdf/water-quality/annual-report/MarineReport2016eng.pdf)
 1261 [quality/annual-report/MarineReport2016eng.pdf](http://wqrc.epd.gov.hk/pdf/water-quality/annual-report/MarineReport2016eng.pdf)

1262 Fabbri, D., Felisatti, O., Lombardo, M., Trombini, C., Vassura, I. 1998. The Lagoon of Ravenna
 1263 (Italy): characterisation of mercury-contaminated sediments. *Science of the Total Environment*,
 1264 213, 121-128.

1265 Fabbri, D., Lombardo, M., Trombini, C., Vassura, I., Zavoli, E., Horvat, M., 2001. Mercury
 1266 contamination of a coastal lagoon (Pialassa Baiona, Ravenna, Italy). *RMZ – Materials and*
 1267 *Geoenvironment*, 48, 186-192.

1268 Fabbri, D., Tartari, D., Trombini, C. 2000. Analysis of poly(vinyl chloride) and other polymers
 1269 in sediments and suspended matter of a coastal lagoon by pyrolysis-gas chromatography-mass
 1270 spectrometry. *Analytica Chimica Acta*, 413, 3-11.

1271 Fernández, E., Álvarez-Salgado, X.A., Beiras, R., Ovejero, A., Méndez, G. 2016. Coexistence of
 1272 urban uses and shellfish production in an upwelling-driven, highly productive marine
 1273 environment: The case of the Ría de Vigo (Galicia, Spain). *Regional Studies in Marine Science*,
 1274 8, 362-370.

1275 Figueiredo, Jr. A.G., Toledo, M.B., Cordeiro, R.C., Godoy, J.M.O., Silva, F.T., Vasconcelos,
 1276 S.C., Santos, R.A. 2014. Linked variations in sediment accumulation rates and sea-level in

1277 Guanabara Bay, Brazil, over the last 6000 years. *Palaeogeography Palaeoclimatology*
1278 *Palaeoecology*, 415, 83–90.

1279 Fletcher, T., Duncan, H., Poelsma, P., Lloyd, S. 2004. Stormwater flow and quality and the
1280 effectiveness of non-propriety stormwater treatment measures: A review and gap analysis,
1281 Technical Report 04/8. December, 2004. Co-operative Research Centre for Catchment
1282 Hydrology, Melbourne, Australia.

1283 Ghani, S.A., El Zokm, G., Shobier, A., Othman, T., Shreadah, M. 2013. Metal pollution to
1284 surface sediments of Abu-Qir Bay and Eastern Harbour of Alexandria, Egypt. *Egypt Journal of*
1285 *Aquatic Research*, 39, 1-12.

1286 Giltrap, M., Macken, A., Davoren, M., Minchin, D., McGovern, E., Foley, B., McHugh, B. 2009.
1287 Use of caged *Nucella lapillus* and *Crassostrea Gigas* to monitor tributyltin-induced bioeffects in
1288 Irish coastal waters. *Environmental Toxicology and Chemistry*, 28 (8), 1671–1678.

1289 Giltrap, M., McHugh, B., Ronan, J., Wilson, J., McGovern, E., 2014. Biological Effects and
1290 Chemical Measurements in Irish Marine Waters. Marine Institute.

1291 Gómez, A.G., Juanes, J.A., Ondiviela, B., Revilla, J.A. 2014. Assessment of susceptibility to
1292 pollution in littoral waters using the concept of recovery time. *Marine Pollution Bulletin*, 81:
1293 140-148. DOI: 10.1016/j.marpolbul.2014.02.004.

1294 Grant, A., Middleton, R. 1990. An assessment of metal contamination of sediments in the
1295 Humber Estuary, U.K. *Estuarine, Coastal and Shelf Science*, 31, 71-85.

1296 Green, M., Timperley, M., Collins, R., Senior, A., Adams, R., Swales, A. 2004. Prediction of
1297 contamination accumulation in the Waitemata Harbour - Results-Methods. NIWA Client Report:
1298 HAM2003-087/1. Report prepared by NIWA for Auckland Council.

1299 Guerra. 2012. Polycyclic aromatic hydrocarbons, polychlorinated biphenyls and trace metals in
1300 sediments from a coastal lagoon (northern Adriatic, Italy). *Water Air and Soil Pollution* 223(1),
1301 85-98.

1302 Guerra, R., Pasteris, A., Ponti, M. 2009. Impacts of maintenance channel dredging in a northern
1303 Adriatic coastal lagoon. I: Effects on sediment properties, contamination and toxicity. *Estuarine*
1304 *Coastal and Shelf Science*, 85, 134-142.

1305 Guerra, R., Pasteris, A., Ponti, M., Fabbri, D., Bruzzi, L. 2007. Impact of dredging in a shallow
1306 coastal lagoon: Microtox (R) Basic Solid-Phase Test, trace metals and *Corophium* bioassay.
1307 *Environmental International*, 33, 469-473.

1308 Guerra, R., Pasteris, A., Seok-hyung, L., No-jin, P., Ok, G., 2014. Spatial patterns of metals,
1309 PCDDs/Fs, PCBs, PBDEs and chemical status of sediments from a coastal lagoon (Pialassa
1310 Baiona, NW Adriatic, Italy). *Marine Pollution Bulletin*, 89, 407-416.

1311 Guo, W., Liu, X., Liu, Z., Li, G. 2010. Pollution and potential ecological risk evaluation of heavy
1312 metals in the sediments around Dongjiang Harbour. *Procedia Environmental Sciences*, 2, 729 –
1313 736.

1314 Hakanson, L. 1980. An ecological risk index for aquatic pollution control: A sedimentological
1315 Approach. *Marine Research*, 14, 975-1001.

1316 Hogg, I.D., Norris, R.H. 1991. Effects of runoff from land clearing and urban development on
1317 the distribution and abundance of macro-invertebrates in pool areas of a river. *Australian Journal*
1318 *of Marine and Freshwater Science*, 42, 507-518.

1319 ISO. 1995. ISO Standard 11466 Soil quality - Extraction of trace elements soluble in aqua regia.

1320 Kabir, M.I., Lee, H., Kim, G. Jun, T. 2011. Correlation assessment and monitoring of the
1321 potential pollutants in the surface sediments of Pyeongchang River, Korea. *International Journal*
1322 *of Sedimentary Research*, 26/2, 152-162.

1323 Karbassi, A.R., Monavari, S.M., Nabi Bidhendi, G.R., Nouri, J., Nematpour, K. 2008. Metal
1324 pollution assessment of sediment and water in the Shur River. *Environmental Monitoring and*
1325 *Assessment*, 147, 107 – 116.

1326 Katz, A., Kaplan I.R. 1981. Heavy metals behaviour in coastal sediments of Southern California:
1327 A critical review and synthesis. *Marine Chemistry*, 10, 261 – 299.

1328 Kaushik, A., Kansal, A., Kumari, S., Kaushik, C. 2009. Heavy metal contamination of River
1329 Yamuna, Haryana, India: assessment by metal enrichment factor of the sediments. *Journal of*
1330 *Hazardous Materials*, 164/1, 265-270.

1331 Kemp, A.L.W., Thomas, R.L., Del, C.J. Jaquet, J.H. 1976. Cultural impact on the geochemistry
1332 of sediments in Lake Erie. *Journal of Fish Research, Board of Canada*, 33, 440 – 462.

1333 Kim, B., F., Angeli, J., Ferreira, P., Sartoretto, J., Miyoshi, C., de Mahiques, M., Figueira, R.
1334 2017. Use of a chemometric tool to establish the regional background and assess trace metal
1335 enrichment at Baixada Santista – southeastern Brazil. *Chemosphere*, 166, 372-379.

1336 Kim, B., F., Angeli, J., Ferreira, P., de Mahiques, M., Figueira, R. 2018. Critical evaluation of
1337 different methods to calculate the Geoaccumulation Index for environmental studies: A new
1338 approach for Baixada Santista – Southeastern Brazil. *Marine Pollution Bulletin*, 127, 548-552.

1339 Kjerfve, B., Ribeiro, C.H., Dias, G.T.M., Fillipo, A.M., Quaresma, V.S. 1997. Oceanographic
1340 characteristics of an impacted coastal bay: Baía de Guanabara, Rio de Janeiro, Brazil.
1341 *Continental Shelf Research*, 17: 1609-1643.

1342 Kjerfve, B., Lacerda, L.D., Dias, G.T.M. 2001. Baía de Guanbara, Rio de Janeiro, Brazil. In:
1343 Seeliger U, Kjerfve B (Eds.). *Coastal marine ecosystems of Latin America*. Berlin, Springer-
1344 Verlag.

1345 Lai, R.W.S., Perkins, M.J., Ho, K.K.Y., Astudillo, J.C., Yung, M.M.N., Russell, B.D., Williams,
1346 G.A., Leung, K.M.Y. 2016. Hong Kong's marine environments: History, challenges and
1347 opportunities. *Regional Studies in Marine Science*, 8, 259–271.

1348 Lee, S.B., Birch, G.F., Lemckert, C. 2011. Field and modelling investigations of fresh-water
1349 plume behaviour in response to infrequent high-precipitation events, Sydney Estuary, Australia.
1350 *Journal of Estuarine and Coastal Shelf Science*, 92, 380-402.

1351 Liu, L., Wang, Z., Ju, F., Zhang, T. 2015. Co-occurrence correlations of heavy metals in
1352 sediments revealed using network analysis. *Chemosphere*, 119: 1305–1313.

1353 Long, E.R., L.G. Morgan. 1990. The Potential for Biological Effects of Sediment-
1354 Sorbed Contaminants Tested in the National Status and Trends program. NOAA
1355 Technical Memorandum NOS OMA 52, Seattle, WA 175 pp & appendices.

1356 Long, E.R., MacDonald, D.D., Smith, S.L., Calder, E.D. 1995. Incidence of adverse biological
1357 effects within ranges of chemical concentrations in marine and estuarine sediments. *Environment*
1358 *Management*, 19, 81-97.

1359 Long, E.D. MacDonald, D.D. 1998. Recommended uses of empirically derived, sediment quality
1360 guidelines for marine and estuarine ecosystems. *Human and Ecological Risk Assessment*, 4, 1019-
1361 1093.

1362 López, I., Álvarez, C., Gil, J.L., García, A., Bárcena, J.F., Revilla, J.A. 2013. A method for the
1363 source apportionment in bathing waters through the modelling of wastewater discharges:
1364 Development of an indicator and application to an urban beach in Santander (Northern Spain).
1365 *Ecological Indicators*. 24, 334-343. DOI: 10.1016/j.ecolind.2012.07.003.

1366 Loring, D.H. 1991. Normalisation of heavy-metal data from estuarine and coastal sediments.
1367 *ICES Journal of Marine Science*, 48, 101 – 115

1368 MacDonald, D.D., Di Pinto, L.M., Field, J., Ingersoll, C.G., Long, E.R., Swartz, R.C. 2000.
1369 Development and evaluation of consensus-based sediment effect concentrations for
1370 polychlorinated biphenyls (PCBs). *Environmental Toxicology and Chemistry*, 19, 1403 – 1413.

1371 Macken, A., Giltrap, M., Foley, B., McGovern, E., McHugh, B., Davoren, M. 2008. An
1372 integrated approach to the toxicity assessment of Irish marine sediments: validation of
1373 established marine bioassays for the monitoring of Irish marine sediments. *Environment*
1374 *International*, 34(7), 1023-1032.

1375 Mao, Q., Shi, P., Yin, K., Gan, J., Qi, Y. 2004. Tides and tidal currents in the Pearl River
1376 Estuary. *Continental Shelf Research*, 24, 1797–1808.

1377 Maranhão, L.A., Abreu, I.M., Santelli, R.E., Cordeiro, R.C., Soares-Gomes, A., Moreira, L.B.,
1378 Morais, R.D., Abessa, D.M. 2009. Sediment toxicity assessment of Guanabara Bay, Rio de
1379 Janeiro, Brazil. *Journal of Coastal Research*, 56, 851-855.

1380 Maranhão, L.A., Abreu, I.M., Santelli, R.E., Cordeiro, R.C., Soares-Gomes, A., Moreira, L.B.,
1381 Morais, R.D., Abessa, D.M. 2010. Acute and chronic toxicity of sediment samples from
1382 Guanabara Bay (RJ) during the rainy period. *Brazilian Journal of Oceanography*, 58, 77-85.

- 1383 McCready, S., Spyrakis, G., Greely, C.R., Birch, G.F., Long, E.L. 2004. Toxicity of surficial
1384 sediments from Sydney Harbour and vicinity, Australia. *Environmental Monitoring and*
1385 *Assessment*, 96, 53-83.
- 1386 McCready, S., Birch, G.F., Long, E.L., Spyrakis, G., Greely, C.R. 2006. Predictive abilities of
1387 numerical sediment quality guidelines for Sydney Harbour, Australia and vicinity. *Environment*
1388 *International*, 32, 38-649.
- 1389 McRae, C., Snape, C.E., Sun, C-G., Fabbri, D., Tartari, D., Trombini, C., Fallick, A.E. 2000. Use
1390 of compound-specific stable isotope analysis to source anthropogenic natural gas-derived
1391 polycyclic aromatic hydrocarbons in a lagoon sediment. *Environmental Science and Technology*,
1392 34, 4684-4686.
- 1393 Mills, G., Williamson, B., Cameron, M., Vaughan, M. 2012, Marine sediment contaminants:
1394 Status and trends assessment 1998 to 2010. Prepared by Diffuse Sources Ltd for Auckland
1395 Council. Auckland council technical report TR2012/041.
- 1396 Mills, G., Williamson, B. 2014. Marine sediment contaminant monitoring programme: review of
1397 data quality and procedures. Prepared by Diffuse Sources Ltd for Auckland Council. Auckland
1398 Council technical report, TR2014/041
- 1399 Moraes, L.A.F. 2012. Remnant Vegetation Analysis of Guanabara Bay Basin, Rio de Janeiro,
1400 Brazil, Using Geographical Information System. In: Moutinho P (Ed.). *Deforestation Around the*
1401 *World*. InTech, Available from: [http://www.intechopen.com/books/deforestation-](http://www.intechopen.com/books/deforestation-aroundtheworld/remnant-vegetation-analysis-of-guanabara-bay-basin-rio-de-janeiro-brazil-using-geographicalinformat)
1402 [aroundtheworld/remnant-vegetation-analysis-of-guanabara-bay-basin-rio-de-janeiro-brazil-](http://www.intechopen.com/books/deforestation-aroundtheworld/remnant-vegetation-analysis-of-guanabara-bay-basin-rio-de-janeiro-brazil-using-geographicalinformat)
1403 [using-geographicalinformat](http://www.intechopen.com/books/deforestation-aroundtheworld/remnant-vegetation-analysis-of-guanabara-bay-basin-rio-de-janeiro-brazil-using-geographicalinformat).
- 1404 Moraes, R.B.C., Pfeiffer, W.C., Guimarães, J.R.D., Borges, A.L.N., Campos, A.N. 2000.
1405 Development of sediment toxicity test with tropical penaeid shrimps. *Environmental Toxicology*
1406 *and Chemistry* 19, 1881-1884, 2000.
- 1407 Morton, B. 1989. Pollution of the coastal waters of Hong Kong. *Marine Pollution Bulletin*, 20,
1408 310–318.
- 1409 Morton, B. 1996. Protecting Hong Kong's marine biodiversity: present proposals, future
1410 challenges. *Environmental Conservation*, 23, 55–65.
- 1411 Müller, G. 1969. Index of geoaccumulation in sediments of the Rhine. *Geojournal*, 2, 108-118.
- 1412 Müller, G. 1979. Schwermetalle in den sedimenten des Rheins-Veränderungen seitt 1971.
1413 *Umschau*, 79, 778-783.
- 1414 Müller, G. 1986. Schadstoffe in sedimenten - Sedimenteals Schadstoffe. *Mitt. Osterr. Geol Ges.*,
1415 79, 107-126.
- 1416 Munksgaard, N.C., Kaestli, M., Gibb, K., Dostine, P., Townsend, S. 2012. Darwin Harbour
1417 Baseline Sediment Survey 2012. Department of Land Resource Management, Darwin, NT.

1418 Munksgaard, N.C., Fortune, J. 2015. Metal concentrations in Rapid Creek sediment cores.
 1419 Environmental Chemistry and Microbiology Unit, Research Institute for the Environment and
 1420 Livelihoods Charles Darwin University, Darwin, NT.

1421 Munksgaard, N.C., Kaestli, M., Gibb K., Dostine, P., Townsend, S. 2013. Darwin Harbour
 1422 baseline Sediment survey, 2012. Northern Territory Dept., of Land and Resources Management.
 1423 Darwin, NT.

1424 Munksgaard, N.C., Hutley, L.B., Metcalfe, K.N., Padovan, A.C., Palmer, C., Gibb, K.S. 2018.
 1425 Environmental challenges in a near-pristine mangrove estuary facing rapid urban and industrial
 1426 development: Darwin Harbour, Northern Australia. *Regional Studies in Marine Science* 25,
 1427 100438, <https://doi.org/10.1016/j.rsma.2018.11.001>.

1428 Murphy, B.T., O'Reilly, S.S., Monteys, X., Reid, B.F., Szpak, M.T., McCaul, M.V., Jordan, S.F.,
 1429 Allen, C.C., Kelleher, B.P. 2016. The occurrence of PAHs and faecal sterols in Dublin Bay and
 1430 their influence on sedimentary microbial communities. *Marine Pollution Bulletin*, 106(1-2), 215-
 1431 224.

1432 Nemerow, N.L. 1991. Stream, lake, estuary and ocean pollution. New York, US.

1433 NLWRA. 2002. Australian Catchment, River and Estuary Assessment. 2002. 1. National Land
 1434 and Water Resources Audit. Federal Government, Canberra, Australia.

1435 NURP. 1983. US Environmental Protection Agency. Results of the National Runoff Program.
 1436 NTIS PB84-1855552, Washington, DC.

1437 Olmos, M.A., Birch, G.F. 2010. A novel method using sedimentary metals and GIS for
 1438 measuring anthropogenic change in coastal lake environments, *Environmental Science and*
 1439 *Pollution Research*, 17/2, 270-287.

1440 Ondiviela, B., Gomez, A.G., Puente, A, Juanes, J.A. 2013. A pragmatic approach to define the
 1441 ecological potential of water bodies heavily modified by the presence of ports. *Environmental*
 1442 *Science and Policy*, 33: 320-331. DOI: [10.1016/j.envsci.2013.07.001](https://doi.org/10.1016/j.envsci.2013.07.001)

1443 OSPAR. 2005. Agreement on background concentrations for contaminants in seawater, biota and
 1444 sediments. OSPAR Agreement 2005-6. Publication number 2005-6.

1445 OSPAR. 2008. Co-ordinated environmental monitoring programme assessment manual for
 1446 contaminants in sediments and biota. OSPAR Commission, 2008. Monitoring and Assessment
 1447 Series. CEMP Assessment Manual. Publication number 379/2008.

1448 Pang, H.J., Lou, Z.H., Jin, A.M., Yan, K.K., Jiang, Y., Yang, X.H., Chen, C.T.A., Chen, X.G.,
 1449 2017. Contamination, distribution and sources of heavy metals in the sediments of Andong tidal
 1450 flat, Hangzhou Bay, China. *Continental Shelf Research*, 110, 72-84.

1451 Pearson, T.H., Rosenberg, R. 1978. Macrobenthic succession in relation to organic enrichment
 1452 and pollution of the marine environment. *Oceanography and Marine Biology Annual Review*,
 1453 16, 229-311.

1454 Pekey, H. 2006. Heavy metal pollution assessment in sediments of the Izmit Bay, Turkey.
 1455 Environmental Monitoring and Assessment, 123, 219 – 231.

1456 Pignotti, E., Guerra, R., Covelli, S., Fabbri, E., Dinelli, E. 2018. Sediment quality assessment in
 1457 a coastal lagoon (Ravenna, NE Italy) based on SEM-AVS and sequential extraction procedure.
 1458 Science of the Total Environment, 635, 216-227.

1459 Ponti, M., Vadrucchi, M.R., Orfanidis, S., Pinna, M. 2009. Biotic indices for ecological status of
 1460 transitional water ecosystems. Transitional Waters Bulletin, 3, 32-90
 1461 <https://doi.org/10.1285/i1825229Xv3n3p32>

1462 Ponti, M., Casselli, C., Abbiati, M. 2011. Anthropogenic disturbance and spatial heterogeneity of
 1463 macrobenthic invertebrate assemblages in coastal lagoons: the study case of Pialassa Baiona
 1464 (northern Adriatic Sea). Helgology Marine Research, 65, 25-42.

1465 Ponti, M., Pasteris, A., Guerra, R., Abbiati, M. 2009. Impacts of maintenance channel dredging
 1466 in a northern Adriatic coastal lagoon. II: Effects on macrobenthic assemblages in channels and
 1467 ponds. Estuary Coastal and Shelf Science, 85, 143-150.

1468 Prego, R., Cobelo-Garcia, A. 2003. Twentieth century overview of heavy metals in the Galician
 1469 Rias (NW Iberian Peninsula). Environmental Pollution, 121, 425 - 452.

1470 Puente, A., Juanes, J.A., García-Castrillo, G., Álvarez, C., Revilla, J.A., Gil, J.L. 2002. Baseline
 1471 study of soft bottom benthic assemblages in the bay of Santander (Gulf of Biscay).
 1472 Hydrobiologia, 475/476, 141-149.

1473 Quelle, C., Besada, V., Andrade, J.M., Gutiérrez, N., Schultze, F., Gago, J., González, J.J. 2011.
 1474 Chemometric tools to evaluate the spatial distribution of trace metals in surface sediments of two
 1475 Spanish rias. Talanta, 87, 197–209.

1476 Riba, I., Del, Valls, T.A., Forja, J.M., Gomez-Parra, A. 2002. Influence of the Aznalcollar
 1477 mining spill on the vertical distribution of heavy metals in sediments from the Guadalquivir
 1478 estuary (SW Spain). Marine Pollution Bulletin, 44, 39 – 47.

1479

1480 Rebello, A., Haekel, W., Moreira, I., Santelli, R., Schroeder, F. 1986. The fate of heavy metals in
 1481 an estuarine tropical system. Marine Chemistry, 18, 215–225.

1482 Remoundou, K., Diaz-Simal, P., Koundouri, P., Rulleau, B. 2015. Valuing climate change
 1483 mitigation: A choice experiment on a coastal and marine ecosystem. Ecosystem Services, 11, 87-
 1484 94. DOI: 10.1016/j.ecoser.2014.11.003.

1485 Rees, H.L., Hyland, J.L., Hyland, K., Mercer, C.S.L., Roff, J.C., Ware, S. 2008. Environmental
 1486 indicators: utility in meeting regulatory needs. An overview. ICES Journal of Marine science
 1487 advanced access publication. September 22, 2008. Indicators.

1488 Rodriguez, J.G., Tueros, I., Borja, A., Belzunce, M.J., Franco, J., Solaun, O., Valencia, V.,
1489 Zuazo, A. 2006. Maximum likelihood mixture estimation to determine metal background values
1490 in estuarine and coastal sediments with in the European Water Framework Directive. *Science of*
1491 *the Total Environment*, 370, 278 – 293.

1492 Roy, R.S., Williams, R.J., Jones, A.R. Yassini, I., Gibbs, P.J., Coates, B. West, R.J. Scanes, P.R.
1493 Hudson, J.P., Nichol, S. 2001. Structure and Function of South-east Australian Estuaries.
1494 *Estuarine, Coastal and Shelf Science*, 53, 351–384.

1495 Rubio, B., Nombela, M.A., Vilas, F. 2000. Geochemistry of major and trace elements in
1496 sediments of the Ría de Vigo (NW Spain): An assessment of metal pollution. *Marine Pollution*
1497 *Bulletin*, 40, 968-980.

1498 Siaka, M., Owens, C.M., Birch G.F. 1998. Evaluation of some digestion methods for the
1499 determination of heavy metals in sediments samples by flame AAS. *Analytical Letters*, 3 1/4,
1500 703-718.

1501 Silva, S.F.G., Brüning, I.M.R.A., Montone, R.C., Taniguchi, S., Cascaes, M.J., Dias, P.S.,
1502 Lavandier, R.C., Hauser-Davis, R.A., Moreira, I. 2013. Polybrominated diphenyl ethers
1503 (PBDES) and polychlorinated biphenyls (PCBS) in mussels and two fish species from the
1504 estuary of the Guanabara Bay, Southeastern Brazil. *Bulletin of Environmental Contamination*
1505 *and Toxicology*, 91, 261–266.

1506 Simpson, S.L., Batley, G.E., Maher, W.A. 2016. Chemistry of sediment contaminants. In:
1507 *Sediment Quality Assessment*. A practical guide. Chap 3. Eds. Simpson, S. L., Batley, G. E.,
1508 CSIRO, Bangor, NSW.

1509 Singh, K.P., Mohan, D., Singh, V.K., Malik, A. 2005. Studies on distribution and fractionation of
1510 heavy metals in Gomti River sediments – a tributary of the Ganges, India. *Journal of Hydrology*,
1511 312, 14 – 27.

1512 Steinberg, P.D., Airoidi, L., Banks, J., Leung, K.M.Y. 2016. Introduction to the special issue on
1513 the World Harbour Project. *Regional Studies in Marine Science*, 8 (2), 217-219.
1514 doi:10.1016/j.rsma.2016.10.001

1515 SMO, 2018. Survey and Mapping Office, Hong Kong geographic data. Retrieved June 19, 2018
1516 from The Government of Hong Kong Special Administrative Region, Survey and Mapping
1517 Office Web site:
1518 https://www.landsd.gov.hk/mapping/en/publications/hk_geographic_data_sheet.pdf

1519 Szefer, P., Skwarzec, B. 1988. Distribution and possible sources of some elements in the
1520 sediment cores of the southern Baltic. *Marine Chemistry*, 23, 109 – 129.

1521 Tang, C.W., Ip, C.C., Zhang, G., Shin, P.K.S., Qian, P., Li, X. 2008. The spatial and temporal
1522 distribution of heavy metals in sediments of Victoria Harbour, Hong Kong. *Marine Pollution*
1523 *Bulletin*, 57, 816–825.

1524 Tanner, P.A., James, J.W.C., Chan, K., Leung, L.S. 1993. Variations in trace metal and total
 1525 organic carbon concentrations in marine sediments from Hong Kong. *Environmental*
 1526 *Technology*, 14, 501–516.

1527 Taylor, S.R. 1964. Abundance of chemical elements in the continental crust: a new table.
 1528 *Geochimica Cosmochimica Acta*, 28, 1273-1285.

1529 Thuong, N.T., Yoneda, M., Ikegami, M., Takakura, M. 2013. Source discrimination of heavy
 1530 metals in sediment and water of the Lich River in Hanoi city using multivariate statistical
 1531 approaches. *Environmental Monitoring and Assessment*, 185/10, 8065 – 8075.

1532 Turekian, K.K. Wedepohl, D.H. 1961. Distribution of the elements major units of the Earth's
 1533 crust. *Bulletin of the Geological Society of America*, 72, 175 – 192.

1534 USEPA. 1983. Results of the National Runoff Program. 1. Final Report NTIS PB84-185552. US
 1535 EPA, Washington, DC.

1536 USEPA. 1994. USEPA 6020 Inductively Coupled Plasma - Mass Spectrometry. United States
 1537 Environmental Protection Agency.

1538 USEPA. 2007a. Method 3051A. Microwave assisted acid digestion of sediments, sludges, soils,
 1539 and oils. United States Environmental Protection Agency, 30pp.

1540 US EPA. 2007b. Method 6020A: Inductively Coupled Plasma - Mass Spectrometry, 30pp.

1541 Usero, J., González-Regalado, E., Gracia, I. 1996. Trace metals in the bivalve mollusc *Chamelea*
 1542 *gallina* from the Atlantic coast of southern Spain. *Marine Pollution Bulletin*, 32, 305-310
 1543 [https://doi.org/https://doi.org/10.1016/0025-326X\(95\)00209-6](https://doi.org/https://doi.org/10.1016/0025-326X(95)00209-6)

1544 Vigurí, J., Verde, J., Irabien, A. 2002. Environmental assessment of polycyclic aromatic
 1545 hydrocarbons (PAHs) in surface sediments of the Santander Bay, Northern Spain. *Chemosphere*,
 1546 48(2): 157-165. DOI: 10.1016/S0045-6535(02)00105-4.

1547 Vigurí, J.R., Irabien, M.J., Yusta, I., Soto, J., Gómez, J., Rodriguez, P., Martinez-Madrid, M.,
 1548 Irabien, J.A., Coz, A. 2007. Physico-chemical and toxicological characterization of the historic
 1549 estuarine sediments: A multidisciplinary approach. *Environment International*, 33 (4), 436-444.
 1550 DOI: 10.1016/j.envint.2006.10.005.

1551 Ying, W., Batley, G.E., Ahsanullah, M. 1982. The ability of sediment extractions to measure the
 1552 bioavailability of metals to three marine invertebrates. *Science of the Total Environment*, 125, 6-
 1553 84.

1554 Wedepohl, K.H. 1995. The composition of the continental crust. – *Geochim and Cosmochim*
 1555 *Acta*, 59, 7, 1217—1232.

1556 Whalley, C., Rowlatt, S., Bennet, M., Lovell, D. 1999. Total arsenic in sediments from the
 1557 western North Sea and the Humber estuary. *Marine Pollution Bulletin*, 38, 394-400.

1558 Whitehead, J., Coughanowr, C., Agius, J., Chrispijn, J., Taylor, U., Wells, F. 2010. State of the
 1559 Derwent Estuary 2009: a review of pollutant sources, loads and environmental quality data from
 1560 2003-2009. Derwent River Program. DPIPW, Tasmania.

1561 Wilson, J.G., Jeffrey, D.W. 1987. Europe-wide industry for monitoring marine quality. In:
 1562 Kramer, K. J. M. (Ed.), Biological indicators of pollution. Royal Irish Academy, Dublin, Ireland.
 1563 225 – 242pp

1564 Xavier de Brito, A.P., Andrade, I.M.R., Brüning, I.M. 2002. Chlorinated pesticides in mussels
 1565 from Guanabara Bay, Rio de Janeiro, Brazil. Marine Pollution Bulletin, 44, 71-81.

1566 Xu, G., Liu J., Pei, S., Kong, X., Hu, G. 2014. Distribution and source of heavy metals in the
 1567 surface sediments from the near-shore area, north Jiangsu Province, China. Marine Pollution
 1568 Bulletin, 83, 275-281.

1569 Zhang, W., Feng, H., Chang, J., Qu, J., Xie, H., Yu, L. 2009. Heavy metal contamination in
 1570 surface sediments of the Yangtze River intertidal zone: an assessment from different indices.
 1571 Environmental Pollution, 157/5, 1533 – 1543.

1572 Zhou, F., Guo, H., Hao, Z. 2007a. Spatial distribution of heavy metals in Hong Kong's marine
 1573 sediments and their human impacts: A GIS-based chemometric approach. Marine Pollution
 1574 Bulletin, 54, 1372–1384.

1575 Zhou, F., Guo, H., Liu, L. 2007b. Quantitative identification and source apportionment of
 1576 anthropogenic heavy metals in marine sediment of Hong Kong. Environmental Geology, 53,
 1577 295–305.

1578 Zoller, W.H., Gladney, E.S., Gordon, G.E. Bors, J.J. 1974. Emissions of trace elements from
 1579 coal-fired power plants: In Hemphill, D. D. Eds., Trace substances in environmental health, V. 8,
 1580 Rolla, University of Missouri.

1581
 1582

Table 1. Data availability, sampling device used and analytical method

Location	Years available	Years used	Sampling device	Analytical method	Extraction technique	Acid	RM	Available data			Comments
								Al	Fe	Size	
Auckland	2010,11,12,13	2008	na	ICP-MS	HNO ₃ /HCl	M	?Y	N	N	N	PEN data SN
Darwin	2012	2012	corer	ICP-MS	HNO ₃ /HClO ₄	S	Y	Y	Y	Y	
Derwent	1998,99,2001,06,09,10,11,12,13	2000	corer	ICP-OES	na	?	Y	N	Y	Y	
Dublin	1991,95,96,97,2003,08,13,14	2006	corer	ICP-OES	HNO ₃	M	?	Y	Y	N	
Hong Kong	1987-2015; 2003-2015	2015/16	v v grab	ICP-MS	HNO ₃	M	?	Y	Y	Y	
Ravenna	1994,2004,09,18	2004	na	GF-AAS	HNO ₃ /HCl	M	Y	N	N	N	
Ria de Vigo	2011	2011	na	SP	HNO ₃	M	Y	Y	Y	Y	
Rio de Janeiro	2005,06 ¹	2005/06	v v grab	ICP-OES	6M HCl	W	N	Y	Y	Y	
Santander	2005,6,7,8,9,10,11,12,13,14,15	2015	v v grab	ICP-MS	HNO ₃ /HCl	M	Y	Y	Y	N	
Sydney	2010-14	2010-14	Box corer	ICP-OES	Aqua regia	W	Y	Y	Y	Y	SN

Notes: PEN=Post Extraction Normalised; SN=size normalised; S=strong; W=weak; M=Moderate; v v=Van Veen; RM=reference material used

ICP=Inductivity coupled plasma; MS=mass spectrometry; OES=optical emission spectrometry;

SP=spectrophotometer; GF-AAS=graphic furnace atomic adsorption spectroscopy; na=not available

HNO₃= nitric acid; HCl=hydrochloric acid; HClO₄=perchloric acid; aqua regia=HCl/HNO₃

¹ mean of four surveys

Table 2. Data availability and reliability

Location	Number of samples	Harbour area (km ²)	Sample density sample/km ²	Map type	Distribution	Reliability
						L M H
Auckland	121	800	0.15	P	Sites mainly in bays & tributaries with central area unsampled	M
Darwin	298	1220	0.24	P	Sites restricted to intertidal flats with tidal channels unsampled	H
Derwent	123	200	0.62	FC	Excellent distribution, all locations and environments covered	H
Dublin	42	3.267	12.9	P	Moderate number of sites mainly in shipping basins	M
Hong Kong						
Harbour	9	63.0	0.14	FC	Excellent coastal dataset but only harbour sites used	L
Ravenna	52	3.62	14.4	P	High density sampling in shipping channels	H
Ria de Vigo	39	156	0.25	FC	Good systematic grid covering all locations & environments	H
Rio de Janeiro	28	449 ¹	0.06	FC	All areas covered but low sample density	M
Santander	10	22.5	4.58	P	Few sites located mainly in marginal areas	L
Sydney	1175	50	23.5	FC	Good distribution, high sample density covers all areas	H

H= high; M=moderate; L=low; FC=filled contour maps; P=point maps

¹ Guanabara Bay area

Table 3. Attributes of concentration and enrichment Factors/Indices used globally

No.	Factor/Index Name	Symbol	Author(s)	Background		Normalis- ation	Multi- element	Classificat- ion Scheme	Score
				Global	Local				
Concentration Factors (no background, or normalisation)									
1	Metal Pollution Index	PI	Usero et al.,1996	N	N	N	Y	N	1
Contamination Factors (background values, but no normalisation)									
2	Contamination Factor	CF	Brady et al., 2015	Y	N	N	N	N	1
3	Nemerow Pollution Index	PI	Nemerow, 1991	N	N	N	Y	N	1
4	Surface Enrichment Factor	SEF	Riba, 2002a	Y	Y	N	N	N	2
Enrichment Factors (apply both background and normalisation procedures)									
5	Enrichment Factor	SEF	Kemp et al., 1976; Rubio et al., 2000	Y	Y	Y	N ¹	N	3
6	Geo-accumation Index (fine sediment)	<i>I_{geo}</i> (fine)	Muller, 1980	Y	Y	Y	N	Y	4
7	Geo-accumation Index (total sediment)	<i>I_{geo}</i> (total)	Xu et al., 2014	Y	Y	Y	N	Y	4
8	Geo accumulation Index (total sediment)	<i>I_{geo}</i> (mud %)	Kim et al., 2018	Y	Y	Y	N	Y	4
9	Degree of Contamination	DC	Hakanson, 1980	Y	N	Y	Y	Y	4
10	Modified Nemerow Pollution Index	mPI	Brady et al., 2015	Y	Y	Y	Y	Y	4
11	Mean Enrichment Quotient	MEQ	Birch & Olmos, 2008; Birch et al., 2013	Y	Y	Y	Y	Y	5
12	Modified Degree of Contamination ²	MDC	Abraham & Parker, 2008	Y	Y	Y	Y	Y	5

¹ combined elements as overlays on maps; ² use 'fine-grained' sediments

Table 4. Mean, minimum and maximum concentrations (µg/g, dry weight) for total sediment

Harbour	Cd (1.2; 9.6)			Cr (81; 370)			Cu (34; 270)			Ni (20.9; 51.6)			Pb (46.7; 218)			Zn (150; 410)		
	Mean	Min	Max	Mean	Min	Max	Mean	Min	Max	Mean	Min	Max	Mean	Min	Max	Mean	Min	Max
Auckland	0.1	0.0	0.1	15	3.1	23	13	3.1	36	na	na	na	22	5.4	44	90	16	210
Darwin	0.1	1	0.5	18	6.6	95	5.4	0.7	23	8.8	1.5	27	10	1.8	50	25	4.6	190
Darwin ¹	0.1	bd	0.9	45	44	230	15	1.0	147	22	3.5	113	29	6	459	79	6.8	1730
Derwent	14	1.0	128	36	2.0	71	112	1.0	591	17	1.0	31	450	6.0	0	2130	22	0
Dublin 2013	3.6	0.1	20	136	25	316	60	3.3	155	90	58	164	138	10	0	660	22	5240
Dublin 2008	0.7	0.1	2.3	24	3.7	47	27	1.3	69	19	1	39	44	1.9	192	152	9.2	470
Dublin 2006	1.1	0.4	2.1	35	9.3	66	49	8.5	111	29	12	55	81	11	243	217	60	420
Hong Kong ²	0.2	0.1	0.7	31	17	67	31	7.0	160	18	10	29	41	19	100	121	51	330
Hong Kong ³	0.3	0.1	1.7	37	22	55	67	23	135	18	14	24	45	29	74	148	72	275
Ravenna	1.6	1.0	2.4	74	46	124	30	8.6	72	na	na	na	49	11	126	106	67	180
Ria de Vigo	0.3	bd	1.3	70	4.2	198	82	4.2	479	29	6.1	43	111	5.0	450	199	13	1490
Rio de Janeiro	0.8	bd	2.4	72	1.4	205	62	0.3	224	21	0.2	113	66	2.7	200	318	4.2	2040
Santander	1.3	0.1	2.6	77	7.5	214	33	2.5	67	46	4.2	126	83	12	145	463	52	1050
Sydney	1.0	bd	52	77	bd	298	133	bd	1060	15	bd	118	210	bd	0	518	bd	0
Mean	1.8	0.3	17	54	15	144	51	5.0	237	28	11	74	98	8.7	502	371	32	2960
Maximum	14.0	1.0	128	136	46	316	133	23	1060	90	58	164	450	29	0	2130	72	0

¹ for <62.5 µm fraction; ² coastal region; ³ for Harbour area only; bd=below detection

ERL= Effects Range Low; ERM=Effects Range Median; Figures in brackets are ERL; ERM values

Table 5. Mean and maximum enrichment based on various normalising techniques

Location	Cd (0.2 µg/g)		Cr (60 µg/g)		Cu (20 µg/g)		Ni (30 µg/g)		Pb (25 µg/g)		Zn (90 µg/g)		MEQ		Normalisor & concentration
	Mean	Max	Mean	Max	Mean	Max	Mean	Max	Mean	Max	Mean	Max	n=6	n=3	
Auckland	na	na	na	na	na	na	na	na	na	na	na	na	na	na	No Al or Fe
Darwin	0.7	11	0.7	3.8	0.7	3.9	0.7	3.8	1.0	7.5	0.9	19.0	0.8	0.9	PEN
Darwin	1.6	11	1.2	4.5	0.9	4.1	1.0	2.1	1.4	5.6	1.0	5.6	1.2	1.1	70K Al
Derwent	77	430	0.8	2.7	5.6	22	0.7	2.1	19	57	23	129	21	16	32K Fe
Dublin 2006	10	45	1	1.6	4.1	7.7	1.8	5.3	5.4	12	4.2	8	4.5	4.6	25K Al
Hong Kong ¹	1.9	24	1.0	1.1	5.1	8.8	1.0	1.3	2.9	5.1	2.6	3.8	2.4	3.5	35K Al
Hong Kong ²	1.2	6.4	0.8	1.6	2.5	11	1.4	2.6	2.5	6.5	2.1	7.8	1.7	2.2	35K Al
Ravenna	na	na	na	na	na	na	na	na	na	na	na	na	na	na	No size, Al, Fe
Ria de Vigo	1.9	7.3	1.4	3.2	4.9	19	1.2	3.9	5.2	19	2.7	10	2.9	4.1	90K Al
Ria de Vigo	1.8	5.8	1.3	3.1	4.9	4.0	1.2	2.7	5.0	17	2.6	16	2.8	4.1	35K Fe
Rio de Janeiro	5.4	16.7	1.5	4.0	4.0	13	0.9	4.4	3.1	9.3	4.6	26	3.2	3.5	20K Al
Santander	9.9	21	3.5	13	2.5	3.3	4.0	16	5.2	7.3	9.5	18	5.8	5.8	27K Al
Sydney	9.5	69	2.4	7.5	9.8	35	0.8	5.3	13	53	7.6	30	4.5	9.6	35K Al
Sydney	5.1	25	2.0	5.8	9.1	36	0.7	2.1	11	52	6.5	21	5.7	8.6	SN
Mean	11	56	1.5	4.3	4.5	14	1.3	4.3	6.2	21	5.6	25	4.7	5.3	
Maximum	77	430	3.5	13	9.8	36	4.0	16	19	57	23	129	21	16	

Figures in brackets are background values; MEQ=Mean Enrichment Quotient; na= not available

MEQ n=6 is for Cd, Cr, Cu, Ni, Pb and Zn; MEQ n=3 is for Cu, Pb and Zn; PEN=post-extraction normalisation; SN=size normalised; K=1000 µg/g

¹ coastal region; ² for Harbour area only

Table 6. Results of enrichment computations using multiple indices

	Concentration Factor	Contamination Factors (background, no normalisation)				Enrichment Factors (background and normalisation applied)						
	MPI	CF	PI	SEF	Igeo (total)	EF	Igeo (AI)	mPI	mDC	MEQ	WoE	Rank
Harbour	Metal pollution index	Contamination Factor	Nemerow pollution index	Surface enrichment factor	Geo-accumulation index total sediment	Enrichment factor	Igeo AI normalised	Modified Nemerow pollution index	Modified degree of contamination	Mean enrichment quotient	Total score	
Auckland	7.5	0.6	49	-0.4	-5.6	na	na	na	0.6	na	na	na
Darwin	5.0	0.3	46	-0.5	-2.0	1.0	-0.8	0.2	0.3	1.1	4	8
Derwent	99	20	2070	23	1.3	25.0	2.2	5.3	20	16.0	20	1
Dublin	31	2.5	117	1.8	0.4	5.0	0.5	0.6	2.5	5.8	11	6
Hong Kong	16	1.1	86	0.1	0.5	1.8	0.5	1.5	1.1	2.2	5	7
Ravenna	26	1.8	81	1.2	0.1	na	na	na	2.2	na	na	na
Ria de Vigo	12	2.4	320	1.7	0.4	6.1	0.9	1.1	2.4	4.1	9	5
Rio de Janeiro	34	2.5	335	1.7	1.9	2.8	0.9	2.5	2.5	2.5	11	4
Santander	40	3.0	208	2.4	0.5	5.0	1.8	1.2	3.0	5.8	12	3
Sydney	50	4.5	1740	4.2	0.8	8.7	1.7	5.6	4.5	8.6	14	2

na=not available; WoE=Weight of evidence based on classification schemes for EF, mDC, mPI and MEQ techniques, see text

(Section 4.3); Rank Least influenced 1 to most impacted 8

Table 7. Mean Effects Range Median (MERM) for individual metals and for metal mixtures (Mean Effects Range Median Quotients, MERMQ)

Harbour	MERM (Cd, Cr, Cu, Ni, Pb, Zn)						MERMQ
	Cd	Cr	Cu	Ni	Pb	Zn	
Auckland	0.01	0.02	0.05	na	0.11	0.24	0.09
Darwin	0.01	0.05	0.02	0.17	0.05	0.06	0.06
Derwent	1.42	0.10	0.41	0.32	2.04	5.20	1.58
Dublin 2006	0.1	0.1	0.18	0.56	0.37	0.53	0.31
Hong Kong ¹	0.02	0.09	0.12	0.35	0.19	0.30	0.17
Hong Kong ²	0.03	0.02	0.25	0.35	0.21	0.36	0.20
Ravenna ³	0.16	na	na	na	0.18	0.43	0.26
Ria de Vigo	0.04	0.15	0.44	0.48	0.6	0.59	0.38
Rio de Janeiro	0.09	0.20	0.23	0.41	0.28	0.78	0.33
Santander	0.13	0.21	0.12	0.87	0.37	1.25	0.51
Sydney	0.09	0.22	0.43	0.27	0.8	0.95	0.53

¹ coastal region; ² for Harbour area only; ³ for Cd, Pb and Zn only; na= not available

Table 8. Effects range low (ERL) and effects range median (ERM) quotients and numbers and percentages of samples >ERL and >ERM

Harbour	Sample No.s	ERLQ (Cd, Cr, Cu, Ni, Pb, Zn)			ERMQ (Cd, Cr, Cu, Ni, Pb, Zn)						MERLQ Cu, Pb, Zn	MERMQ Cu, Pb, Zn
		Mean	Samples >ERL ¹		Mean ERMQ	Samples >ERM ¹		No. samples exceeding ERM				
			No.	%		No.	%	1-3	3.1-5	5.1-6		
Auckland	121	0.48	0	0	0.08	0	0	0	0	0	0.48	0.12
Darwin	298	0.21	2	<1	0.06	0	0	3	0	0	0.18	0.04
Derwent	123	6.50	102	83	1.60	111	90	51	20	40	8.70	2.56
Dublin 2006	42	1.23	36	88	0.34	2	5	2	0	0	1.43	0.45
Hong Kong	9	0.67	9	100	0.23	0	0	0	0	0	0.87	0.27
Ravenna ³	52	0.89	26	54	0.20	0	0	0	0	0	0.88	0.20
ria de Vigo	39	1.46	34	87	0.35	6	15	5	1	0	2.04	0.43
Rio de Janeiro	28	1.31	17	60	0.33	4	15	4	1	0	0.78	0.40
Santander	10	1.57	9	90	0.51	8	80	8	0	0	2.07	0.92
Sydney	207	2.00	195	94	0.53	109	53	107	2	0	2.75	0.51

¹ samples with any one metal >ERL or > ERM; ² for mean Cd, Cr, Cu, Ni, Pb and Zn; ³ for mean Cr, Cu, Pb and Zn only

ERL= Effects Range Low; ERM=Effects Range Median; PEL= Probable Effects Level; Q=Quotient

Table 9. Harbour ranking for Mean Enrichment Quotient (MEQ), ten environmental indices and Mean Effects Median Quotient (MERMQ)

Harbour	MEQ	Environmental Indices	MERMQ	Rank ¹	
				Total	Place
Auckland	na	na	9	na	na
Darwin	8	8	10	26	1
Derwent	1	1	1	3	8
Dublin 2006	4	6	6	16	4
Hong Kong	7	7	8	24	2
Ravenna	na	na	7	na	na
Ria de Vigo	5	5	4	14	5
Rio de Janeiro	6	4	5	15	3
Santander	3	3	3	9	6
Sydney	2	2	2	6	7

¹ Least influenced 1 to most impacted 8; na= not available

Table 10. Magnitude of anthropogenic change (MAC) and Ecological risk posed by sedimentary contaminants (ERA)

Harbour	Magnitude of anthropogenic change (MAC) ¹		Ecological risk posed by sedimentary metals (ERA) ²	
	MEQ	Enrichment/Modification	MERMQ	Ecological Risk
Auckland	na	na	0.09	Minimal risk
Darwin	1.2	Not enriched	0.06	Minimal risk
Derwent	21	Highly enriched	1.58	High risk
Dublin	5.7	Slightly enriched	0.31	Slight risk
Hong Kong Harbour	1.7	Slightly enriched	0.20	Slight risk
Ravenna	na	na	0.26	Slight risk
Ria de Vigo	2.8	Slightly enriched	0.38	Slight risk
Rio de Janeiro	3.2	Moderately enriched	0.33	Slight risk
Santander	5.8	Highly enriched	0.51	Moderate risk
Sydney	5.7	Highly enriched	0.53	Moderate risk

MEQ= Mean enrichment quotient; MERMQ= Mean effects median quotient; na=not available

¹ MEQ <1.5 - not enriched; 1.5-3.0 - slightly enriched; 3.0-5.0 - moderately enriched; >5.0 - highly enriched

² MERMQ <0.1 - minimal risk; 0.1 - 0.5 - slight risk; 0.5 - 1.5 - moderate risk; >1.5 high risk

Figure Captions

Figure 1. The ten world harbour estuaries assessed in the current WHP

Figure 2. Sample sites in six of the WHP locations

Figure 3. Distribution of fine (<62.5 µm) sediment in six of the WHP locations

Figure 4. Distribution of Zn in total sediment for six of the WHP locations

Figure 5. Distribution of Zn enrichment for six of the WHP locations

Figure 6. Distribution of Mean Enrichment Quotient (MEQ) (Cu, Pb and Zn) for six of the WHP locations

Figure 7. Distribution of Ecological Risk Assessment (ERA) for six of the WHP locations. ERL=Effects Range Low; ERM=Effects Range Median

Figure 8. Distribution of Mean Effects Range Median Quotient (MERMQ) for six of the WHP locations

Figure 9. Ecological Risk for world harbour estuaries assessed in the current WHP expressed as Effects Range Median (ERM) with Mean Effects Range Median Quotient (MERMQ for six metals) (a) including the Derwent River and (b) excluding the Derwent River to emphasise remaining locations



Figure 1.

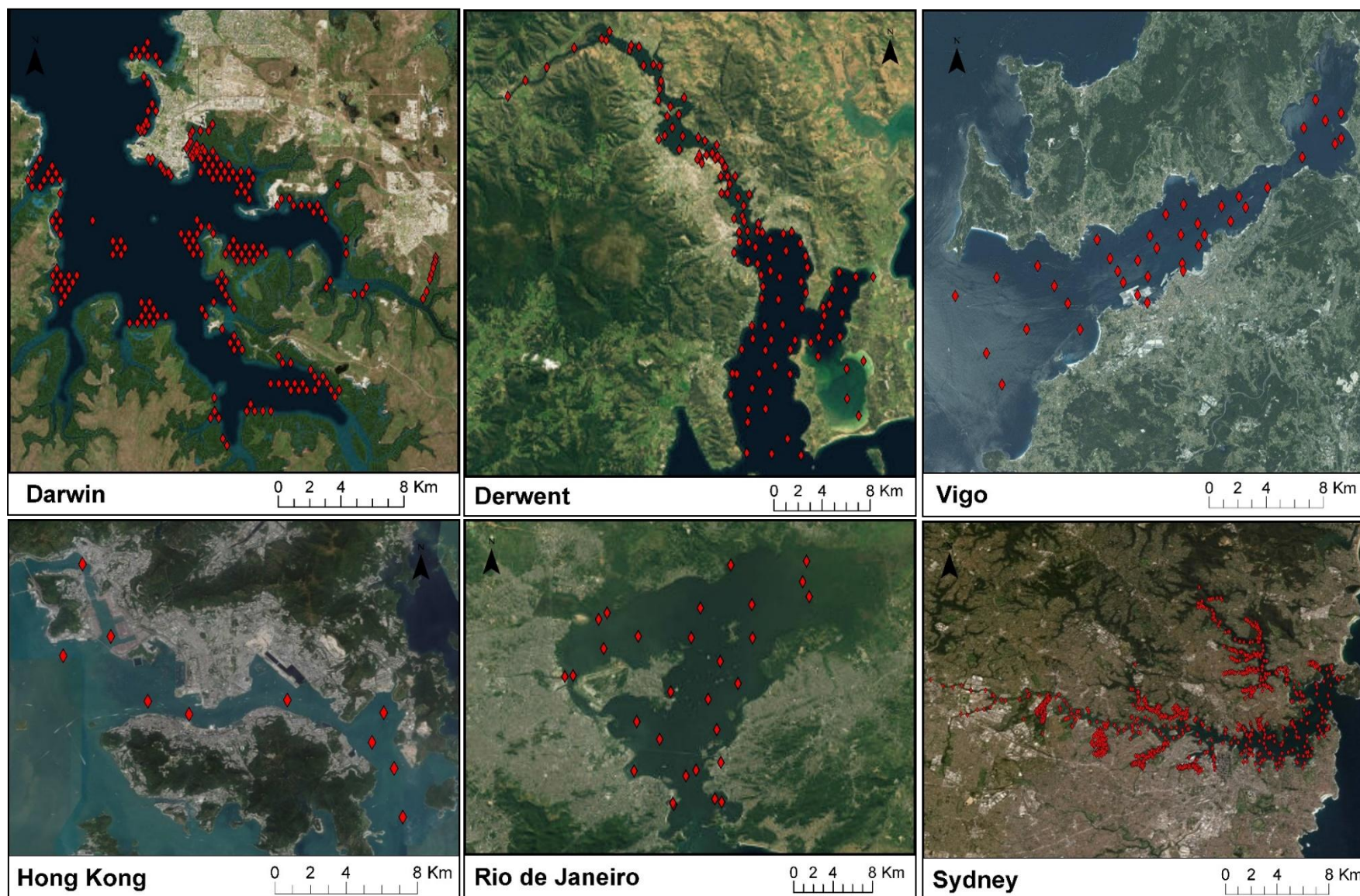


Figure 2.

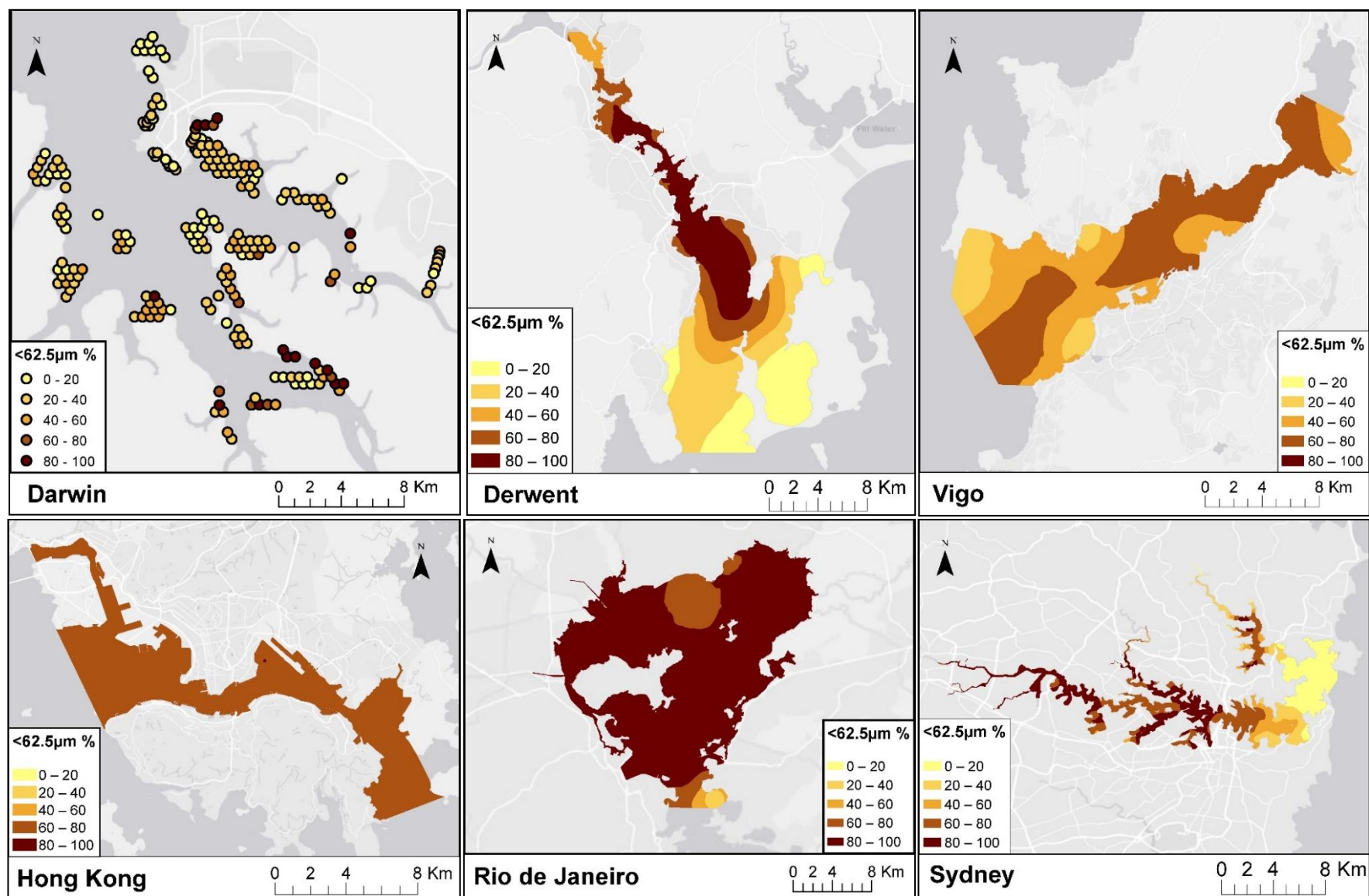


Figure 3.

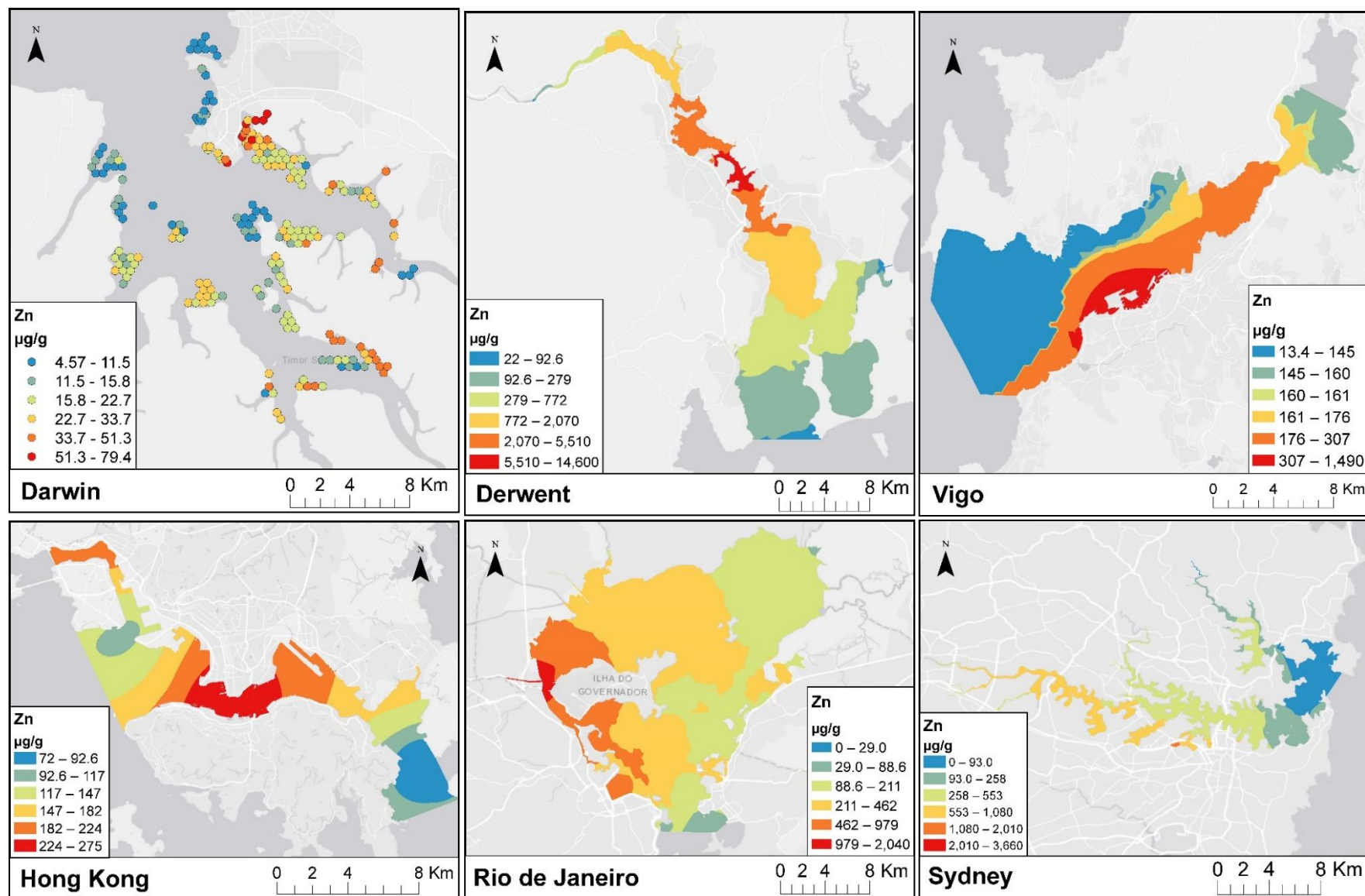


Figure 4.

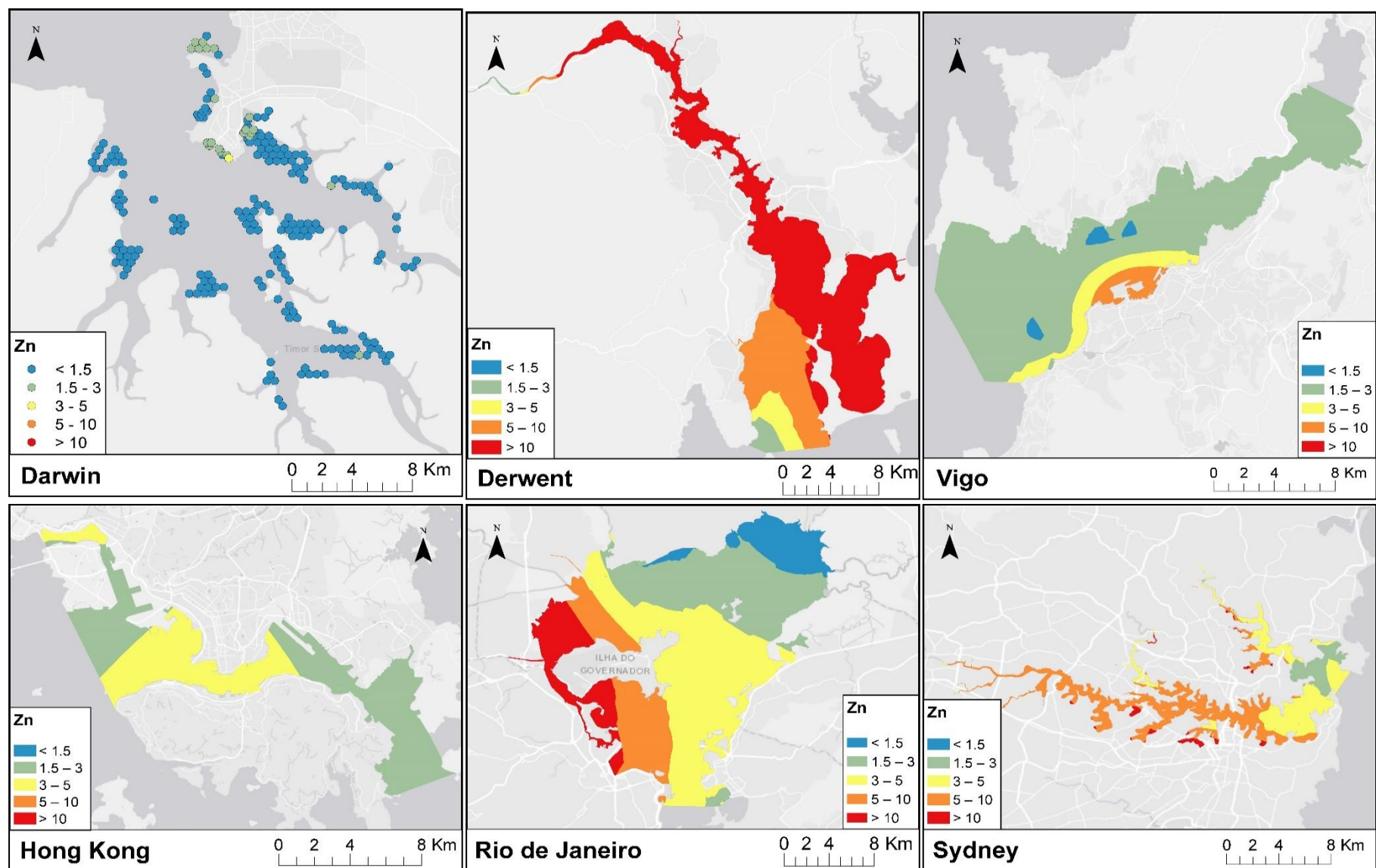


Figure 5.

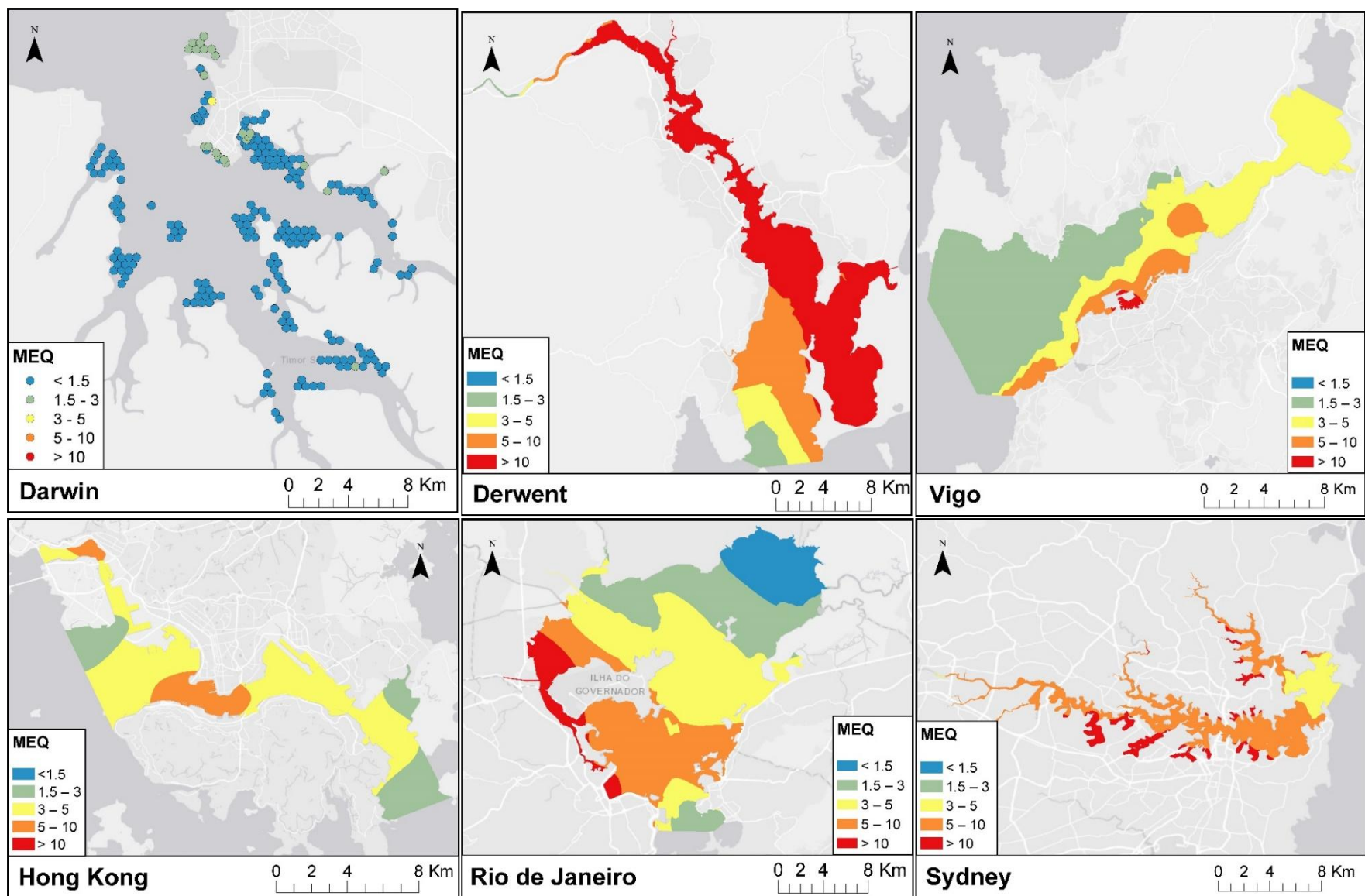


Figure 6.

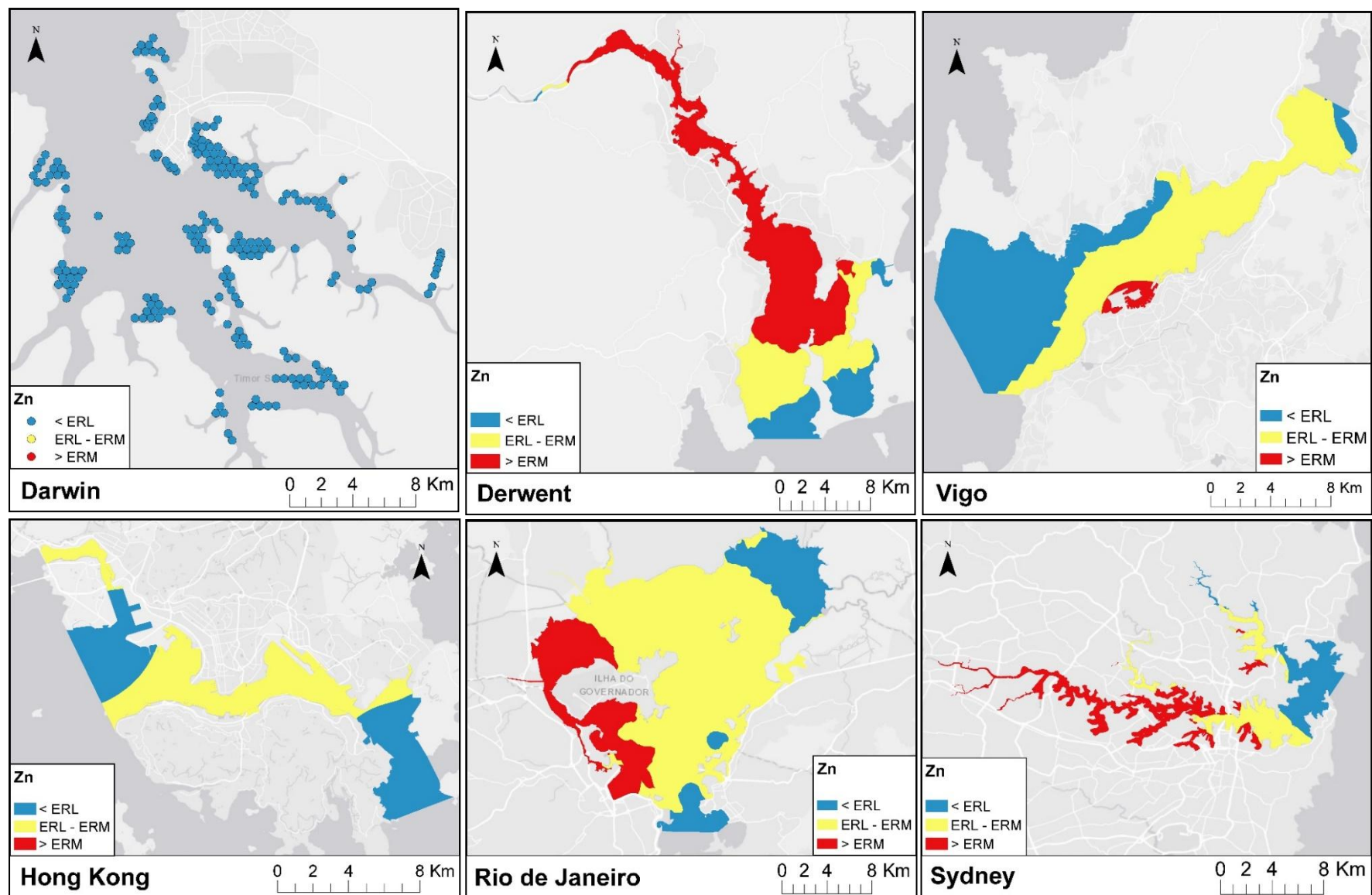


Figure 7.

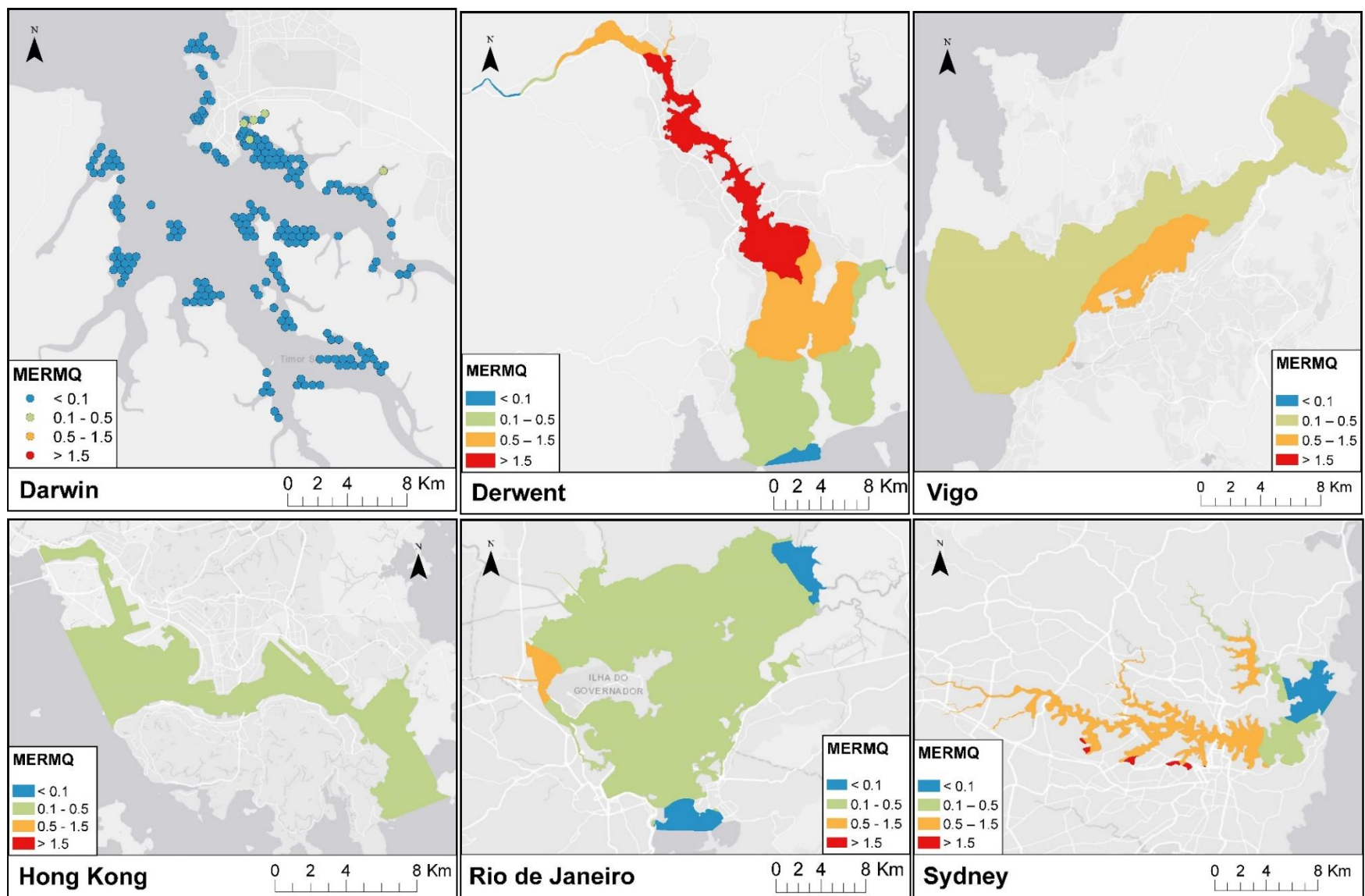


Figure 8.

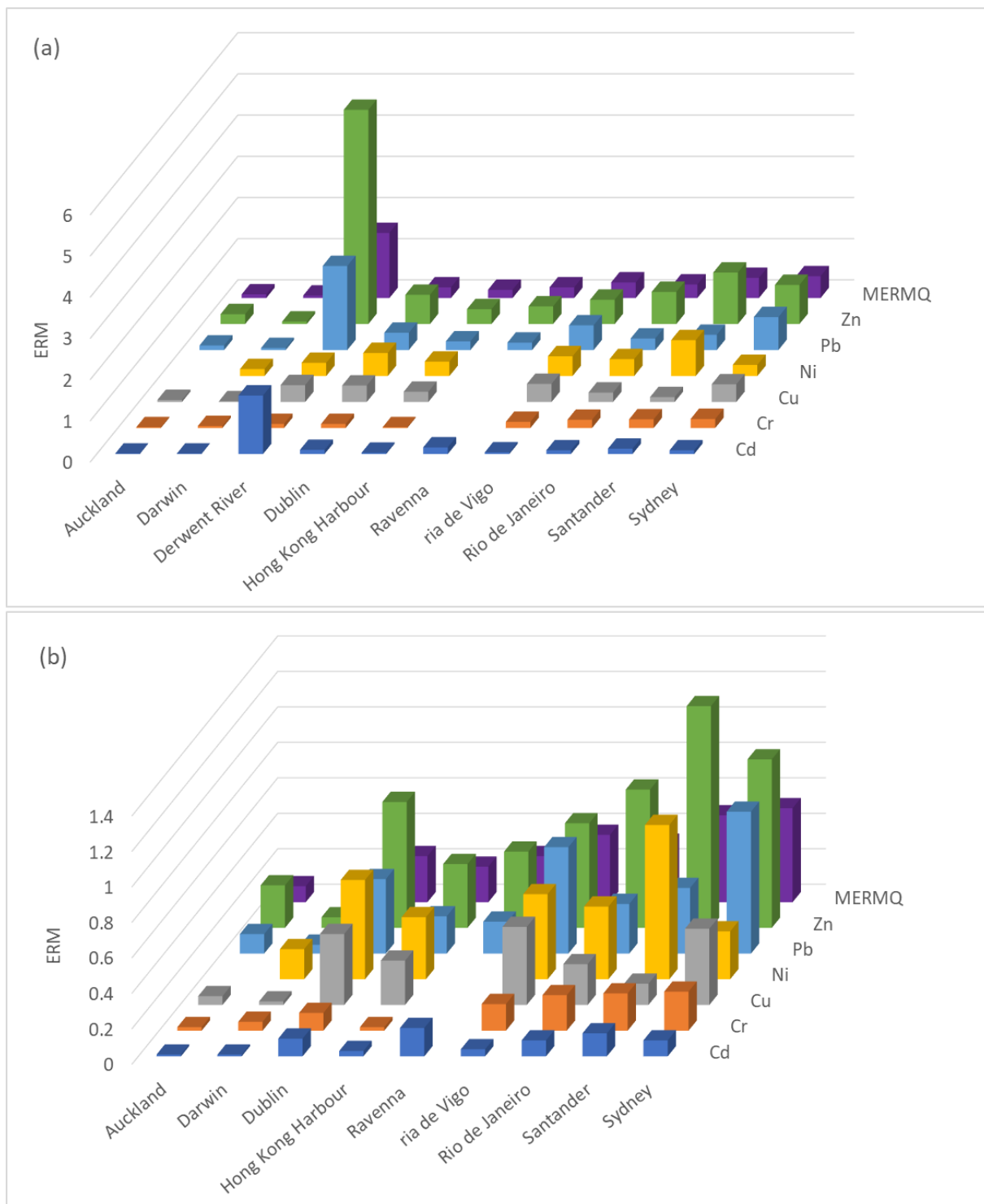


Figure 9.

Credit Author Statement

G F Birch wrote most of the paper and undertook most of the computations

J-H Lee drafted the maps and diagrams and undertook some of the computations. He also read many versions of the early drafts

E Tanner managed the project, provided advise on the locations and co-authors and read many drafts of early versions of the paper

The remaining co-authors provided data on the harbours and read drafts of early versions of the paper