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Global Environmental Change and Noncommunicable Disease Risks

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Abstract

Multiple global environmental changes (GECs) now under way, including climate change, biodiversity loss, freshwater depletion, tropical deforestation, overexploitation of fisheries, ocean acidification, and soil degradation, have substantial, but still imperfectly understood, implications for human health. Noncommunicable diseases (NCDs) make a major contribution to the global burden of disease. Many of the driving forces responsible for GEC also influence NCD risk through a range of mechanisms. This article provides an overview of pathways linking GEC and NCDs, focusing on five pathways: (a) energy, air pollution, and climate change; (b) urbanization; (c) food, nutrition, and agriculture; (d) the deposition of persistent chemicals in the environment; and (e) biodiversity loss.

INTRODUCTION

Noncommunicable Diseases

Noncommunicable diseases (NCDs), principally cardiovascular diseases, cancers, chronic respiratory diseases, diabetes, and mental health conditions, together with neurologic, endocrine, gastrointestinal, renal, allergic, and autoimmune disorders, have commanded increasing attention in recent years, and with good reason. These diseases kill more than 38 million people each year, accounting for 70% of global deaths (95, 156). Among the victims are more than 14 million adults who, in the words of the World Health Organization (WHO), “die too young”—that is, between the ages of 30 and 70 (155). NCDs also account for 21 of the leading 30 causes of age-standardized years lived with disability (YLDs) in 2016, or 80.6% [95% confidence interval (CI) 78.2–82.5] of YLDs, according to the Global Burden of Disease study (145). Leading contributors to disability include headache, low back pain and osteoarthritis, depression and anxiety, diabetes, asthma, and vitamin A deficiency.

Four aspects of NCDs are especially salient. First, they are not only, or even mostly, a problem of wealthy nations. Low- and middle-income countries (LMICs) account for 86% of the burden of premature deaths from NCDs (19, 155), reflecting what has been known for nearly half a century as the “epidemiologic transition” (101). Second, the economic implications of NCDs are substantial, with a projected cost to the global economy of \$47 trillion over the next 20 years (14). Particularly in LMICs, NCDs slow economic development and trap millions of people in poverty. Third, many risk factors for NCDs are environmental in origin or may be influenced by the environment. For example, about 50% of the disease burden from chronic obstructive pulmonary disease and about one-quarter of the ischemic heart disease burden have been attributed to environmental factors, particularly air pollution (79). However, environmental contributions to NCDs are routinely overlooked; for example, the 2013 WHO NCD Global Action Plan largely failed to mention environmental factors (155) [although a later report from WHO did address the issue (158)]. Fourth, NCD research, prevention, and treatment are severely underfunded, particularly in LMICs, relative to their population burden (3).

A debate has arisen concerning the framing of NCDs, and even the term NCD. Some have argued against the name because it starts with “non,” defining an entity by what it is not and seemingly making it a “nonissue.” Others have pointed to increasing evidence that NCDs are, in fact, communicable, through social networks, viruses, and/or cultural and economic conditions (4). Suggested replacement terms include “biosocial and development diseases” (160) and “life-long disease” (112). These terms, however, may overlook a set of causal factors whose importance is increasingly clear but largely overlooked to date: those related to global environmental changes.

Global Environmental Changes

Since at least the middle of the last century, unprecedented global environmental changes (GECs) have occurred. These changes undermine many of the natural systems that underpin human civilization and have the potential to reverse the progress in health and development achieved in recent decades. These changes include climate change, biodiversity loss, freshwater depletion, tropical deforestation, overexploitation of fisheries, ocean acidification, soil degradation, changes in nutrient (nitrogen, phosphorus, and others) cycling, and environmental loading with persistent toxic substances. **Figure 1** summarizes some of these environmental trends. Because these human-induced impacts on earth systems are so extensive, they have been proposed to signal a new geological epoch, the Anthropocene (131). The far-reaching implications of these changes

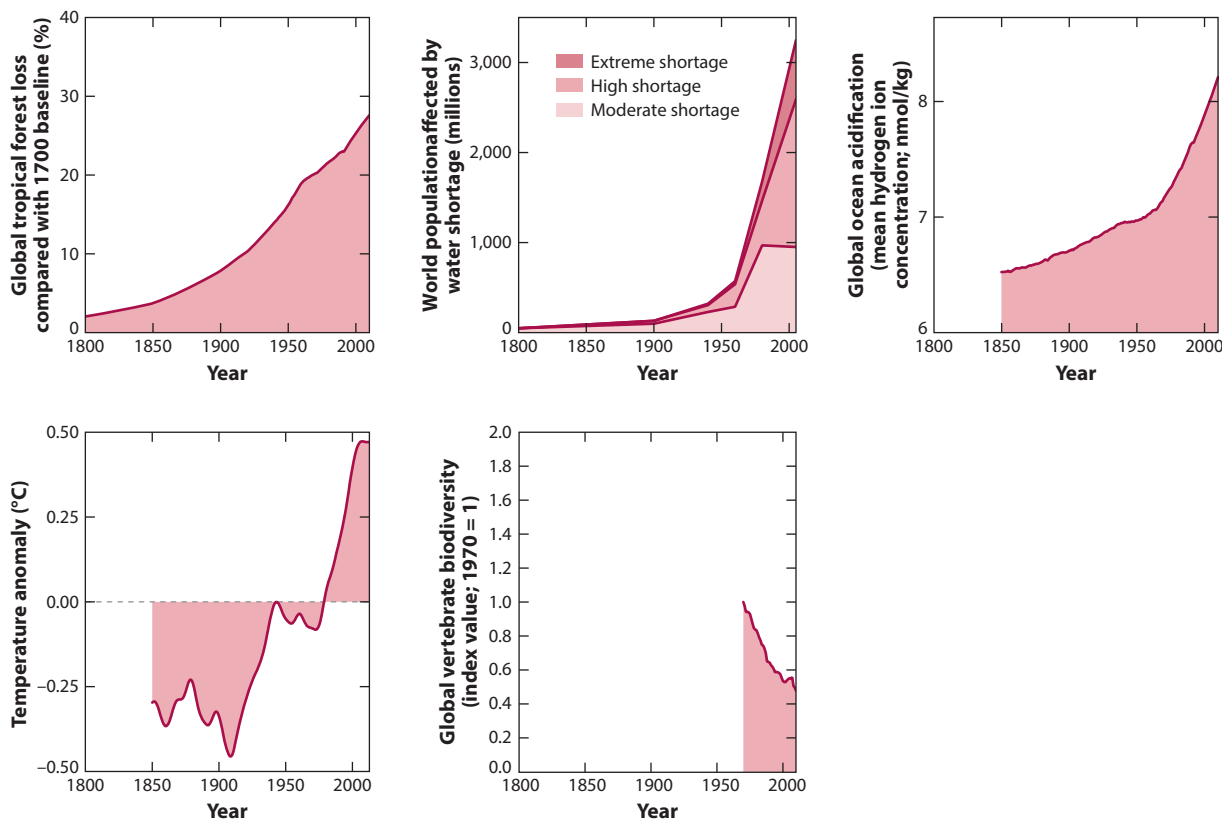


Figure 1

Selected global environmental trends. Figure adapted from Reference 154.

for human health have been described by The Rockefeller Foundation/Lancet Commission on Planetary Health (154).

GECs can affect a range of health outcomes through a complex web of pathways. **Figure 2** shows a subset of these pathways, focused on climate change and related processes (152). A diagram showing other processes, from changes in nitrogen cycling to biodiversity loss, would be considerably more complex. Key pathways to health impacts include extreme events (floods, droughts, intense storms, wildfires, and heat waves), changes in the incidence and distribution of both vector-borne and water-related diseases, reduction in food availability and quality, and socially mediated effects such as increased poverty, conflict, and population displacement (152, 154).

The pathways from GECs to NCD risk can also be represented through the DPSEEA (driving forces, pressures, states, exposures, effects, actions) framework (**Figure 3**) (46). Strategies to prevent and treat NCDs are often narrowly focused on clinical interventions or behavior changes. These approaches are essential but not sufficient; the failure to address many of the key driving forces increasing the risks of NCDs will likely limit the impacts of these preventive strategies (103).

This article outlines pathways by which GECs, and the driving forces responsible for them, influence the risks of NCDs. It considers five pathways: (a) energy, air pollution, and climate change; (b) urbanization; (c) food, nutrition, and agriculture; (d) persistent environmental chemicals; and

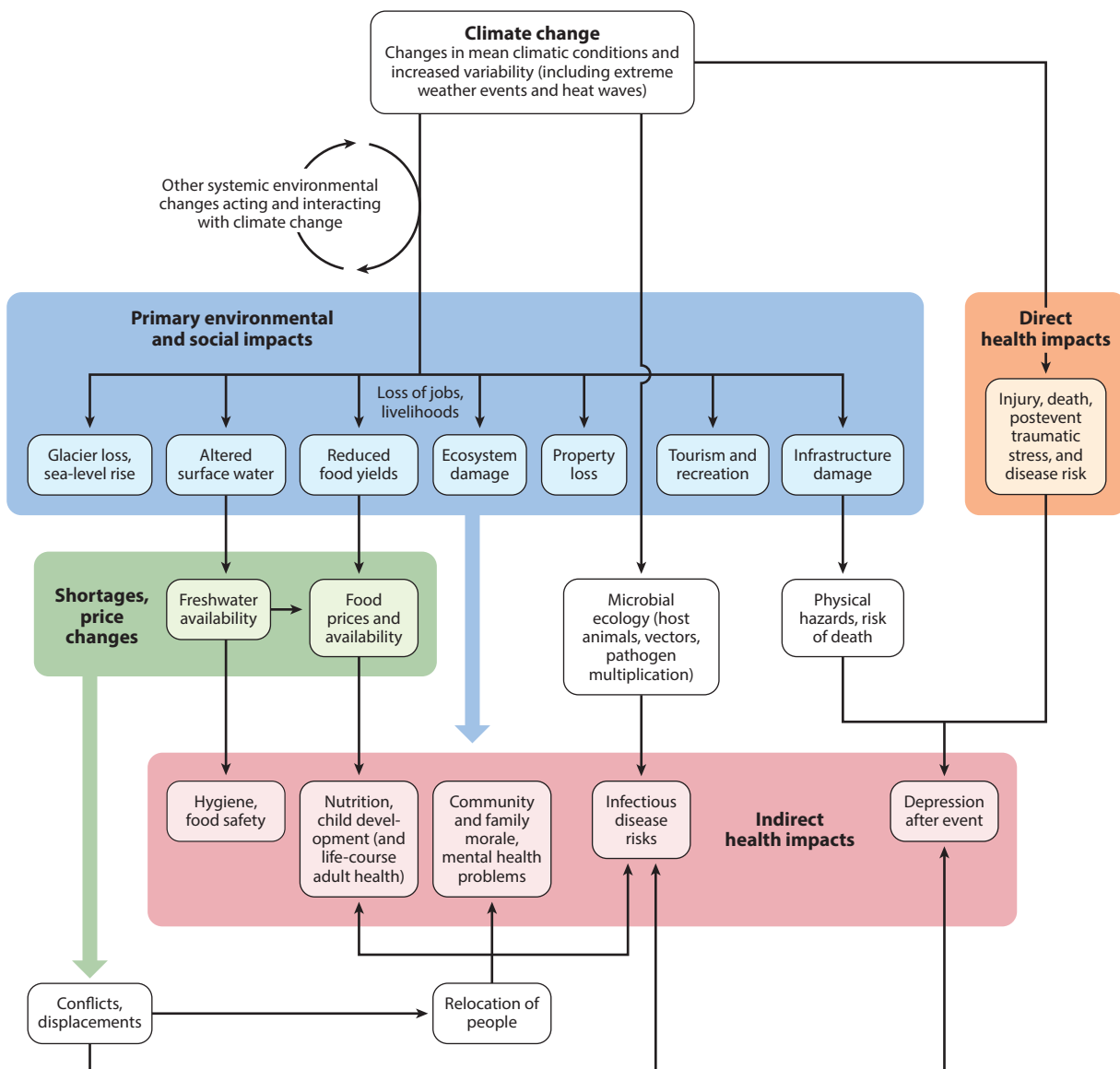


Figure 2

Pathways from climate change to health outcomes. The orange field shows direct health impacts, such as the effects of severe weather events. The blue field shows social impacts that follow climate-related environmental changes; many of these relate to reduced access to needed resources such as water, to infrastructure damage, and to loss of livelihoods. Less direct health impacts are shown in the lower part of the figure. These are mediated by the primary environmental and social impacts of climate change. The red field shows five categories of indirect health impacts. Tertiary effects, indicated by upward-pointing arrows, arise from more diffuse disruptions, dislocations, and conflicts, which are likely to increase with advancing climate change. The figure is not intended to be comprehensive. Figure adapted with permission from Reference 92.

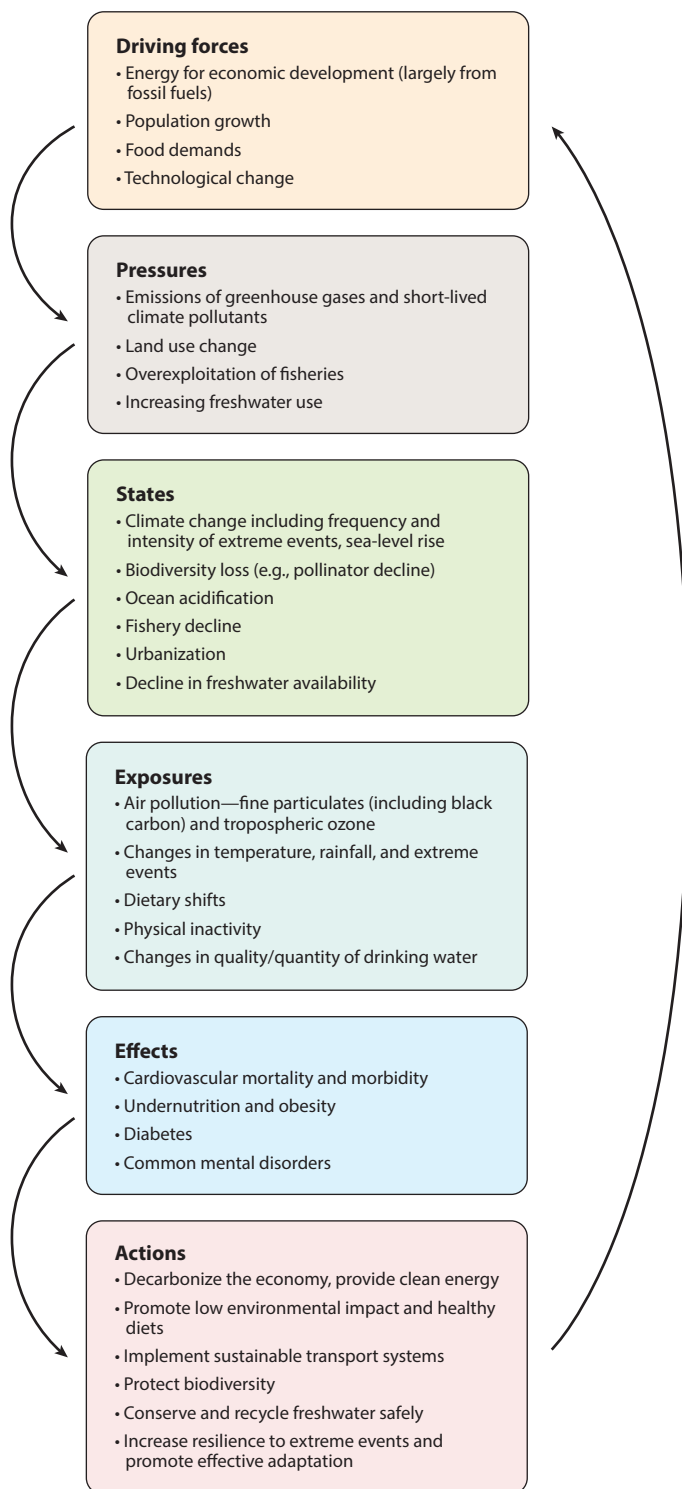


Figure 3

The DPSEEA (driving forces, pressures, states, exposures, effects, actions) framework as applied to global environmental changes. Figure adapted with permission from a figure by James Milner.

(e) biodiversity loss (including fisheries depletion). Land use changes, a sixth category, are considered in the context of urbanization, agriculture, and biodiversity loss.

While GECs contribute to the risk of NCDs, the association has potentially positive implications as well. The measures required to limit GECs and to promote sustainable development also affect NCD risk, often in beneficial ways (61, 152). Well-designed policies to reduce environmental change may yield ancillary health benefits (cobenefits) and avoid unanticipated health risks (coharms) (61, 144).

THE IMPACT OF GLOBAL CHANGES ON NCDs

Energy, Air Pollution, and Climate Change

A principal driver of GEC is the combustion of the fossil fuels coal, oil, and gas; of these, coal has the largest effect. The dependence on fossil fuels for energy is a leading source of air pollution (66). It is also the single largest contributor to climate change. Two sets of pathways link energy use to NCDs, one through air pollution, the other through climate change.

Air pollution. Air pollution related to fossil fuel combustion is a major contributor to morbidity and mortality worldwide.¹ Key pollutants include fine particulate matter (PM), ozone, oxides of nitrogen, oxides of sulfur, hydrocarbons, and metals; many of these are related to each other through complex atmospheric chemistry. The Global Burden of Disease (GBD) Study estimates that ambient fine particulate matter (PM_{2.5}) alone ranked fifth globally as a mortality risk factor in 2015, accounting for an estimated 4.2 million deaths (7.6% of global deaths) and 103.1 million disability-adjusted life years (DALYs) (4.2% of global DALYs) in 2015 (27, 52). These deaths are not uniformly distributed; the greatest burdens are in China and India (with about 1.1 million annual deaths each), Russia (137,000 deaths), Pakistan (135,000 deaths), and Bangladesh (122,000 deaths). Of note, the WHO estimates of the burden of ambient air pollution are somewhat lower than those of the GBD Study: 3 million deaths and 85 million DALYs (157). The reasons for the different estimates have been reviewed by the Lancet Commission on Pollution (79). A more recent study, on the other hand, estimated a substantially larger burden: 8.9 million deaths globally in 2015 (18), in part because new evidence suggests that even low levels of PM air pollution confer risks to health. Unless there is a rapid transition toward clean renewable sources of energy, economic growth is projected to increase this burden substantially in some regions, such as southeast Asia, over the coming decades (77).

The excess mortality from ambient PM exposure is attributable primarily to NCDs: ischemic heart disease, cerebrovascular disease, chronic obstructive pulmonary disease, and lung cancer, with a small proportion of excess deaths—perhaps one in ten—due to lower respiratory infection (64). Black carbon, a short-lived climate pollutant (SLCP) and an important component of PM, appears to be a better indicator of harmful exposure than undifferentiated PM in short-term studies, but the evidence from long-term studies is inconclusive (68). Air pollution exposure, especially to PM, may aggravate cardiac arrhythmias (44) and heart failure (123). Systematic reviews suggest an association between air pollution exposure and diabetes risk (37, 63), and while the evidence is not conclusive, a recent analysis calculated that in 2016, ambient PM_{2.5} contributed to approximately 3.2 million incident cases of diabetes, 8.2 million DALYs caused by diabetes, and 206,000 deaths from diabetes (16). Evidence indicates that PM exposure may be neurotoxic across the life span, contributing to neurodevelopmental delays in children and to cognitive decline in older adults (25, 33), although these associations are inconsistent across studies and cannot be considered definitive.

¹Household air pollution is also a major global health hazard, but it is not discussed here because its link to planetary changes is less direct.

Ozone is formed from atmospheric precursors—hydrocarbons (methane and volatile organic compounds) and oxides of nitrogen—many of which are combustion products. Like PM, tropospheric ozone is also associated with excess mortality, although lesser in magnitude than PM (141). In the GBD data, exposure to ozone caused an additional 254,000 (95% CI 97,000–422,000) deaths and a loss of 4.1 million (95% CI 1.6 million–6.8 million) DALYs from chronic obstructive pulmonary disease in 2015 (27). More recent estimates using updated exposure–response relationships suggest at least a fourfold higher burden, with 1.04–1.23 million respiratory deaths in adults attributable to long-term ozone exposures. The largest increases in estimated attributable mortality were in northern India, southeast China, and Pakistan (90). The association between ozone exposure and mortality seems to relate both to short-term high exposures and to long-term exposure (7, 90). Short-term ozone exposure also triggers exacerbations of airway disease (asthma and chronic obstructive pulmonary disease), accounting for substantial numbers of emergency room visits and hospitalizations (69).

Climate change. Climate change presents another, overlapping set of pathways through which energy use and resulting GECs affect health (126), and NCDs in particular (46). Multiple direct and indirect mechanisms operate.

One pathway operates through the effects of climate-related disasters on health care. Climate change increases the intensity and frequency of extreme weather events. A systematic review of the impacts of cyclone, flood, and storm-related disasters on health care for people with NCDs, based on 48 studies from developed countries, found that people with cancer, diabetes, and cardiovascular diseases sustained increased risks of exacerbation of their health problems following such disasters. These outcomes were due to a range of factors including disruption of transport, weakened health systems including drug supply chains, loss of power, and evacuations of populations (115).

Another pathway operates through heat. The increasing hot weather that comes with climate change is associated with increases in mortality, in emergency presentations, and in hospital admissions due to cardiovascular, respiratory, and renal disease (126). Increased heat-related mortality with advancing climate change will likely outweigh any declines in cold-related mortality, particularly in tropical and subtropical regions and in Southern Europe (51, 126).

Climate change may increase the risk of cardiovascular disease through other mechanisms than the direct effects of heat. First, warmer weather is associated with sleep disturbance (100), which in turn is a risk factor for cardiovascular disease (21). Second, hot weather is associated with a reduction in physical activity (99), and reduced physical activity is a risk factor for cardiovascular disease (13). On the other hand, cold weather also reduces physical activity, so, in cold regions that become warmer, physical activity may increase. The net effect globally is likely to be negative (99). Third, rising sea levels, together with excessive groundwater withdrawals, land use changes, agricultural practices, and other factors, can lead to saline intrusion of groundwater in coastal areas (138). This, in turn, increases the salt intake of affected populations. A study in coastal Bangladesh suggested that pregnant women were exposed to more than double the limits recommended by the WHO/FAO (Food and Agriculture Organization of the United Nations) for salt intake in drinking water and that this exposure increased the risk of preeclampsia and hypertension in pregnancy (73). A further study of nonpregnant adults in coastal Bangladesh showed that sodium concentrations in drinking water were strongly associated with blood pressures after controlling for personal, lifestyle, and environmental factors and that changing to a less saline drinking water source reduced blood pressure (119). On the other hand, some evidence has linked warmer weather with lower blood pressure, which may act to reduce cardiovascular disease risk (147).

Climate change may also aggravate cancer risk. First, as noted above, physical activity declines with heat; sedentariness increases risk of some cancers. Second, evidence indicates that climate

change increases the production of aflatoxin, a liver carcinogen that contaminates some foods such as peanuts (11).

Climate change may increase the risk of kidney disease. Kidney stone formation seems to vary with temperature, perhaps a result of relative dehydration and resulting urinary concentration (136). Chronic kidney disease of unknown origin has been observed in working populations in hot places, such as among sugarcane cutters in Nicaragua (153), although a recent systematic review found inconsistent associations between heat and chronic kidney disease (88).

Climate change may affect NCD risk through a variety of nutritional pathways. These are discussed below in the section on food and nutrition.

Mental illness and distress are substantial contributors to the global burden of disease, and climate change may contribute in several ways (24). One way is a mechanism already mentioned: reduced physical activity due to heat. Physical activity is associated with reduced depression (91) and anxiety (133). Another pathway is the deleterious effect of disasters—which are increasingly common because of climate change—on mental health. Mental health problems in some circumstances dominate the health burden following disasters (56). Hurricane Katrina provides an example. Although not definitively attributed to climate change, this disaster exemplified the mental health burden that can follow climate-related disasters. Research conducted several months after Hurricane Katrina showed that approximately 49% of those surveyed in New Orleans, and approximately 26% in other hurricane-affected areas, suffered from a DSM-IV (*Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition*) anxiety or mood disorder, of which half or more was post-traumatic stress disorder (PTSD) (48). By two years after the hurricane, the prevalence of PTSD and depression had actually increased (72). Similar disease burdens have been documented following other kinds of disasters, which are predicted to become more frequent with climate change, such as severe storms (1), floods (42), heat waves (62), and wildfires (35). Risk factors for postdisaster depression and PTSD across the life span have been studied in both wealthy and resource-poor settings; these include predisaster mental and physical health, personality structure and beliefs, features of the disaster experience itself, postdisaster social support and services, postdisaster displacement, and others (135). Slow-moving climate-related disasters may also threaten mental health. For example, research in Australia during the recent decade-long drought revealed an increase in anxiety, depression, and possibly suicidality among rural populations (98).

Of note, the two sets of pathways discussed here that link energy generation and NCDs—through air pollution and through climate change—are not independent. Certain air pollutants function as SLCPs and promote climate change while also threatening health via direct toxic effects (e.g., black carbon, tropospheric ozone) or by giving rise to ozone (methane) (124).

Urbanization

The city has become the prototypical human habitat, representing a global shift in both demographic and land use patterns. More than half of humanity now lives in urban areas, and with nearly all global population growth occurring in cities, that proportion is expected to reach two-thirds by 2050 (143). Much attention focuses on megacities of more than 10 million people, such as Mexico City and São Paulo, Cairo and Lagos, Karachi and Delhi, and Manila and Jakarta, but nearly half the world's urban dwellers live in a growing number of relatively small cities with fewer than 500,000 inhabitants and where population increase is faster (143).

Urbanization has emerged with different trajectories and in different ways, in different places. Europe and the Americas urbanized relatively early and have plateaued, whereas Asia and Africa continue to undergo rapid urbanization. Fast-growing cities of the Global South confront a range

of health and environmental challenges, including deficiencies in basic infrastructure (piped water, sewage, solid waste management, electricity, transportation, housing) and hazardous exposures (extremely poor air quality, noise, and unsafe roadways), which compounds problems of poverty, poor governance, and inadequate social services (117). In wealthy settings, especially in North America, Europe, and Australia, but increasingly in other regions, urban environmental health challenges reflect excessive automobile dependence, with associated urban sprawl, and resulting problems such as poor air quality, sedentary lifestyles, and injury risk (47). The burden is economically significant; the externalized costs of urban sprawl are about \$400 billion annually in the United States alone, owing to the increased cost of providing public services, the higher capital investment needed for infrastructure, and the costs of increased traffic congestion, crashes, and air pollution, a substantial portion of these being health costs (54). Finally, some problems are common to cities in both wealthy and poor nations; these include extreme social stratification, neighborhoods of concentrated poverty, insufficient green space, food deserts, and vulnerability to disasters.

The relationship between GEC and urbanization is bidirectional. Poorly designed cities contribute to GEC. Conversely, GEC imposes burdens on cities, and many of these burdens directly threaten the health and well-being of urban residents.

What is the impact of urbanization on GEC (and on NCD risk)? Economic activity in cities generates between 70% and 85% of global gross domestic product and about 75% of energy-related greenhouse gas (GHG) emissions (54, 142). Cities offer major opportunities to reduce per capita environmental impacts relative to rural areas, due to short travel distances, reduced per capita living space, and efficiencies in delivering goods and services, but poorly designed cities forfeit these potential advantages. Moreover, because cities source most of their energy and goods from outside the city limits, a full accounting of the impact of cities on GEC extends well beyond city boundaries (105). Several pathways are well documented, and they often feature contributions both to GEC and to NCDs.

First, the combination of automobile-dependent transportation systems and urban sprawl, and the concentration of industrial and domestic energy needs, means that large quantities of fossil fuels are burned in metropolitan areas. This practice contributes to climate change, as well as to regional air pollution.

Urban air quality is often poor, especially in cities in LMICs. According to the WHO (http://www.who.int/phe/health_topics/outdoorair/databases/cities/en/), more than 80% of people living in urban areas that monitor air pollution are exposed to air-quality levels above WHO guidelines, and 98% of cities in LMICs with more than 100,000 inhabitants do not meet WHO air-quality standards. A systematic review found that, globally, 25% of urban ambient PM air pollution comes from traffic, 15% from industrial activities, 20% from domestic fuel burning, 22% from unspecified sources of human origin, and 18% from natural dust and salt (71). In cities where indoor solid fuels are commonly used, exposure to ambient air pollution is compounded by household air pollution exposure, an important source of black carbon that is a powerful SLCP (85). The contributions of air pollution to NCDs have been discussed above.

Second, automobile-dependent transportation systems contribute both to GHG emissions and to sedentary lifestyles, as walking, cycling, and mass transit give way to private vehicular travel. A recent study of 14 cities in 10 countries identified 4 environmental attributes significantly associated with physical activity: net residential density, intersection density, public transport density, and number of parks. (In contrast with some other studies, land use mix and distance to transit did not show this association.) The difference in physical activity levels between the most and least activity-friendly neighborhoods ranged from 68 to 89 minutes per week (of a recommended total of 150 minutes per week) (116). A recent prospective study in Perth, Australia, confirmed the role

of street connectivity in predicting walking and also found associations for land use mix, nearby public transit stops, and a variety of local destinations (76). Sedentary lifestyles increase the risk of, and physical activity is protective against, a range of NCDs, including cardiovascular disease, some cancers, hypertension, obesity, depression and anxiety, osteoporosis, gall bladder disease, and others (81, 150). The links among automobile dependency, sedentary behaviors, and GHG emissions may be bidirectional, as increasing obesity makes walking more difficult and may be associated with higher transport-related GHG emissions (57).

Finally, several common features of urban life are indirectly linked to GEC but impact NCD risk. Noise is a common exposure in cities, caused by traffic, machinery, and other sources (which are also GEC contributors). Noise is a stressor that contributes to a range of NCDs, including hypertension, cardiovascular disease, anxiety, sleep disturbance, and hearing loss (111). City living may provide fewer opportunities for contact with nature and green space compared with rural living. Systematic reviews suggest that such salutogenic exposure reduces the risk of obesity (78), diabetes, hypertension, and hyperlipidemia (17), depression and anxiety (49), and cardiovascular mortality (50), although the quality and consistency of the evidence vary. Crowding is another common feature of urban life, especially in impoverished parts of cities. While crowding is difficult to define with precision and is context dependent (134), it is associated with similar cardiovascular and mental health effects as found with noise. Children may be especially susceptible to the effects of both noise and crowding (36, 129). [Urban density is associated with reduced transport-related GHG emissions (84), underlining the importance of evidence-based design that achieves the benefits of density while avoiding disbenefits such as noise and subjective crowding.] Inadequate housing is also a problem for many cities, especially those undergoing rapid growth and those in poor nations. Substantial portions of urban populations live in slums—more than half the population of cities such as Nairobi, Mumbai, and Mexico City, totaling an estimated 881 million people globally and rising (38). In such settings, the effects of inadequate housing—chronic stress, contaminated indoor air due to the use of solid fuels, exposure to temperature extremes, injury risks, and more—are amplified by failures in community infrastructure and health services. Respiratory disease and mental disorders are likely results, although few studies have examined the burden of NCDs in slums (38).

Conversely, GEC exerts impacts on cities (and on urban health, including NCD risk). First, droughts, floods, and other environmental disruptions propel rural-to-urban migration, a pattern documented across the Global South (9, 151). The resulting contribution to rapid urban growth intensifies such problems as air pollution, crowding, noise, and strained infrastructure and services. Second, environmental change may particularly threaten cities, through such pathways as heat [amplified by the urban heat island effect (104)], flood risk (60), water scarcity (60), and diminished air quality (102). These vulnerabilities may increase the risk of NCDs such as cardiopulmonary disease (from heat and reduced air quality) and mental illness (from heat and disasters) for people in urban areas. This risk is especially pertinent in poor cities such as those in the Global South, which lack the infrastructure needed for resilience (53), and particularly for people in informal settlements, because of their location (flood plains, steep slopes, etc.) and their lack of infrastructure (120).

There are complex social, economic, cultural, and physical determinants of NCDs in urban environments. Improving understanding of these risks is a priority for research (125). The challenge will be to provide housing and infrastructure that support health and livelihoods, particularly in rapidly growing smaller and midsized cities in LMICs, while avoiding the costly errors of some wealthy cities. Transforming existing cities so that they flourish at much lower levels of environmental impact represents a major policy challenge.

Food, Nutrition, and Agriculture

The relationships among GEC, diet, nutrition, and NCDs are complex and multidirectional. Like cities, food systems both contribute to environmental change and are vulnerable to its effects. Here we discuss three examples of interacting processes. First, the global dietary transition, featuring growing demand for both dietary animal products and processed foods, has direct health consequences, as well as environmental consequences that indirectly affect health. Second, environmental change affects agricultural productivity, threatening nutritional status in some regions, with implications for NCDs. Third, environmental change affects the nutritional content of some foods, with implications for NCDs.

The global dietary transition has been under way for several decades (107). It is marked by the consumption of energy-dense, less diverse, and increasingly processed foods, including animal products, oils and fats, refined carbohydrates, and sugar-sweetened beverages, and by behavioral changes such as increased snacking and increased eating outside the home. This transition is well demonstrated by evidence from LMICs (59), and, with increasing prosperity, population growth, and aging, it is expected that the NCD burden will continue to increase in the absence of strong preventive actions (26). This dietary pattern has direct implications for NCDs, promoting obesity, cardiovascular disease, diabetes, and some cancers (5, 15, 80, 106, 107). It also has environmental impacts that, in turn, may loop back to exert indirect impacts on human health. For example, diets heavy in meat (particularly ruminant meat) require substantially more land, water, and energy throughput to produce than plant-based diets (159). The FAO (39) has estimated that livestock production accounts for 18% of global GHG emissions. In Latin America, conversion of land to pasture for cattle is a major driver of deforestation (particularly in Brazil where it accounts for about 80% of forest loss); this land use change contributes to greenhouse gas emissions, which in turn contribute to NCD risk and to biodiversity loss (30).

A second food-related pathway from GEC to NCDs is the effect of climate change on agricultural productivity. Climate change reduces crop yields particularly at low and mid latitudes. Compared with a scenario without climate change, crop yields (wheat, rice, and maize) could decline by a median of 0.2% per decade for the rest of the century, even as demand increases by an estimated 14% per decade (109). Recent analysis suggests even larger losses, especially in temperate zones, due to the flourishing of insect pests with warmer weather (31). Climate change could therefore substantially increase the risk of undernutrition and stunting, particularly severe stunting (86). Whereas stunting has plateaued in Africa since 1990 at a prevalence of about 40%, evidence has shown a dramatic decrease in Asia from 49% in 1990 to 28% in 2010, suggesting that vulnerability may be highest in Africa (29). Stunting has serious implications, including impaired cognitive development and reduced economic prospects for those affected; these impacts persist in subsequent generations (146). Some evidence suggests that stunting increases the risk of obesity in later life, although this point remains controversial (140).

A recent modeling study of the effects of climate change on crop yield by 2050 suggested a net annual increase of about 529,000 premature deaths (95% CI averaged over all climate scenarios 314,000–736,000) compared with no climate change, mainly as a result of increased NCD mortality (130). Reductions in fruit and vegetable consumption were the major contributor to increased mortality, and most climate-related deaths were projected to occur in South and East Asia. Decreased caloric intake was projected to increase deaths related to undernutrition, but these were approximately balanced by reduced deaths from overweight and obesity.

A third food-related pathway from GEC to NCDs is the effect of environmental change on the nutritional value of crops. A meta-analysis of data on 130 varieties of plants found that elevated carbon dioxide (CO₂) levels reduced the overall concentration of 25 important minerals in plants,

including calcium, potassium, zinc, and iron, by 8% on average (87). An estimated 175 million (95% CI 162–186) more people, mainly in Africa and South Asia, could be placed at new risk of zinc deficiency by 2050 as a result of elevated CO₂ levels (127). Strong evidence indicates that elevated CO₂ levels also reduce protein and increase carbohydrate levels particularly in C3 crops. One study found that elevated CO₂ was associated with a ~6% decrease in protein in wheat grains and an ~8% decrease in rice grains but demonstrated no significant effect in soybeans (93). The health implications of these changes are under investigation and are likely to depend on the nutritional profile of the affected populations. A study in the United States showed that the substitution of dietary carbohydrate for dietary protein increased the risk of hypertension, lipid disorders, and 10-year coronary heart disease risk (6). In low-income countries, if protein intake is low, there may be additional substantial impacts on public health.

Oil palm cultivation exemplifies the complex relationships among GEC, diet, nutrition, and NCDs. Production is increasing as a result of demand for biofuels in Europe and food in India, Indonesia, and China. Oil palm plantations are consistently less biodiverse than primary forests; they contain about 50% of the vertebrate species found in primary forests and show reduced species richness compared with secondary forests (118). To clear tropical forests in Indonesia for palm oil (as well as timber) production, fire is commonly used; the resulting smoke contains fine PM that contributes to a global burden of more than 250,000 deaths annually (70). Clearing tropical forests to produce palm oil is also deleterious in terms of climate change. And dietary palm oil contains highly saturated fatty acids, which have been proposed to be a risk factor for heart disease and other NCDs (10). A systematic review and meta-analysis (41) of short-term dietary intervention studies suggests, however, that both favorable and unfavorable changes in coronary heart disease/cardiovascular disease risk markers occurred when palm oil was substituted for primary dietary fats and that additional evidence is needed to inform firm conclusions.

Two additional pathways from GEC to NCDs are the effects of fisheries depletion and pollinator loss. These are discussed below in the section on biodiversity.

Persistent Chemicals in the Environment

Another feature of the Anthropocene has been widespread contamination of the global environment by chemicals (32, 79). Although many features of this contamination, from its geographic extent and levels in various biota to its full impacts on human health, are not fully understood, the links with NCDs are coming into sharper focus. Two examples are illustrative: endocrine disruptors and metals.

Endocrine disruptors include diverse classes of chemicals that affect endocrine pathways, by either blocking or activating receptors in sex hormone, thyroid, or other pathways. Many endocrine disruptors are synthetic organic chemicals such as polychlorinated biphenyls, bisphenols [e.g., bisphenol A (BPA)], organochlorine pesticides, brominated flame retardants, and perfluorinated substances (perfluorooctanoic acid and perfluorooctane sulfonate). Many such chemicals persist in the environment over long periods of time and are therefore called persistent organic pollutants (POPs). While there is considerable variation among these chemicals, three points are relevant. First, they are widely distributed in global ecosystems. Second, human exposure is widespread. Third, evidence suggests numerous associations with NCD risk.

With regard to environmental distribution, POPs have been found in nearly every ecosystem assessed, including in Arctic and Antarctic regions remote from any sites of manufacturing or use (149). Once these substances enter the environment, global transport is widespread, in both biological and physical media. Fat-soluble chemicals are concentrated (biomagnified) as they move from lower to higher trophic levels of food webs (89). A recently recognized contributor is the

presence of plastics in various ecosystems, from oceans to lakes to coastlines (8). Plastic fragments and microplastics are now commonly found in the gastrointestinal tracts of many marine birds and fish (114). Component monomers such as propylene and ethylene are relatively nontoxic, but associated plasticizers such as BPA and other adsorbed organic chemicals may have substantial biological activity (113). Of note, concomitant global trends affect the distribution of POPs. For example, climate change, operating through such processes as altered ocean currents, changes in temperature, and erosion of soils following severe weather events, may both liberate more pollutants and accelerate their breakdown (94, 148).

When POPs are widely distributed through ecosystems, human exposure is unavoidable. Measurement of population tissue levels of POPs has revealed nearly ubiquitous body burdens (22, 108). While levels of various contaminants vary by location, age, dietary patterns, and other factors, few humans escape exposure to, and accumulation of, at least some POPs.

Evidence suggests that these exposures play a role in several NCDs through both epigenetic and nonepigenetic mechanisms (65). POPs exposure has been associated with metabolic conditions such as adiposity, insulin resistance, and dyslipidemias, although evidence is inconsistent and methodologic shortcomings are common in published studies (67, 82, 137). POPs exposure has also been associated with the risk of some cancers, especially non-Hodgkin's lymphoma (45) and hormone-responsive cancers such as those of the breast, ovaries, and prostate; there is considerable animal evidence, but human epidemiologic evidence is less definitive (58). POPs may also increase the risks of thyroid disease, neurobehavioral disorders, and reproductive dysfunction (58). For each of these outcomes, the role of ambient environmental contamination relative to local exposures, such as in the workplace, remains to be defined.

Metals represent a second example of widespread chemical contamination. Human exposure to metals is typically a local phenomenon; examples include the child who consumes lead from aging paint in substandard housing or the worker exposed to mercury in artisanal gold mining. However, more dispersed mobilization of metals has occurred, with impacts that are regional if not global. For example, in China, industrial activity, mining, and the use of inadequately treated wastewater for irrigation have contaminated soil across large portions of the country with lead, cadmium, chromium, and other metals (74). Approximately 250,000 km² of farmland—an area equivalent to the arable farmland of Mexico—is contaminated, and in 2017 Chinese officials reportedly designated 35,000 km² of farmland as too polluted to permit any agricultural use (34). As another example, mercury emissions have been a feature of coal combustion since the dawn of the Industrial Revolution. Mercury contamination is concentrated near point sources such as power plants, smelters, and cement, iron, and steel plants; however, this is also a global phenomenon, as the mercury in air emissions travels great distances, even intercontinentally, before settling back to earth (121). The metals, collectively, contribute to many NCDs, including neurobehavioral abnormalities (lead, mercury), cardiovascular disease (lead, cadmium), renal disease (lead, cadmium), and some cancers (arsenic, chromium) (97).

Biodiversity Loss

Biodiversity loss has accelerated dramatically during the Anthropocene (96). Although this loss has numerous potential impacts on human health, including on NCDs (12), two examples are especially illustrative: the effects of pollinator loss and of fisheries depletion.

Pollination by insects is an important form of reproduction for more than 35% of the annual global food production by volume. At least 87 major types of food crops, and up to 40% of the world's supply of some micronutrients, such as vitamin A, depend on pollination by insects (75). Pollinators are declining in many parts of the world probably for a combination of reasons,

including habitat loss, pesticide use, and parasitic infestation. Pollinator loss can reduce the amount of fruits, vegetables, and nuts and seeds in the diet and lead to vitamin A and folate deficiencies. A recent analysis projects that a 50% loss of pollination would cause an additional ~0.7 million deaths worldwide, mostly as a result of increased ischemic heart disease and stroke due to reduced fruit and vegetable consumption (128).

Fisheries depletion has emerged as a global problem, with about 90% of fisheries at or beyond maximum sustainable levels of exploitation (40). Climate change will intensify this problem in coming decades (28). For many human populations, fish is a leading dietary source of protein, of micronutrients (often in highly bioavailable forms), and of omega-3 fatty acids (mainly from oily fish). Dietary omega-3 fatty acids may reduce ischemic heart disease risk, although research findings remain inconsistent (110). In addition, through gene regulation, anti-inflammatory effects, or other mechanisms, omega-3 fatty acids may play a role in preventing and/or treating other NCDs such as cancer (83) and arthritis (122). However, reductions in fish stocks could limit these potential benefits. For example, the United Kingdom is unable to meet healthy diet guidelines for its population with its domestic catch, and intake fell to only 19% of the recommended level in 2012 (139). One study projected that more than 10% of the global population could face micronutrient and fatty-acid deficiencies due to fish declines over the coming decades, especially in LMICs near the equator (55). Aquaculture is, to some extent, replacing wild fish catches, but sustainable aquaculture must address a range of challenges including disease and chemical contamination in farmed fish, unsustainable feedstocks, water pollution near fish farms, and genetic contamination of wild fish (20).

CONCLUSIONS

The Importance of Systems Understanding

Although we have presented five aspects of GEC—energy, air pollution, and climate change; urbanization; food, nutrition, and agriculture; chemical contamination; and biodiversity loss—in separate discussions, they are by no means distinct. A complete understanding of their effects on NCDs requires consideration of their interconnections (**Figure 4**). For example, energy, pollution, and biodiversity are linked. Heavy reliance on petroleum as an energy source results

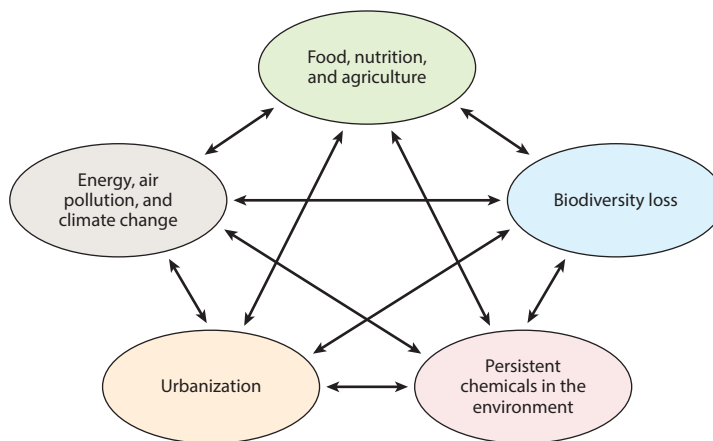


Figure 4

Interactions among various global environmental changes.

in contamination of marine environments by petroleum-derived pollutants, which are toxic to developing fish and which compound threats to some fisheries (23). Similarly, urbanization is linked with climate change in many ways. Cities occupy only 2% of the world's landmass, but as noted above they account for the lion's share of global energy use and global CO₂ emissions. The form of urbanization can, in turn, determine a city's experience of climate change. For example, evidence from the United States suggests that sprawling metropolitan areas are at substantially higher risk of extreme heat events than are more compact urban areas (132). Ample, well-designed urban green space and blue space can not only promote health directly, but also help reduce the heat island effect (104). A third example is the link between urbanization and food systems. Rural-to-urban migration in LMICs is associated with rapid changes in dietary patterns, including the adoption of Western-style processed foods; one study found the obesity prevalence in Accra, Ghana, to be 4–5 times higher, and the diabetes prevalence 2–3 times higher, than in rural areas of Ghana (2). The various pathways through which GECs affect NCD risk are interrelated in complex ways, and solutions must take account of these complexities.

Planetary Health: Reducing NCD Risk and Achieving Environmental Sustainability

The preceding discussion has offered an analysis of links between planetary changes and NCD risk. In each of the five areas discussed, strategies are available that simultaneously promote health and well-being and environmental sustainability (61, 79, 154). Clean, renewable energy sources can replace fossil fuels. The built environment, from the scale of buildings to the scale of entire metropolitan areas, including both stationary assets such as buildings, parks, and utilities, and the mobile components of transportation systems, can be designed in ways that optimize human health as well as environmental performance. The food system, from farm to table, can be shifted toward more sustainable food production, consumption of healthier foods, and less waste. Green chemistry aims to design molecules that are less persistent, less toxic, and as functional as many legacy chemicals, if not more so. And management strategies for land, water, and other ecosystem components can optimize biodiversity conservation, for example through forest conservation, as well as reduce air pollution from landscape fires. In light of the growing evidence that GECs can affect NCD risk through a range of pathways, it is imperative that the evidence base linking the two be strengthened and greater investment be devoted to policies that can reduce both environmental change and NCD risk.

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LITERATURE CITED

1. Adams ZW, Sumner JA, Danielson CK, McCauley JL, Resnick HS, et al. 2014. Prevalence and predictors of PTSD and depression among adolescent victims of the Spring 2011 tornado outbreak. *J. Child Psychol. Psychiatry* 55:1047–55
2. Agyemang C, Meeks K, Beune E, Owusu-Dabo E, Mockenhaupt FP, et al. 2016. Obesity and type 2 diabetes in sub-Saharan Africans - Is the burden in today's Africa similar to African migrants in Europe? The RODAM study. *BMC Med.* 14:166
3. Allen L. 2017. Non-communicable disease funding. *Lancet Diabetes Endocrinol.* 5:92

4. Allen LN, Feigl AB. 2017. What's in a name? A call to reframe non-communicable diseases. *Lancet Global Health* 5:e129–30
5. Anand SS, Hawkes C, de Souza RJ, Mente A, Dehghan M, et al. 2015. Food consumption and its impact on cardiovascular disease: importance of solutions focused on the globalized food system: a report from the workshop convened by the World Heart Federation. *J. Am. Coll. Cardiol.* 66:1590–614
6. Appel LJ, Sacks FM, Carey VJ, Obarzanek E, Swain JF, et al. 2005. Effects of protein, monounsaturated fat, and carbohydrate intake on blood pressure and serum lipids: results of the OmniHeart randomized trial. *JAMA* 294:2455–64
7. Atkinson RW, Butland BK, Dimitroulopoulou C, Heal MR, Stedman JR, et al. 2016. Long-term exposure to ambient ozone and mortality: a quantitative systematic review and meta-analysis of evidence from cohort studies. *BMJ Open* 6:e009493
8. Avio CG, Gorbi S, Regoli F. 2017. Plastics and microplastics in the oceans: from emerging pollutants to emerged threat. *Mar. Environ. Res.* 128:2–11
9. Barrios S, Bertinelli L, Strobl E. 2006. Climatic change and rural–urban migration: the case of sub-Saharan Africa. *J. Urban Econ.* 60:357–71
10. Basu S, Babiarz KS, Ebrahim S, Vellakkal S, Stuckler D, Goldhaber-Fiebert JD. 2013. Palm oil taxes and cardiovascular disease mortality in India: economic-epidemiologic model. *BMJ* 347:f6048
11. Battilani P, Toscano P, Van der Fels-Klerx HJ, Moretti A, Camardo Leggieri M, et al. 2016. Aflatoxin B₁ contamination in maize in Europe increases due to climate change. *Sci. Rep.* 6:24328
12. Bernstein AS. 2014. Biological diversity and public health. *Annu. Rev. Public Health* 35:153–67
13. Biswas A, Oh PI, Faulkner GE, Bajaj RR, Silver MA, et al. 2015. Sedentary time and its association with risk for disease incidence, mortality, and hospitalization in adults: a systematic review and meta-analysis. *Ann. Intern. Med.* 162:123–32
14. Bloom DE, Cafiero ET, Jané-Llopis E, Abrahams-Gessel S, Bloom LR, et al. 2011. *The Global Economic Burden of Non-Communicable Diseases*. Geneva: World Econ. Forum, Harvard Sch. Public Health
15. Bouvard V, Loomis D, Guyton KZ, Grosse Y, El Ghissassi F, et al. 2015. Carcinogenicity of consumption of red and processed meat. *Lancet Oncol.* 16:1599–600
16. Bowe B, Xie Y, Li T, Yan Y, Xian H, Al-Aly Z. 2018. The 2016 global and national burden of diabetes mellitus attributable to PM_{2.5} air pollution. *Lancet Planet. Health* 2:e301–12
17. Brown SC, Lombard J, Wang K, Byrne MM, Toro M, et al. 2016. Neighborhood greenness and chronic health conditions in Medicare beneficiaries. *Am. J. Prev. Med.* 51:78–89
18. Burnett R, Chen H, Szyszkowicz M, Fann N, Hubbell B, et al. 2018. Global estimates of mortality associated with long-term exposure to outdoor fine particulate matter. *PNAS* 115(38):9592–97
19. Bygbjerg IC. 2012. Double burden of noncommunicable and infectious diseases in developing countries. *Science* 337:1499–501
20. Cao L, Wang W, Yang Y, Yang C, Yuan Z, et al. 2007. Environmental impact of aquaculture and countermeasures to aquaculture pollution in China. *Environ. Sci. Pollut. Res. Int.* 14:452–62
21. Cappuccio FP, Cooper D, D'Elia L, Strazzullo P, Miller MA. 2011. Sleep duration predicts cardiovascular outcomes: a systematic review and meta-analysis of prospective studies. *Eur. Heart J.* 32:1484–92
22. CDC (Cent. Dis. Control Prev.). 2018. *Fourth national report on human exposure to environmental chemicals. Updated tables*. Rep., CDC, Atlanta. <https://www.cdc.gov/exposurereport/index.html>
23. Cherr GN, Fairbairn E, Whitehead A. 2017. Impacts of petroleum-derived pollutants on fish development. *Annu. Rev. Anim. Biosci.* 5:185–203
24. Clayton S, Manning C, Krygsman K, Speiser M. 2017. *Mental Health and Our Changing Climate: Impacts, Implications, and Guidance*. Washington, DC: Am. Psychol. Assoc., EcoAmerica
25. Clifford A, Lang L, Chen R, Anstey KJ, Seaton A. 2016. Exposure to air pollution and cognitive functioning across the life course—a systematic literature review. *Environ. Res.* 147:383–98
26. Clonan A, Roberts KE, Holdsworth M. 2016. Socioeconomic and demographic drivers of red and processed meat consumption: implications for health and environmental sustainability. *Proc. Nutr. Soc.* 75:367–73

27. Cohen AJ, Brauer M, Burnett R, Anderson HR, Frostad J, et al. 2017. Estimates and 25-year trends of the global burden of disease attributable to ambient air pollution: an analysis of data from the Global Burden of Diseases Study 2015. *Lancet* 389:1907–18
28. Comte L, Olden JD. 2017. Climatic vulnerability of the world's freshwater and marine fishes. *Nat. Clim. Change* 7:718–22
29. de Onis M, Blössner M, Borghi E. 2012. Prevalence and trends of stunting among pre-school children, 1990–2020. *Public Health Nutr.* 15:142–48
30. De Sy V, Herold M, Achard F, Beuchle R, Clevers JGPW, et al. 2015. Land use patterns and related carbon losses following deforestation in South America. *Environ. Res. Lett.* 10:124004
31. Deutsch CA, Tewksbury JJ, Tigchelaar M, Battisti DS, Merrill SC, et al. 2018. Increase in crop losses to insect pests in a warming climate. *Science* 361:916–19
32. Diamond ML, de Wit CA, Molander S, Scheringer M, Backhaus T, et al. 2015. Exploring the planetary boundary for chemical pollution. *Environ. Int.* 78:8–15
33. Dimakakou E, Johnston HJ, Streftaris G, Cherrie JW. 2018. Exposure to environmental and occupational particulate air pollution as a potential contributor to neurodegeneration and diabetes: a systematic review of epidemiological research. *Int. J. Environ. Res. Public Health* 15:1704
34. Economist. 2017. The most neglected threat to public health in China is toxic soil. *Economist*, June 8. <https://www.economist.com/briefing/2017/06/08/the-most-neglected-threat-to-public-health-in-china-is-toxic-soil>
35. Eisenman D, McCaffrey S, Donatello I, Marshal G. 2015. An ecosystems and vulnerable populations perspective on solastalgia and psychological distress after a wildfire. *EcoHealth* 12:602–10
36. Evans GW, Lercher P, Meis M, Ising H, Kofler WW. 2001. Community noise exposure and stress in children. *J. Acoust. Soc. Am.* 109:1023–27
37. Eze IC, Hemkens LG, Bucher HC, Hoffman B, Schindler C, et al. 2015. Association between ambient air pollution and diabetes mellitus in Europe and North America: systematic review and meta-analysis. *Environ. Health Perspect.* 123(5):381–89
38. Ezeh A, Oyebo O, Satterthwaite D, Chen Y-F, Ndugwa R, et al. 2017. The history, geography, and sociology of slums and the health problems of people who live in slums. *Lancet* 389:547–58
39. FAO (Food Agric. Organ.). 2006. *Livestock's Long Shadow: Environmental Issues and Options*. Rome: FAO
40. FAO (Food Agric. Organ.). 2016. *The State of World Fisheries and Aquaculture*. Rome: FAO
41. Fattore E, Bosetti C, Brighenti F, Agostoni C, Fattore G. 2014. Palm oil and blood lipid-related markers of cardiovascular disease: a systematic review and meta-analysis of dietary intervention trials. *Am. J. Clin. Nutr.* 99:1331–50
42. Fernandez A, Black J, Jones M, Wilson L, Salvador-Carulla L, et al. 2015. Flooding and mental health: a systematic mapping review. *PLOS ONE* 10:e0119929
43. Field CB, Barros VR, Dokken DJ, Mach KJ, Mastrandrea MD, eds. 2014. *Climate Change 2014: Impacts, Adaptation and Vulnerability. Part A: Global and Sectoral Aspects. Contribution of Working Group II to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change*. Cambridge, UK/New York: Cambridge Univ. Press
44. Folino F, Buja G, Zanutto G, Marras E, Allocca G, et al. 2017. Association between air pollution and ventricular arrhythmias in high-risk patients (ARIA study): a multicentre longitudinal study. *Lancet Planet. Health* 1:e58–64
45. Freeman MD, Kohles SS. 2012. Plasma levels of polychlorinated biphenyls, non-Hodgkin lymphoma, and causation. *J. Environ. Public Health* 2012:258981
46. Friel S, Bowen K, Campbell-Lendrum D, Frumkin H, McMichael AJ, Rasanathan K. 2011. Climate change, noncommunicable diseases, and development: the relationships and common policy opportunities. *Annu. Rev. Public Health* 32:133–47
47. Frumkin H, Frank LD, Jackson RJ. 2004. *Urban Sprawl and Public Health: Designing, Planning, and Building for Healthy Communities*. Washington, DC: Island Press
48. Galea S, Brewin CR, Gruber M, Jones RT, King DW, et al. 2007. Exposure to hurricane-related stressors and mental illness after Hurricane Katrina. *Arch. Gen. Psychiatry* 64:1427–34

49. Gascon M, Triguero-Mas M, Martínez D, Dadvand P, Fors J, et al. 2015. Mental health benefits of long-term exposure to residential green and blue spaces: a systematic review. *Int. J. Environ. Res. Public Health* 12:4354–79
50. Gascon M, Triguero-Mas M, Martínez D, Dadvand P, Rojas-Rueda D, et al. 2016. Residential green spaces and mortality: a systematic review. *Environ. Int.* 86:60–67
51. Gasparrini A, Guo Y, Sera F, Vicedo-Cabrera AM, Huber V, et al. 2017. Projections of temperature-related excess mortality under climate change scenarios. *Lancet Planet. Health* 1:e360–67
52. GBD Risk Factors Collab. 2016. Global, regional, and national comparative risk assessment of 79 behavioural, environmental and occupational, and metabolic risks or clusters of risks, 1990–2015: a systematic analysis for the Global Burden of Disease Study 2015. *Lancet* 388:1659–724
53. Giugni M, Simonis I, Bucchignani E, Capuano P, De Paola F, et al. 2015. The impacts of climate change on African cities. In *Urban Vulnerability and Climate Change in Africa: A Multidisciplinary Approach*, ed. S Pauleit, A Coly, S Fohlmeister, P Gasparini, G Jørgensen, et al., pp. 37–75. Cham, Switz.: Springer Int.
54. Global Comm. Econ. Clim. 2015. *Seizing the global opportunity: partnerships for better growth and a better climate. The 2015 New Climate Economy report*. Rep., New Clim. Econ., New York/London. <http://newclimateeconomy.report/2015/>
55. Golden CD, Allison EH, Cheung WWL, Dey MM, Halpern BS, et al. 2016. Nutrition: fall in fish catch threatens human health. *Nature* 534:317–20
56. Goldmann E, Galea S. 2014. Mental health consequences of disasters. *Annu. Rev. Public Health* 35:169–83
57. Goodman A, Brand C, Ogilvie D. 2012. Associations of health, physical activity and weight status with motorised travel and transport carbon dioxide emissions: a cross-sectional, observational study. *Environ. Health* 11:52
58. Gore AC, Chappell VA, Fenton SE, Flaws JA, Nadal A, et al. 2015. EDC-2: The Endocrine Society's second scientific statement on endocrine-disrupting chemicals. *Endocr. Rev.* 36:E1–150
59. Green R, Milner J, Joy EJ, Agrawal S, Dangour AD. 2016. Dietary patterns in India: a systematic review. *Br. J. Nutr.* 116:142–48
60. Güneralp B, Güneralp İ, Liu Y. 2015. Changing global patterns of urban exposure to flood and drought hazards. *Glob. Environ. Change* 31:217–25
61. Haines A, McMichael AJ, Smith KR, Roberts I, Woodcock J, et al. 2009. Public health benefits of strategies to reduce greenhouse-gas emissions: overview and implications for policy makers. *Lancet* 374:2104–14
62. Hart CR, Berry HL, Tonna AM. 2011. Improving the mental health of rural New South Wales communities facing drought and other adversities. *Aust. J. Rural Health* 19:231–38
63. He D, Wu S, Zhao H, Qiu H, Fu Y, et al. 2017. Association between particulate matter 2.5 and diabetes mellitus: a meta-analysis of cohort studies. *J. Diabetes Investig.* 8:687–96
64. Hoek G, Krishnan RM, Beelen R, Peters A, Ostro B, et al. 2013. Long-term air pollution exposure and cardio-respiratory mortality: a review. *Environ. Health* 12:43
65. Hou L, Zhang X, Wang D, Baccarelli A. 2012. Environmental chemical exposures and human epigenetics. *Int. J. Epidemiol.* 41:79–105
66. IEA (Int. Energy Agency). 2016. *World Energy Outlook Special Report 2016: Energy and Air Pollution*. Paris: IEA
67. Jaacks LM, Staimez LR. 2015. Association of persistent organic pollutants and non-persistent pesticides with diabetes and diabetes-related health outcomes in Asia: a systematic review. *Environ. Int.* 76:57–70
68. Janssen NAH, Gerlofs-Nijland ME, Lanki T, Salonen RO, Cassee F, et al. 2012. *Health Effects of Black Carbon*. Copenhagen: World Health Organ.
69. Ji M, Cohan DS, Bell ML. 2011. Meta-analysis of the association between short-term exposure to ambient ozone and respiratory hospital admissions. *Environ. Res. Lett.* 6:024006
70. Johnston FH, Henderson SB, Chen Y, Randerson JT, Marlier M, et al. 2012. Estimated global mortality attributable to smoke from landscape fires. *Environ. Health Perspect.* 120:695–701
71. Karagulian F, Belis CA, Dora CFC, Prüss-Ustün AM, Bonjour S, et al. 2015. Contributions to cities' ambient particulate matter (PM): a systematic review of local source contributions at global level. *Atmos. Environ.* 120:475–83

72. Kessler RC, Galea S, Gruber MJ, Sampson NA, Ursano RJ, Wessely S. 2008. Trends in mental illness and suicidality after Hurricane Katrina. *Mol. Psychiatry* 13:374–84
73. Khan AE, Scheelbeek PFD, Shilpi AB, Chan Q, Mojumder SK, et al. 2014. Salinity in drinking water and the risk of (pre)eclampsia and gestational hypertension in coastal Bangladesh: a case-control study. *PLOS ONE* 9:e108715
74. Khan S, Cao Q, Zheng YM, Huang YZ, Zhu YG. 2008. Health risks of heavy metals in contaminated soils and food crops irrigated with wastewater in Beijing, China. *Environ. Poll.* 152:686–92
75. Klein A-M, Vaissière BE, Cane JH, Steffan-Dewenter I, Cunningham SA, et al. 2007. Importance of pollinators in changing landscapes for world crops. *Proc. R. Soc. B* 274:303–13
76. Knuiman MW, Christian HE, Divitini ML, Foster SA, Bull FC, et al. 2014. A longitudinal analysis of the influence of the neighborhood built environment on walking for transportation: the RESIDE study. *Am. J. Epidemiol.* 180:453–61
77. Koplit SN, Jacob DJ, Sulprizio MP, Myllyvirta L, Reid C. 2017. Burden of disease from rising coal-fired power plant emissions in Southeast Asia. *Environ. Sci. Technol.* 51:1467–76
78. Lachowycz K, Jones AP. 2011. Greenspace and obesity: a systematic review of the evidence. *Obes. Rev.* 12:e183–89
79. Landrigan PJ, Fuller R, Acosta NJR, Adeyi O, Arnold R, et al. 2017. The Lancet Commission on pollution and health. *Lancet* 391:10119
80. Larsson SC, Orsini N. 2014. Red meat and processed meat consumption and all-cause mortality: a meta-analysis. *Am. J. Epidemiol.* 179:282–89
81. Lear SA, Hu W, Rangarajan S, Gasevic D, Leong D, et al. 2017. The effect of physical activity on mortality and cardiovascular disease in 130 000 people from 17 high-income, middle-income, and low-income countries: the PURE study. *Lancet* 390:p2643–54
82. Lee DH, Porta M, Jacobs DR Jr., Vandenberg LN. 2014. Chlorinated persistent organic pollutants, obesity, and type 2 diabetes. *Endocr. Rev.* 35:557–601
83. Lee JY, Sim T-B, Lee J-E, Na H-K. 2017. Chemopreventive and chemotherapeutic effects of fish oil derived omega-3 polyunsaturated fatty acids on colon carcinogenesis. *Clin. Nutr. Res.* 6:147–60
84. Lee S, Lee B. 2014. The influence of urban form on GHG emissions in the U.S. household sector. *Energy Policy* 68:534–49
85. Leung DY. 2015. Outdoor-indoor air pollution in urban environment: challenges and opportunity. *Front. Environ. Sci.* 2:69
86. Lloyd SJ, Kovats RS, Chalabi Z. 2011. Climate change, crop yields, and undernutrition: development of a model to quantify the impact of climate scenarios on child undernutrition. *Environ. Health Perspect.* 119:1817–23
87. Loladze I. 2014. Hidden shift of the ionome of plants exposed to elevated CO₂ depletes minerals at the base of human nutrition. *eLife* 3:e02245
88. Lunyera J, Mohottige D, von Isenburg M, Jeuland M, Patel UD, Stanifer JW. 2016. CKD of uncertain etiology: a systematic review. *Clin. J. Am. Soc. Nephrol.* 11:379–85
89. Mackay D, Fraser A. 2000. Bioaccumulation of persistent organic chemicals: mechanisms and models. *Environ. Poll.* 110:375–91
90. Malley CS, Henze DK, Kuylenstierna JCI, Vallack HW, Davila Y, et al. 2017. Updated global estimates of respiratory mortality in adults ≥ 30 years of age attributable to long-term ozone exposure. *Environ. Health Perspect.* 125:087021
91. Mammen G, Faulkner G. 2013. Physical activity and the prevention of depression: a systematic review of prospective studies. *Am. J. Prev. Med.* 45:649–57
92. McMichael AJ. 2013. Globalization, climate change, and human health. *New Engl. J. Med.* 368:1335–43
93. Myers SS, Zanoibetti A, Kloog I, Huybers P, Leakey ADB, et al. 2014. Increasing CO₂ threatens human nutrition. *Nature* 510:139–42
94. Nadal M, Marquès M, Mari M, Domingo JL. 2015. Climate change and environmental concentrations of POPs: a review. *Environ. Res.* 143:177–85

95. Naghavi M, Abajobir AA, Abbafati C, Abbas KM, Abd-Allah F, et al. 2017. Global, regional, and national age-sex specific mortality for 264 causes of death, 1980–2016: a systematic analysis for the Global Burden of Disease Study 2016. *Lancet* 390:1151–210
96. Newbold T, Hudson LN, Arnell AP, Contu S, De Palma A, et al. 2016. Has land use pushed terrestrial biodiversity beyond the planetary boundary? A global assessment. *Science* 353:288–91
97. Nordberg GF, Fowler BA, Nordberg M, eds. 2015. *Handbook on the Toxicology of Metals*. San Diego: Academic
98. O'Brien LV, Berry HL, Coleman C, Hanigan IC. 2014. Drought as a mental health exposure. *Environ. Res.* 131:181–87
99. Obradovich N, Fowler JH. 2017. Climate change may alter human physical activity patterns. *Nat. Hum. Behav.* 1:0097
100. Obradovich N, Migliorini R, Mednick SC, Fowler JH. 2017. Nighttime temperature and human sleep loss in a changing climate. *Sci. Adv.* 3:e1601555
101. Omran AR. 1971. The epidemiologic transition: a theory of the epidemiology of population change. *Milbank Mem. Fund. Q.* 49:509–38
102. Orru H, Ebi KL, Forsberg B. 2017. The interplay of climate change and air pollution on health. *Curr. Environ. Health Rep.* 4:504–13
103. Pearce N, Ebrahim S, McKee M, Lamprey P, Barreto ML, et al. 2015. Global prevention and control of NCDs: limitations of the standard approach. *J. Public Health Policy* 36:408–25
104. Phelan PE, Kaloush K, Miner M, Golden J, Phelan B, et al. 2015. Urban heat island: mechanisms, implications, and possible remedies. *Annu. Rev. Environ. Resour.* 40:285–307
105. Pincetl S. 2017. Cities in the age of the Anthropocene: climate change agents and the potential for mitigation. *Anthropocene* 20:74–82
106. Popkin BM. 2015. Nutrition transition and the global diabetes epidemic. *Curr. Diab. Rep.* 15:64
107. Popkin BM, Adair LS, Ng SW. 2012. Global nutrition transition: the pandemic of obesity in developing countries. *Nutr. Rev.* 70:3–21
108. Porta M, Puigdomènech E, Ballester F, Selva J, Ribas-Fitó N, et al. 2008. Monitoring concentrations of persistent organic pollutants in the general population: the international experience. *Environ. Int.* 34:546–61
109. Porter JR, Xie L, Challinor AJ, Cochrane K, Howden M, et al. 2014. Food security and food production systems. See Ref. 43, pp. 485–533
110. Rangel-Huerta OD, Gil A. 2017. Omega 3 fatty acids in cardiovascular disease risk factors: an updated systematic review of randomised clinical trials. *Clin. Nutr.* 37:72–77
111. Recio A, Linares C, Banegas JR, Díaz J. 2016. Road traffic noise effects on cardiovascular, respiratory, and metabolic health: an integrative model of biological mechanisms. *Environ. Res.* 146:359–70
112. Rigby M. 2017. Renaming non-communicable diseases. *Lancet Glob. Health* 5:e653
113. Rochman CM, Hoh E, Hentschel BT, Kaye S. 2013. Long-term field measurement of sorption of organic contaminants to five types of plastic pellets: implications for plastic marine debris. *Environ. Sci. Technol.* 47:1646–54
114. Rochman CM, Tahir A, Williams SL, Baxa DV, Lam R, et al. 2015. Anthropogenic debris in seafood: plastic debris and fibers from textiles in fish and bivalves sold for human consumption. *Sci. Rep.* 5:14340
115. Ryan B, Franklin RC, Burkle FM Jr., Aitken P, Smith E, et al. 2015. Identifying and describing the impact of cyclone, storm and flood related disasters on treatment management, care and exacerbations of non-communicable diseases and the implications for public health. *PLOS Curr. Disast.* 7. <https://doi.org/10.1371/currents.dis.62e9286d152de04799644dcca47d9288>
116. Sallis JF, Cerin E, Conway TL, Adams MA, Frank LD, et al. 2016. Physical activity in relation to urban environments in 14 cities worldwide: a cross-sectional study. *Lancet* 387:2207–17
117. Satterthwaite D. 2011. Editorial: Why is urban health so poor even in many successful cities? *Environ. Urban.* 23:5–11
118. Savilaakso S, Garcia C, Garcia-Ulloa J, Ghazoul J, Groom M, et al. 2014. Systematic review of effects on biodiversity from oil palm production. *Environ. Evidence* 3:4

119. Scheelbeek PFD, Chowdhury MAH, Haines A, Alam DS, Hoque MA, et al. 2017. Drinking water salinity and raised blood pressure: evidence from a cohort study in coastal Bangladesh. *Environ. Health Perspect.* 125:057007
120. Scovronick N, Lloyd SJ, Kovats RS. 2015. Climate and health in informal urban settlements. *Environ. Urban.* 27:657–78
121. Selin NE. 2009. Global biogeochemical cycling of mercury: a review. *Annu. Rev. Environ. Resour.* 34:43–63
122. Senftleber NK, Nielsen SM, Andersen JR, Bliddal H, Tarp S, et al. 2017. Marine oil supplements for arthritis pain: a systematic review and meta-analysis of randomized trials. *Nutrients* 9:42
123. Shah AS, Langrish JP, Nair H, McAllister DA, Hunter AL, et al. 2013. Global association of air pollution and heart failure: a systematic review and meta-analysis. *Lancet* 382:1039–48
124. Shoemaker JK, Schrag DP, Molina MJ, Ramanathan V. 2013. What role for short-lived climate pollutants in mitigation policy? *Science* 342:1323–24
125. Smit W, Hancock T, Kumaresen J, Santos-Burgoa C, Sánchez-Kobashi Meneses R, Friel S. 2011. Toward a research and action agenda on urban planning/design and health equity in cities in low and middle-income countries. *J. Urban Health* 88:875–85
126. Smith KR, Woodward A, Campbell-Lendrum D, Chadee DD, Honda Y, et al. 2014. Human health: impacts, adaptation, and co-benefits. See Ref. 43, pp. 709–54
127. Smith MR, Myers SS. 2018. Impact of anthropogenic CO₂ emissions on global human nutrition. *Nature Climate Change* 8:834–39
128. Smith MR, Singh GM, Mozaffarian D, Myers SS. 2015. Effects of decreases of animal pollinators on human nutrition and global health: a modelling analysis. *Lancet* 386:1964–72
129. Solari CD, Mare RD. 2012. Housing crowding effects on children's wellbeing. *Soc. Sci. Res.* 41:464–76
130. Springmann M, Mason-D'Croz D, Robinson S, Garnett T, Godfray HC, et al. 2016. Global and regional health effects of future food production under climate change: a modelling study. *Lancet* 387:1937–46
131. Steffen W, Crutzen PJ, McNeill JR. 2007. The Anthropocene: Are humans now overwhelming the great forces of nature? *AMBIO: J. Hum. Environ.* 36:614–21
132. Stone B, Hess JJ, Frumkin H. 2010. Urban form and extreme heat events: Are sprawling cities more vulnerable to climate change than compact cities? *Environ. Health Perspect.* 118:1425–28
133. Stubbs B, Vancampfort D, Rosenbaum S, Firth J, Cosco T, et al. 2017. An examination of the anxiolytic effects of exercise for people with anxiety and stress-related disorders: a meta-analysis. *Psychiatry Res.* 249:102–8
134. Sunega P, Lux M. 2016. Subjective perception versus objective indicators of overcrowding and housing affordability. *J. Hous. Built Environ.* 31:695–717
135. Tang B, Liu X, Liu Y, Xue C, Zhang L. 2014. A meta-analysis of risk factors for depression in adults and children after natural disasters. *BMC Public Health* 14:623
136. Tasian GE, Pulido JE, Gasparrini A, Saigal CS, Horton BP, et al. 2014. Daily mean temperature and clinical kidney stone presentation in five U.S. metropolitan areas: a time-series analysis. *Environ. Health Perspect.* 122:1081–87
137. Taylor KW, Novak RF, Anderson HA, Birnbaum LS, Blystone C, et al. 2013. Evaluation of the association between persistent organic pollutants (POPs) and diabetes in epidemiological studies: a national toxicology program workshop review. *Environ. Health Perspect.* 121:774–83
138. Taylor RG, Scanlon B, Döll P, Rodell M, van Beek R, et al. 2013. Ground water and climate change. *Nat. Clim. Change* 3:322–29
139. Thurstan RH, Roberts CM. 2014. The past and future of fish consumption: Can supplies meet healthy eating recommendations? *Mar. Pollut. Bull.* 89:5–11
140. Timaeus IM. 2012. Stunting and obesity in childhood: a reassessment using longitudinal data from South Africa. *Int. J. Epidemiol.* 41:764–72
141. Turner MC, Jerrett M, Pope CA 3rd, Krewski D, Gapstur SM, et al. 2016. Long-term ozone exposure and mortality in a large prospective study. *Am. J. Respir. Crit. Care Med.* 193:1134–42
142. U. N.-Habitat. 2011. *Cities and Climate Change: Global Report on Human Settlements 2011*. London/Washington, DC: U. N.-Habitat, Earthscan

143. U. N., Dep. Econ. Soc. Aff., Popul. Div. 2014. *World Urbanization Prospects. The 2014 Revision*. New York: U. N.
144. Ürge-Vorsatz D, Herrero ST, Dubash NK, Lecocq F. 2014. Measuring the co-benefits of climate change mitigation. *Annu. Rev. Environ. Resour.* 39:549–82
145. Vos T, Abajobir AA, Abbafati C, Abbas KM, Abate KH, et al. 2017. Global, regional, and national incidence, prevalence, and years lived with disability for 328 diseases and injuries for 195 countries, 1990–2016: a systematic analysis for the Global Burden of Disease Study 2016. *Lancet* 390:1211–59
146. Walker SP, Chang SM, Wright A, Osmond C, Grantham-McGregor SM. 2015. Early childhood stunting is associated with lower developmental levels in the subsequent generation of children. *J. Nutr.* 145:823–28
147. Wang Q, Li C, Guo Y, Barnett AG, Tong S, et al. 2017. Environmental ambient temperature and blood pressure in adults: a systematic review and meta-analysis. *Sci. Total Environ.* 575:276–86
148. Wang X, Sun D, Yao T. 2016. Climate change and global cycling of persistent organic pollutants: a critical review. *Sci. China Earth Sci.* 59:1899–911
149. Wania F, Mackay D. 1995. A global distribution model for persistent organic chemicals. *Sci. Total Environ.* 160:211–32
150. Warburton DER, Nicol CW, Bredin SSD. 2006. Health benefits of physical activity: the evidence. *CMAJ* 174:801–9
151. Warn E, Adamo SB. 2014. The impact of climate change: migration and cities in South America. *WMO Bull.* 63:10–14
152. Watts N, Adger WN, Agnolucci P, Blackstock J, Byass P, et al. 2015. Health and climate change: policy responses to protect public health. *Lancet* 386:1861–914
153. Wesseling C, Aragón A, González M, Weiss I, Glaser J, et al. 2016. Kidney function in sugarcane cutters in Nicaragua—a longitudinal study of workers at risk of Mesoamerican nephropathy. *Environ. Res.* 147:125–32
154. Whitmee S, Haines A, Beyrer C, Boltz F, Capon AG, et al. 2015. Safeguarding human health in the Anthropocene epoch: report of The Rockefeller Foundation–Lancet Commission on planetary health. *Lancet* 386:1973–2028
155. WHO (World Health Organization). 2013. *Global action plan for the prevention and control of non-communicable diseases 2013–2020*. Rep., WHO, Geneva. http://apps.who.int/iris/bitstream/handle/10665/94384/9789241506236_eng.pdf;jsessionid=71BCEA94B3F85737AB42F3C84216E54A?sequence=1
156. WHO (World Health Organization). 2014. *Global status report on noncommunicable diseases, 2014*. Rep., WHO, Geneva. http://apps.who.int/iris/bitstream/handle/10665/148114/9789241564854_eng.pdf;sequence=1
157. WHO (World Health Organization). 2016. *Ambient air pollution: a global assessment of exposure and burden of disease*. Rep., WHO, Geneva. <http://apps.who.int/iris/bitstream/handle/10665/250141/9789241511353-eng.pdf;sequence=1>
158. WHO (World Health Organization). 2017. *Preventing noncommunicable diseases (NCDs) by reducing environmental risk factors*. Rep., WHO, Geneva. <http://apps.who.int/iris/bitstream/handle/10665/258796/WHO-FWC-EPE-17.01-eng.pdf;sequence=1>
159. Wu G, Bazer FW, Cross HR. 2014. Land-based production of animal protein: impacts, efficiency, and sustainability. *Ann. N. Y. Acad. Sci.* 1328:18–28
160. Zou G, Decoster K, McPake B, Witter S. 2017. Renaming non-communicable diseases. *Lancet Global Health* 5:e656



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