nature ecology & evolution

Effects of urban-induced mutations on ecology, evolution and health

Received: 23 October 2023

Accepted: 13 March 2024

Published online: 19 April 2024

Check for updates

Marc T. J. Johnson ^{® 1,2}⊠, Irtaqa Arif^{1,2}, Francesco Marchetti³, **Jason Munshi-South** Φ^4 , Rob W. Ness^{1,2}, Marta Szulkin Φ^5 , Brian C. Verrelli Φ^6 , Carole L. Yauk⁷, Daniel N. Anstett⁸, Warren Booth⁹, Aude E. Caizergues^{1,2}, **Elizabeth J. Carlen**^{Φ **¹⁰, Anthony Dant¹¹, Josefa González** Φ ¹²,} **César González Lagos 13,14, Madeleine Oman 1,2, Megan Phifer-Rixey15, Diana J. Rennison  16, Michael S. Rosenberg ⁶ & Kristin M. Winchell ¹⁷**

Increasing evidence suggests that urbanization is associated with higher mutation rates, which can afect the health and evolution of organisms that inhabit cities. Elevated pollution levels in urban areas can induce DNA damage, leading to de novo mutations. Studies on mutations induced by urban pollution are most prevalent in humans and microorganisms, whereas studies of non-human eukaryotes are rare, even though increased mutation rates have the potential to afect organisms and their populations in contemporary time. Our Perspective explores how higher mutation rates in urban environments could impact the ftness, ecology and evolution of populations. Most mutations will be neutral or deleterious, and higher mutation rates associated with elevated pollution in urban populations can increase the risk of cancer in humans and potentially other species. We highlight the potential for urban-driven increased deleterious mutational loads in some organisms, which could lead to a decline in population growth of a wide diversity of organisms. Although benefcial mutations are expected to be rare, we argue that higher mutation rates in urban areas could infuence adaptive evolution, especially in organisms with short generation times. Finally, we explore avenues for future research to better understand the efects of urban-induced mutations on the ftness, ecology and evolution of city-dwelling organisms.

Mutation is the original source of all genetic variation. Despite its importance, variation in mutation rates is often overlooked or considered of negligible significance in empirical studies of ecology and evolution, particularly in eukaryotes^{[1](#page-7-0)}. Mutation rates can be influenced by the environment^{[2](#page-7-1),[3](#page-7-2)} and can evolve through time^{[4](#page-7-3),[5](#page-7-4)}. Neglecting to consider mutation may be especially problematic in cities, where emerging evidence suggests that pollution elevates mutation rates $6,7$ $6,7$.

One of the most consistent differences between urban and non-urban environments that could influence mutation rates is chemical pollution. Transportation, industry, wastewater management, home heating, landfills and pesticide application are all activities in urban areas commonly associated with elevated air, water and soil pollution[8](#page-8-0)–[10](#page-8-1). Although less frequent in urban areas, nuclear plants, nuclear testing and warfare can also result in highly mutagenic ionizing radiation¹¹. Studies on the mutagenic effects of radiation also provide general insight into how highly mutagenic pollutants can influence organisms in cities. Although pollution is not unique to urban areas, the concentration and diversity of pollutants are often highest in cities, exposing organisms to harmful stressors in unprecedented ways $8-10$ $8-10$ $8-10$.

A full list of affiliations appears at the end of the paper. \boxtimes e-mail: marc.johnson@utoronto.ca

Pollutant Chemical species Sources Medium Refs. PM $PM_{2.5}$ PM₁₀: inorganic ionic compounds, metal oxides, organic and elemental carbon Combustion by-products from traffic and industrial emissions, residential heating and reactions between pollutants Air [8](#page-8-0)[,18](#page-8-3) Volatile organic compounds Aldehydes, ketones, aromatics and alkanes Household products, building materials and combustion sources Air [18](#page-8-3)[,132](#page-10-0) PAHs Examples include benzo[a]pyrene, benzo[a] anthracene, chrysene, benzo[b]fluoranthene, benzo[k]fluoranthene Combustion by-products from industrial, residential and transport emissions Air/water/soil [33](#page-8-5)[,133](#page-10-1)–[135](#page-10-2) Sulfur oxides (SO_x) SO₂, sulfur trioxide (SO₃) Fossil fuel combustion, other industrial processes Air [8](#page-8-0).18 CO – Fossil fuel combustion, transport emissions Air [8](#page-8-0)[,136](#page-10-3) NO_x Nitrous oxide (NO), nitrogen dioxide (NO₂) Transport and industrial emissions Air [8](#page-8-0)[,137](#page-10-4)[,138](#page-10-5) Pesticides **Organophosphates, pyrethroids, carbamates,** polychlorinated biphenyls, polybrominated biphenyls, persistent organic pollutants Pesticide use in urban areas Water/soil [139](#page-10-6) Heavy metals Mercury, arsenic, cadmium, chromium and lead Industrial processes, mining Water/soil [9](#page-8-33)[,137](#page-10-4) High salt Salt (NaCl) Road salting Soil/water [140](#page-10-7)

Table 1 | Common urban chemical mutagens and carcinogens

For each pollutant, we indicate the chemical species, the most common anthropogenic sources, the medium in which the pollutant is typically encountered (air, water or soil) and references.

Urban chemical pollutants can cause physiological and genotoxic stress to organisms that may result in mutations. Such pollution is known to result in respiratory illnesses in humans¹², reduced photosynthesis and cell damage in plants 13 , higher mortality in fishes and amphibians 14 , and decreased fledgling success in birds^{[15](#page-8-23)}. Exposure to some pollutants can damage DNA and induce de novo mutations (hereafter simply called 'mutations') $16-19$. Although carcinogenic pollutants are known to cause somatic mutations (mutations in non-reproductive germ cell tissue), the fitness effects of these mutations and the prevalence of pollution-induced germline mutations are poorly understood outside of laboratory settings. Moreover, whether urban-induced higher mutation rates lead to an increased number of deleterious mutations, population decline or accelerated adaptive evolution has not been previously considered (but see ref. [20](#page-8-26)).

Our goal is to provide a forward-looking Perspective on the potential for elevated mutation rates in cities to influence the ecology and evolution of populations. Studies of the effects of urbanization on evolution have focused on genetic drift, gene flow and natural selection, and the potential for elevated mutation rates in cities to influence the ecology and evolution of populations is largely unexplored and of high priority for future research $^{21-24}$ $^{21-24}$ $^{21-24}$. We begin by reviewing urban pollutants and the damage they cause to DNA. Next, we consider how pollution affects somatic and germline mutations and the potential importance of these mutations for ecology and evolution. Although urban pollution can affect all organisms in cities, most existing examples come from research on humans. We consider the effects of pollution on humans and non-human organisms throughout this Perspective, and we use the extensive literature on humans as a model to understand the wider ecological and evolutionary impacts for all organisms. Looking beyond humans is important because although cities reduce and homogenize species diversity, urban habitats still harbour substantial biodiver- $sity^{25-27}$ $sity^{25-27}$ $sity^{25-27}$, and many of these species in cities are of conservation concern or have fundamental ecosystem roles²⁸. We end by discussing existing knowledge gaps and directions for future research.

Urban pollutants and damage to DNA

Air, water and soil in cities are consistently associated with a diverse mixture of pollutants (Table [1](#page-1-0) and Box [1\)](#page-2-0). The sources of most outdoor air pollutants in cities are combustion by-products from transportation, power generation, home heating/cooking and industry $29,30$ $29,30$. These by-products include pollutants such as polycyclic aromatic hydrocarbons (PAHs), nitrogen oxides (NO_x), sulfur dioxide (SO₂), carbon monoxide (CO) and various metal species (for example, Hg, Cu, Pb and Sn). These compounds can bind to particulate matter (PM), which can then be deposited in soil^{[18,](#page-8-3)[31](#page-8-4)-33}. Soil can also become contaminated with genotoxicants from industrial by-products, manufacturing, mining and road salting³⁰. Air pollutants, soil leaching, run-off and sewage all contribute to water pollution³⁴, which can lead to elevated levels of pesticides^{35,[36](#page-8-9)}, polychlorinated biphenyls³⁷, pharmaceutical products $38-40$ $38-40$ and microplastics $30,41,42$ $30,41,42$ $30,41,42$ in aquatic habitats.

Pollution in urban settings varies in both time and space in complex ways. The levels and types of urban pollution have changed throughout the history of industrial and urban growth. For example, during the past 20 years, the level of PM_{2.5} (PM with diameters <2.5 μ m) in Shanghai, China, has increased by over 200%, yet it decreased by nearly 30% in New York, USA, and remained consistently low in Melbourne, Australia (Fig. [1\)](#page-3-0). These changes through time are often influenced by changes in governmental policies (for example, the United States' Clean Air Act and the European Union's Ambient Air Quality Directive) and technological change, such as conversion from leaded to unleaded fuels. Urban pollutants also vary spatially in their concentrations and composition (Fig. [1](#page-3-0) insets). For example, industrial steel production often leads to some of the highest concentrations of PAHs^{[43](#page-8-15)}, whereas high vehicle traffic is typically associated with higher PM, ozone, CO and NO*x* (Table [1](#page-1-0)). Socio-economic variation among neighbourhoods often covaries with pollution levels, whereby poorer neighbourhoods are frequently in the most polluted areas, causing disparity in exposure to potentially harmful genotoxicants^{44,[45](#page-8-17)}. Non-urban areas also frequently experience pollution due to anthropogenic activities, including resource extraction, agriculture, forestry and nuclear radiation. However, we focus on urban areas because they are the fastest-growing ecosystem on Earth, and they are consistently associated with elevated pollution made up of diverse mixtures of chemicals that potentially harm organisms including causing damage to DNA (genotoxicants) (Box [1\)](#page-2-0).

The genotoxic effects of pollutants include chemical interactions that form DNA adducts (chemicals that bind to DNA) and reactive oxygen species that damage DNA (Box [1\)](#page-2-0). When such damage is improperly repaired, it can cause small-scale and large-scale mutations. Small-scale mutations include single nucleotide substitutions and small insertions/deletions (indels). Large-scale mutations involve large indels, duplications, translocations, inversions and aneuploidy $46-48$.

BOX 1

Genotoxicity of urban pollutants and induction of mutations

Chemical pollutants are the primary cause of DNA damage induced by urban pollution. Ionizing radiation is less common but is a more extreme mechanism of DNA damage in and around cities. When an organism is exposed to a chemical pollutant, the pollutant can cause DNA damage and mutation through several steps:

- (1) Pollutants can enter the cell via diffusion 157 or receptor-mediated endocytosis^{[158](#page-11-1)}
- (2) Once inside the cell:
	- (a) Pollutants (for example, PAHs) can form bonds with nitrogenous DNA bases, resulting in DNA adducts¹⁵
	- (b) Presence and interaction of pollutants with cellular processes or proteins causes increases in reactive oxygen species that can oxidize DNA and proteins^{[160](#page-11-3),1}
- (3) Chemically induced DNA lesions may be subject to error-prone DNA repair processes that cause mutations, or if the amount of damage exceeds the cell's capacity for DNA repair, it can result in mutations or chromosome damage^{[162](#page-11-5)}
- (4) Air pollutants can also cause oxidative stress via chronic inflammation and subsequent formation of reactive oxygen species 18

Ionizing radiation and radiomimetic compounds can alter DNA sequences through a diferent mechanism:

- (1) Radiation directly deposits energy in DNA, causing strand breaks, or it creates free radicals that damage DNA and proteins^{163[,164](#page-11-7)}
- (2) Free radical DNA damage includes apurinic/apyrimidinic sites and deamination of DNA bases (among others), both of which have unique mutagenic mechanisms 16
- (3) Lack of repair or error-prone repair of this damage can cause chromosomal aberrations and mutations

DNA replication errors such as unequal crossovers that can result in gene duplication and deletion are also possible. The location of DNA damage (coding versus non-coding regions), the molecular function of damaged DNA (regulatory versus structural) and whether coding mutations are synonymous or non-synonymous all can influence the molecular, physiological and fitness consequences of damage. The fitness effects of mutation can in turn impact the ecology and evolution of populations^{[49](#page-8-34)-51} (see 'Ecological and evolutionary consequences').

The effects of urban-induced mutations may differ between species because of variation in ploidy, cellular complexity, mutation rate, reproductive system, population size and generation time. For example, many animals, higher plants and some eukaryotic microorganisms live primarily as diploids or polyploids, which can mask the fitness effects of recessive mutations at low frequencies $52,53$ $52,53$ $52,53$. Similarly, many multicellular organisms have differentiated germ and somatic cells, such that pollution-induced mutations in somatic cells will not generally be passed on to subsequent generations. By contrast, organisms with no distinction between germ and soma, such as some plants and fungi, may accumulate inherited mutations more rapidly if mutations arise in the cells that ultimately form gametic tissue $54,55$ $54,55$. Moreover, mutation rates vary by orders of magnitude, with bacteria and microbial eukaryotes having the lowest rates, vascular plants and animals having moderate rates, and viruses having the highest mutation rates[4,](#page-7-3)[56](#page-8-40). Recombination in sexual organisms can allow more efficient purging of harmful mutations by selection than in asexual populations $57,58$ $57,58$. Finally, large populations with rapid generation times are expected to purge or fix environmentally induced mutations that affect fitness more rapidly than small long-lived populations 59 . In the sections that follow, we expand on how such variation among species may lead to different ecological and evolutionary consequences of urban-induced mutations.

Somatic mutations

The primary consequence of genotoxic exposure is the induction of somatic mutations that can adversely affect molecular, cellular and tissue function. Somatic mutations are not transmitted to the next generation unless they occur in germ cell progenitors, such as plant apical meristems⁶⁰, so they typically affect only the exposed individual's health and fitness. The causal role of chemically induced mutations in cancer development is well known in certain cases, such as lung cancer due to tobacco smoke⁶¹ (Table [2](#page-4-0)). These examples show that exposure to genotoxicants can cause mutations in tumour suppressor genes or proto-oncogenes that can function as cancer drivers, causing cellular proliferation, tumour development and genetic instability 62 . Moreover, exposure to mutagens during key life stages, especially embryogenesis and organogenesis, may increase the probability of clonal expansion of mutation-bearing cells^{17[,63,](#page-9-5)64}. Data supporting the association between environmentally induced mutations and non-cancerous diseases are almost entirely lacking, despite knowledge of mutations across the genome caused by genotoxicant exposure and a growing understanding of the role of somatic cell mutagenicity in disease more generally (for example, ageing, neurological and cardiac diseases) $65,66$ $65,66$. Thus, there is currently no knowledge on the rates and functional consequences of pollution-induced somatic mutations for individuals, populations and species beyond the established association with cancer.

The study of mutagenesis is challenging because mutations are rare events at a genomic scale. This difficulty is compounded in the case of somatic mutations because the occurrence of mutations varies among tissues within a single individual. However, a variety of studies provide empirical evidence supporting an association between specific urban pollutants and elevated somatic cell mutation rates. The invention of the *Salmonella* mutation assay, often called the 'Ames assay', has been a transformative tool in the study of environmental mutagenesi[s67](#page-9-9)[,68](#page-9-10). In brief, the assay assesses how frequently *Salmonella* strains lacking the ability to metabolize histidine—due to engineered base-pair substitutions or frameshift mutations—exhibit revertant mutations to restore histidine metabolism when challenged by a toxicant^{[17](#page-8-42),67}. This simple bacterial assay has revealed that the air, soil and water in urban environments is replete with mutagens⁶⁷. Beyond *Salmonella*, observational and experimental cytogenetic studies show that numerous chemical pollutants cause chromosomal abnormalities in diverse organisms, including structural aberrations and aneuploidy^{[16](#page-8-24),[17](#page-8-42),69}. Additional lines of evidence are based on the types and distribution of mutations (the mutation spectrum) observed in human cancers used to infer mutagenic exposures⁷⁰ as well as the COSMIC database⁷¹. Overall, laboratory models (for example, *Salmonella*, mice and plants) exposed to environmental media or extracts demonstrate the widespread mutagenicity of many chemical pollutants in urban areas⁷⁰.

The most extensive evidence of pollution-induced somatic cell mutagenicity is from studies on combustion-related by-products found in urban air pollution, contaminated soils and sediments. The weight of evidence for the mutagenicity of outdoor air pollution is high, with many specific agents declared "carcinogenic to humans" by the International Agency for Research on Cancer¹⁸. This agency's monographs thoroughly describe how these urban pollutants cause mutagenicity in laboratory organisms as diverse as bacteria, plants and rodents $18,72$ $18,72$ $18,72$. For example, the mutation spectrum observed in lung tumours of

carcinogenic pollutants. Concentrations of PM_{2.5} across terrestrial Earth in 2019–2020, with inset panels illustrating that concentrations are frequently highest in and around cities^{[152](#page-11-9),153}. PM_{2.5} concentrations have been changing through time (top right inset), increasing in some cities (for example, Shanghai, China) and decreasing in others (for example, New York, USA)¹⁵⁴. The stacked bar charts show how the composition of major carcinogenic pollutants

varies among countries^{[155](#page-11-12),156}. High concentrations of PM_{2.5} outside urban areas are caused by a combination of anthropogenic sources such as long-distance dispersal of industrial pollution, burning of crops in agricultural regions, forest fires and naturally occurring fine dust picked up by strong winds from bare soil, especially in arid regions (for example, Saharan and sub-Saharan Africa).

non-smokers associated with air pollution is broadly consistent with exposure to bulky-DNA-adduct-forming chemicals such as benzo[a] pyrene $73,74$ $73,74$. Additional evidence for the mutagenicity of air pollution comes from humans exposed to high levels of combustion by-products in residential and occupational settings, whereby individuals exhibit cytogenetic damage to various cell types^{75,76}, and the urine from such individuals is mutagenic to bacterial cells^{[77,](#page-9-19)[78](#page-9-20)}. Moreover, soil and sediments that contain combustion-related contaminants are mutagenic to organisms that frequently come into contact with these substrates, such as bacteria and plants^{[17](#page-8-42),69}. Undoubtedly, inhabitants of any urban ecosystem are exposed to mutagenic particulate pollutants associated with combustion emissions.

There are many other examples of mutagenic contaminants found in urban settings, from metals to pesticides, organochlorines and benzene (Table [1\)](#page-1-0). These genotoxicants have the potential to impact the somatic cell mutation burden, contributing to the decreased health of individuals and populations $18,79$ $18,79$. The vast majority of mutagenicity testing is conducted in the laboratory on individual chemicals at high $doses¹⁹$ $doses¹⁹$ $doses¹⁹$, leading to a major gap in our understanding of how lifelong, low-dose exposures to mixtures of mutagens affect mutation rates and disease outcomes. Moreover, the complex interactions between sociodemographic factors and mutagenic environmental mixtures inherent to cities have yet to be explored.

The study of environmentally induced somatic cell mutations has been considerably hampered by the lack of tools available out-side of the laboratory. Although single-cell deep-sequencing^{[80](#page-9-22)} and error-corrected sequencing $81,82$ $81,82$ methodologies exist, these have mostly been applied in clinical settings and have yet to be extended to studies on environmental exposures in natural populations. The high levels of pollution in urban areas offer an opportunity to address these obstacles using field experiments, in addition to laboratory experiments, that apply genomic technologies to directly quantify mutation frequency and spectrum in a diverse array of organisms (see 'Future directions').

Germline mutations

Unlike somatic mutations, germline mutations are inherited between generations. For this reason, it is primarily germline mutations that can influence the evolution of populations. Although germline mutations are rare at the individual level, even the smallest increase in the mutation rate can have large consequences for populations 83 83 83 .

Table 2 | Cancers associated with urban-induced mutations

Examples of the most common cancers associated with urban-induced mutations, including changes in rates of cancer in urban and non-urban populations. For each example, we indicate the study region, the pollutant studied and the main findings.

Laboratory and field studies suggest that exposure to many common urban pollutants can induce germline mutations. For example, over 80 chemical agents have been identified as germline mutagens in laboratory mice 19 . In humans, the best evidence of the impact of pollutants on germ cell mutagenesis comes from studies demonstrating an increased incidence of chromosomal abnormalities in human sperm¹⁹. Such abnormalities may explain the significant correlation between paternal blood dioxin levels due to occupational exposure and increased mutation rates in their offspring 84 . When considering exposure to radiation as an example of extreme exposure to a mutagen, children of parents exposed to ionizing radiation following the Chernobyl nuclear plant accident exhibited increased rates of tandem repeat mutations^{[85](#page-9-27)}. Similar inherited mutations have been observed in plants⁸⁶ and barn swallows⁸⁷. However, increases in inherited single nucleotide variants have yet to be conclusively demonstrated for humans exposed to radiation⁸⁸. In non-polluted areas, a recent study reported a reduced mutation rate in an Amish population, which has been interpreted as traditional rural lifestyles leading to low mutation rates because of reduced exposure to chemical mutagens^{[89](#page-9-31)}. Very few studies have examined non-human populations outside of laboratory conditions, and they show that birds and rodents exhibit increased heritable mutation rates in repetitive DNA regions when exposed to ambient industrial air pollution $6,7,90,91$ $6,7,90,91$ $6,7,90,91$ $6,7,90,91$.

In addition to pollution, urban and rural human populations diverge in their demographic patterns in ways that are expected to influence germline mutation rates. In recent decades, there has been a trend for delayed childbearing in many countries. In both developed and developing nations, this delay is more pronounced in urban set-tings than in rural settings^{[92](#page-9-34),93}. Studies of human parents and offspring over the past decade have consistently demonstrated an age-related increase in mutation rates, especially in fathers 94 . It is estimated that fathers transmit ~1.2 additional mutations for each year of age, versus ~0.4 new mutations per year of age in the mother. The higher paternal contribution is partially ascribed to the continuous production of sperm as men age, whereas no new oocytes are generated once a female individual is born. Surprisingly, the urban-biased shift towards delaying the age of reproduction is the only clear example of how urban living is associated with elevated germline mutation rates, other than urban pollution inducing mutations in repetitive regions of birds and mice. The consistency of divergence in parental age between urban and rural populations in developed and developing nations requires further investigation, as this major source of increased mutation rates could also result from differences in socio-economic factors and cultural differences throughout the world. There is also evidence that non-human organisms exhibit demographic shifts in urban habitats^{[95](#page-9-37)}, but whether this is associated with changes in mutation rates requires investigation.

Despite the circumstantial evidence mentioned above for an effect of urban pollution and demographics on increased germline mutation rates, a direct link between urban pollution and mutations has yet to be definitively demonstrated using modern genome sequencing techniques. We therefore lack information on how and when urban pollution increases germline mutation rates, the targets of mutation and especially their phenotypic and fitness effects.

Ecological and evolutionary consequences

Alterations to the rate and spectrum of both somatic and germline mutations due to urban pollution could have important ecological and evolutionary repercussions. Theoretical and empirical studies show that the majority of new functionally significant mutations are deleterious and removed by purifying selection⁹⁶. If deleterious mutations are elevated in urban settings, either due to a higher rate or as a larger fraction of deleterious mutations, we expect an increased mutation load (reduced fitness due to the burden of deleterious mutations relative to an unmutated individual) that will decrease population mean fitness $97,98$ $97,98$. Whether urban species in fact suffer a demographic decline depends on several factors including the strength of selection, effective population size (N_e) and generation time (Fig. [2\)](#page-5-0). Keightley⁹⁹ estimated that the decline in human fitness due to mutation could reach 0.01% per generation, and the decline would change linearly with changes in mutation rate. This estimate does not include the countering force of purifying selection. It is therefore likely that organisms with long generation times will experience little effect on population mean fitness in the short term. Conversely, organisms such as microorganisms that have short generation times may experience changes in fitness over contemporary timescales.

Although evolutionary responses depend on inherited germline mutations, somatic mutations also have important consequences for the health and fitness of individuals that contribute to long-term population viability. In multicellular organisms, somatic mutations can create a mosaic of cells with slightly different genotypes^{[100](#page-9-42)}. These mutations can lead to developmental instability, which is particularly detrimental in organisms with strict body plans such as animals 101 (Table [2](#page-4-0)). The genomic diversity within an individual can also produce competition among cell lineages that can be harmful, as in the case

Fig. 2 | The potential for elevated mutation rates in cities to affect the evolution of a population relative to a fitness optimum. When a population starts at a fitness optimum (dashed horizontal black line) in an urban environment (blue lines), any increase in the mutation rate (∆*µ*) can lead to a net increase in deleterious mutations within a population, moving the population further from the fitness optimum. If urban pollution elevates mutation rates in urban areas (that is, high ∆*µ*, indicated by the solid blue line), then we predict a population will move further from the fitness optimum through time. If ∆*µ* is low but still >0, then this effect can be relatively small. By contrast, when a population is initially maladapted to an urban environment (red lines), such that it starts far away from the fitness optimum, then higher mutation rates in urban areas (solid red line) can lead to rapid adaptation such that the population quickly evolves towards the fitness optimum. The rate of this evolution will be slower when ∆*µ* is lower (red dashed line). Such adaptive evolution could lead to evolutionary rescue, but such dynamics are only likely over contemporary time when N_e is high and generation times are short (as in viruses, bacteria and eukaryotic microorganisms). At equilibrium, populations are below the fitness optimum because elevated mutation rates in urban areas increase a population's mutation load. Moreover, populations experiencing higher ∆*µ* are predicted to have lower fitness than those with lower ∆*µ* because most new mutations will be deleterious when a population is close to its fitness optimum. A population may remain maladapted (scenario not shown) when N_e is low and generation times are long, which could lead to extinction if population growth rates are negative.

of cancers. There is also clear evidence for intra-organismal selection for healthy cell lineages that can reduce the overall impact of deleterious mutations, including in marine tunicates and long-lived perennial plants $100,101$ $100,101$. These different phenomena hint at complex interactions between development, life history and genetic systems when determining the relative impact of elevated somatic mutation rates in urban settings. Given the evidence that urban habitats have elevated concentrations of numerous mutagens (Table [1](#page-1-0)), the impact of somatic mutations may become very important to predicting the sustainability of some urban populations (see 'Applied impacts').

Theory generally predicts an advantage for reduced mutation rates because most non-neutral mutations are deleterious^{[102,](#page-10-14)[103](#page-10-15)}. We might therefore expect that urban populations will be under selection to reduce mutation rates in the presence of mutagens. The ability and time it takes for selection to reduce mutation rates will depend on numerous factors such as the mating system, N_e and target size (the amount of nucleotide sequence that can reduce mutation rate) for mutation modifiers¹⁰⁴. The drift-barrier hypothesis¹⁰⁵ predicts that directional selection will reduce mutation rates until a point at which the strength of genetic drift (1/ N_e) overcomes the selective advantage (*s*) of smaller improvements in mutation rate (when N_e s < 1). This hypothesis is supported by recent comparative genomic analyses that show that species with higher long-term N_e and shorter generation times tend to have lower mutation rates per generation $^{\textrm{\tiny{S}}}$. There is an equilibrium point beyond which if mutation rates are sufficiently high, selection to reduce the mutation rate should overcome drift. Nevertheless, if urban environments reduce an organism's N_e , resulting in a loss of genetic diversity^{[22](#page-8-43)}, we may expect a higher equilibrium mutation rate.

Despite the genetic load created by deleterious mutations, mutation also provides the raw variation necessary for adaptation. These contrasting effects of mutation lead to the possibility that mutation-fuelled adaptation can result in an "evolutionary rescue["98](#page-9-40)[,106](#page-10-18) (that is, an increase in the population growth rate of small populations due to adaptation) of populations subject to environmental challenges in urban environments (Fig. [2\)](#page-5-0). For example, pathogens whose fitness in a new host is so low as to preclude persistence may benefit from higher mutation rates, where the higher the mutation rate, the larger the probability of evolutionary rescue^{[107](#page-10-19)}. However, this situation is highly context-dependent—once a population approaches its fitness optimum, any new mutations are likely to be deleterious. It is reasonable to speculate that urban environments will pose such strong selective pressures that some populations will benefit from elevated mutational input during initial establishment (Fig. [2\)](#page-5-0). The extent to which mutation will provide variation to tackle new selective challenges will depend on how elevated the mutation rate is in urban areas, how close a population is to a fitness optimum, N_e and generation time (Fig. [2](#page-5-0)). If elevated mutation rates have beneficial implications for species colonizing urban environments, it may also mean that cities could facilitate rapid adaptation to pesticides, herbicides and antibiotics or provide the raw variation needed for pathogens to switch hosts.

It is plausible that elevated patterns of mutation in cities could facilitate speciation, especially if mutations induced by urban pollution cause chromosomal changes that affect mating compatibility, ecology or physiology. Elevated mutation rates in cities could help to fuel population divergence among urban and non-urban populations via local adaptation and accelerate genetic drift due to population fragmentation¹⁰⁸. Under these conditions, higher mutation rates in urban settings would increase the possibility of generating mutations that are compatible with population-specific local alleles at other loci but incompatible with alleles in populations adapted to non-urban environments. Alleles that are compatible only with the genetic background in which they arose are called Bateson–Dobzhansky–Muller incompatibilities and often form the genetic basis of speciation 109 . Such incompatibilities may be particularly likely to occur if urban pollutants increase the frequency of chromosomal abnormalities or large structural mutations, including inversions, translocations, polyploidy or elevated activity of transposable elements. It is these types of large-scale structural mutations that are most commonly associated with genes that influence reproductive isolation and large changes in ecology and physiology¹¹⁰. Even in the absence of reproductive isolation, reduced vigour of urban and non-urban hybrids could alter the fitness of nearby populations. Overall, because elevated mutation rates in urban areas have the potential to lead to increased divergence¹⁰⁸, we believe that cities offer unique opportunities to study the process of speciation in real time.

Applied impacts

Given that urbanization can increase mutation rates, we expect numerous applied consequences associated with the health and conservation of organisms inhabiting cities. The anticipated health effects on humans and non-human species include cancers and other diseases linked to somatic and germline mutations. The conservation consequences relate to how elevated mutation rates are expected to influence the fitness and long-term population growth of urban-dwelling species (Fig. [2](#page-5-0)).

Urban pollution causes numerous types of cancer in humans and other organisms. Contemporary urban pollution elevates lung $74,111$ $74,111$, breast¹¹² and other forms of cancer¹¹³ by 10% to 1,000% above baseline incidence rates (Table [2\)](#page-4-0). The magnitude of these effects varies among cities and over time because of variation in the types and concentrations of pollutants (Fig. [1\)](#page-3-0). Admittedly, most research on the health

Fig. 3 | Potential biosentinel species for studying urban-associated mutations. a–**l**, Proposed biosentinels include *Salmonella enterica* (**a**), *Caenorhabditis elegans* (**b**), *Drosophila melanogaster* (**c**), *Arabidopsis thaliana* (**d**), *Trifolium repens* (**e**), *Flavoparmelia caperata* (a lichen) (**f**), *Fundulus majalis* (**g**), *Passer domesticus* (**h**), *Columba livia* (**i**), *Mus musculus* (**j**), *Rattus norvegicus* (**k**) and *Canis lupus familiaris* (**l**). An image of humans (*Homo sapiens*) is not shown but is included in the schematics below. These species represent a range of traditional laboratory model organisms used for studying genetic and evolutionary processes, as well as emerging models for studying ecological responses to pollution or evolution in urban areas. **m**,**n**, Some species offer a combination of fast generation time and excellent genomic resources for

effects of urban pollution has been done on humans and rodents. How urban pollution affects somatic mutations and cancers in non-model organisms is poorly understood, especially outside of laboratory settings, and represents a gap in knowledge^{[114](#page-10-26)-[116](#page-10-27)} (see 'Future directions'). Although heritable germline mutations have the potential to magnify cancer risk in offspring due to pollution exposure in parents, there is currently no evidence outside the laboratory of environmentally induced heritable mutations causing cancer, even for ionizing radiation^{[19](#page-8-25)[,88](#page-9-30),11[7](#page-7-6)}. However, observational studies of birds⁷ and labora-tory studies of rodents^{[6](#page-7-5),91} confirm that air pollution from steel mills can induce heritable germline mutations in repetitive DNA regions, which suggests that urban-induced mutations in cancer driver genes could also be inherited. Understanding how, when and where urban pollution leads to inherited mutations that influence cancer risk is an important goal for future research (see 'Future directions').

mutagenic studies (**m**), whereas others are more directly relevant to humans (that is, with respect to health and well-being) and urbanization (that is, owing to their relative abundance in urban versus non-urban habitats) given their commensal status with humans (**n**). Credits: Phanie - Sipa Press/Alamy Stock Photo (**a**), Science Photo Library/Alamy Stock Photo (**b**), Itsik Marom/Alamy Stock Photo (**c**), thrillerfillerspiller/Alamy Stock Photo (**d**), Nigel Cattlin/Alamy Stock Photo (**e**), Clarence Holmes Wildlife/Alamy Stock Photo (**f**), Robert S. Michelson/Tom Stack & Assoc./Alamy Stock Photo (**g**), robertharding/Alamy Stock Photo (**h**), M. Johnson (**i**), Tim Mander/Alamy Stock Photo (**j**), Dave Bevan/ Alamy Stock Photo (**k**), K. L. Howard/Alamy Stock Photo (**l**).

Multiple socio-ecological factors associated with urban lifestyles could interact with pollution to elevate mutation rates. The previously mentioned shift to older parental age among people in urban compared with non-urban communities is the best-known cause of higher germline mutation rates in urban populations^{[94](#page-9-36)}. Urban mutagenic pollution probably interacts with and amplifies this demographic effect on mutation rates. Human urban populations also exhibit increased rates of obesity and associated cancers due to a large proportion of processed foods in urban diets and relatively sedentary lifestyles^{[118](#page-10-29)}. Wildlife species also exhibit altered diets in cities that incorporate more anthropogenic food sources such as sugar, corn and wheat. Such diet shifts have been linked to higher body mass and hyperglycaemia in some species^{[119](#page-10-30)-121}. Food additives and contaminants in processed foods may influence germline mutation rates 122 , as could shifts in urban gut microbiomes 123 . Exposure to environmental pollutants and lack of access to high-quality diets may be biased towards certain urban demographics. Analysing urban mutagenesis and other evolutionary processes is thus an important step to address concerns about environmental justice $24,124,125$ $24,124,125$ $24,124,125$ $24,124,125$.

Elevated mutation rates in cities have the potential to influence the dynamics of urban populations (Fig. [2\)](#page-5-0). Given that most mutations are neutral or deleterious, it is likely that urban-induced mutations will frequently negatively affect individual fitness and population growth rate $97,98$ $97,98$. Determining whether such negative demographic effects will be sufficiently large to outweigh the influence of other factors requires careful quantification and modelling. We expect that the urban pollution-induced mutational load will be one of many factors threatening the persistence of populations and may become a conservation concern for rare or declining native species in cities. By contrast, we predict that populations of pests and other organisms that maintain large populations are less likely to be negatively affected by elevated mutation rates.

It is unlikely that urban-induced mutations will positively influence conservation through evolutionary rescue for most species. Only organisms with rapid generation times and high N_e are expected to experience positive long-term fitness effects of elevated mutation rates, and even then, only when selection is strong (Fig. [2](#page-5-0)). Such scenarios are most likely to apply to viruses, bacteria and some eukaryotic microorganisms (for example, yeast and algae), raising the possibility that elevated mutation rates in cities could promote the spread of pathogenic organisms¹⁰⁷. Field and laboratory experiments that examine how urbaninduced changes in mutation rates affect known and emerging diseases and pests could have important implications for public health.

Future directions

Our Perspective illustrates that water, soil and air pollution in urban areas increases mutation rates, but the magnitude and mutational spectrum of this increase, as well as its ecological and evolutionary consequences, remain unresolved. These gaps represent important problems requiring attention, which we outline as research questions below.

What is the magnitude of increase in somatic and germline mutation rates, and what are the types of mutations caused by urban pollution?

Although it is important to refine how somatic mutation rates are influenced by urban pollution, the greatest need remains establishing whether and under what circumstances urban pollution causes ger-mline mutations in wild populations^{[19](#page-8-25)}. Conventional genomic technologies are poorly suited for quickly surveying the mutagenic properties of changing environments such as urban areas. New error-corrected sequencing approaches enable the study of rare mutations within a heterogenous population of cells $126,127$ $126,127$. These methods can facilitate more rapid and definitive tests of how urban pollution affects mutation rates because they rely on uniquely labelling individual DNA molecules prior to sequencing, which allows the removal of PCR and sequencing errors associated with standard next-generation sequencing. This enables, for the first time, the accurate quantification of rare mutations directly in the exposed organism.

What are the fitness effects of urban-induced mutations, and how do these influence the ecology and evolution of populations?

Answering this question will require a combination of laboratory and field experiments, coupled with genome sequencing. Laboratory experiments could establish how mutations caused by specific urban pollutants influence individual fitness, population growth and (mal) adaptation. Field experiments could follow the fitness of individuals that exhibit the presence or absence of mutations. Such experiments could be expanded on by experimentally recreating mutations via transgenic or CRISPR manipulations. Finally, identification of somatic and germline mutations from human and wild urban populations of diverse organisms (Fig. [3](#page-6-0)) could be used to infer fitness and health effects on the basis of how the types and locations of mutations are expected to disrupt homeostasis using deep learning models of DNA sequence evolution across thousands of species 128 .

How do urban-induced mutations vary among species?

There is a need to expand the investigation of mutations caused by pollution to a wider diversity of organisms beyond humans given the indiscriminate threats of urban pollutants to all species. We propose a global research programme that uses a range of organisms as biosentinels (organisms to assay mutations induced by pollution), where the species chosen would vary in their relevance to humans, prevalence in urban areas, generation time and genomic resources (Fig. [3](#page-6-0)). A biosentinel programme could detect mutagenic effects even when specific mutagens are difficult to identify^{[129](#page-10-39),[130](#page-10-40)}. Bacteria, plants and human cell lines have all been proposed as urban biosentinels¹³¹. *Salmonella* has been the vanguard biosentinel because it responds readily to both known and unknown mutagens^{[68](#page-9-10)}, and we see it as a possible bacterial model moving forward (Fig. [3](#page-6-0)). Existing plant (*Arabidopsis*) and animal (*Drosophila* and *Caenorhabditis elegans*) model organisms offer a rich genomic toolkit, although given their marginal importance to humans and/or prevalence in urban areas, non-model organisms that have been the focus of studies in urban areas should also be included, such as white clover, dogs and various birds. Rodents, particularly house mouse (*Mus musculus*) and Norway rat (*Rattus norvegicus*), are important pests in urban areas that are commonly used in laboratories, offering a biosentinel model that more closely resembles human physiology^{[16](#page-8-24)}. The deployment of such biosentinels could provide a rapid and accurate view of how urban-induced mutations affect the biology of urban-dwelling species, including humans.

Conclusions

Our Perspective highlights the potential broad-ranging mutagenic effects of urban pollution on virtually all life in cities. These mutagenic effects are expected to influence the fitness, ecology and evolution of wild populations, but these effects are largely unstudied outside of laboratory settings, and even there, only a small subset of species have been studied. Given the many mutagens that are prevalent in urban areas and their potentially large impacts on human and wildlife fitness, we argue that the study of urban mutagenesis is in urgent need of attention and should be prioritized in future research on health, ecology and evolution.

References

- 1. Fenster, C. B. & Murren, C. J. Commentary: mutation: source of variation in evolutionary ecology. *Evol. Ecol.* **34**, 311–314 (2020).
- 2. MacLean, R. C., Torres-Barceló, C. & Moxon, R. Evaluating evolutionary models of stress-induced mutagenesis in bacteria. *Nat. Rev. Genet.* **14**, 221–227 (2013).
- 3. Fitzgerald, D. M., Hastings, P. & Rosenberg, S. M. Stress-induced mutagenesis: implications in cancer and drug resistance. *Annu. Rev. Cancer Biol.* **1**, 119–140 (2017).
- 4. Lynch, M. et al. Genetic drift, selection and the evolution of the mutation rate. *Nat. Rev. Genet.* **17**, 704–714 (2016).
- 5. Bergeron, L. A. et al. Evolution of the germline mutation rate across vertebrates. *Nature* **615**, 285–291 (2023).
- 6. Somers, C. M., McCarry, B. E., Malek, F. & Quinn, J. S. Reduction of particulate air pollution lowers the risk of heritable mutations in mice. *Science* **304**, 1008–1010 (2004).
- 7. Yauk, C. L. & Quinn, J. S. Multilocus DNA fingerprinting reveals high rate of heritable genetic mutation in herring gulls nesting in an industrialized urban site. *Proc. Natl Acad. Sci. USA* **93**, 12137–12141 (1996).

- 8. *Ambient Air Pollution: A Global Assessment of Exposure and Burden of Disease* (World Health Organization, 2016).
- 9. *Global Assessment of Soil Pollution—Summary for Policy Makers* (FAO and UNEP, 2021).
- 10. *A Snapshot of the World's Water Quality: Towards a Global Assessment* (UNEP, 2016).
- 11. Filburn, T., Bullard, S. & Bullard, S. G. *Three Mile Island, Chernobyl and Fukushima* (Springer, 2016).
- 12. Seaton, A., Godden, D., MacNee, W. & Donaldson, K. Particulate air pollution and acute health efects. *Lancet* **345**, 176–178 (1995).
- 13. Seyyednejad, S., Niknejad, M. & Koochak, H. A review of some diferent efects of air pollution on plants. *Res. J. Environ. Sci.* **5**, 302–309 (2011).
- 14. Casey, R., Shaw, A., Massal, L. & Snodgrass, J. Stormwater retention ponds in suburban Maryland, USA. *Bull. Environ. Contam. Toxicol.* **74**, 273–280 (2005).
- 15. Chatelain, M. et al. Urban metal pollution explains variation in reproductive outputs in great tits and blue tits. *Sci. Total Environ.* **776**, 145966 (2021).
- 16. Claxton, L. D. & Woodall, G. M. Jr A review of the mutagenicity and rodent carcinogenicity of ambient air. *Mutat. Res. Rev. Mutat. Res.* **636**, 36–94 (2007).
- 17. White, P. A. & Claxton, L. D. Mutagens in contaminated soil: a review. *Mutat. Res. Rev. Mutat. Res.* **567**, 227–345 (2004).
- 18. IARC *Outdoor Air Pollution* IARC Monographs on the Evaluation of Carcinogenic Risks to Humans Vol. 109 (WHO Press, 2016).
- 19. Marchetti, F., Douglas, G. R. & Yauk, C. L. A return to the origin of the EMGS: rejuvenating the quest for human germ cell mutagens and determining the risk to future generations. *Environ. Mol. Mutagen.* **61**, 42–54 (2020).
- 20. Bromham, L., Hua, X., Lanfear, R. & Cowman, P. F. Exploring the relationships between mutation rates, life history, genome size, environment, and species richness in flowering plants. *Am. Nat.* **185**, 507–524 (2015).
- 21. Diamond, S. E. & Martin, R. A. Evolution in cities. *Annu. Rev. Ecol. Evol. Syst.* **52**, 519–540 (2021).
- 22. Johnson, M. T. J. & Munshi-South, J. Evolution of life in urban environments. *Science* **358**, aam8327 (2017).
- 23. Szulkin, M., Munshi-South, J. & Charmantier, A. *Urban Evolutionary Biology* (Oxford Univ. Press, 2020).
- 24. Verrelli, B. C. et al. A global horizon scan for urban evolutionary ecology. *Trends Ecol. Evol.* **37**, 1006–1019 (2022).
- 25. Aronson, M. F. et al. A global analysis of the impacts of urbanization on bird and plant diversity reveals key anthropogenic drivers. *Proc. R. Soc. B* **281**, 20133330 (2014).
- 26. Knapp, S. et al. Phylogenetic and functional characteristics of household yard floras and their changes along an urbanization gradient. *Ecology* **93**, S83–S98 (2012).
- 27. Rogers, A. M., Yong, R. Q. Y. & Holden, M. H. The house of a thousand species: the untapped potential of comprehensive biodiversity censuses of urban properties. *Ecology* **105**, e4225 (2023).
- 28. Lambert, M. R. & Donihue, C. M. Urban biodiversity management using evolutionary tools. *Nat. Ecol. Evol.* **4**, 903–910 (2020).
- 29. Leung, D. Y. Outdoor-indoor air pollution in urban environment: challenges and opportunity. *Front. Environ. Sci.* **2**, 69 (2015).
- 30. *Towards a Pollution-Free Planet: Background Report* (UNEP, 2017).
- 31. Baensch-Baltruschat, B., Kocher, B., Stock, F. & Reiferscheid, G. Tyre and road wear particles (TRWP)—a review of generation, properties, emissions, human health risk, ecotoxicity, and fate in the environment. *Sci. Total Environ.* **733**, 137823 (2020).
- 32. Nirmalkar, J., Haswani, D., Singh, A., Kumar, S. & Raman, R. S. Concentrations, transport characteristics, and health risks of PM2.5-bound trace elements over a national park in central India. *J. Environ. Manage.* **293**, 112904 (2021).
- 33. IARC *Some Non-heterocyclic Polycyclic Aromatic Hydrocarbons and Some Related Exposures* IARC Monographs on the Evaluation of Carcinogenic Risks to Humans Vol. 92 (WHO Press, 2010).
- 34. Martínez-Bravo, M. & Martínez-del-Río, J. in *Encyclopedia of the UN Sustainable Development Goals: Sustainable Cities and Communities* (eds Leal Filho, W. et al.) 905–915 (Springer, 2020).
- 35. Nagy, K., Rácz, G., Matsumoto, T., Ádány, R. & Ádám, B. Evaluation of the genotoxicity of the pyrethroid insecticide phenothrin. *Mutat. Res. Genet. Toxicol. Environ. Mutagen.* **770**, 1–5 (2014).
- 36. Annabi, E., Ben Salem, I. & Abid-Essefi, S. Acetamiprid, a neonicotinoid insecticide, induced cytotoxicity and genotoxicity in PC12 cells. *Toxicol. Mech. Methods* **29**, 580–586 (2019).
- 37. Agudo, A. et al. *Polychlorinated Biphenyls and Polybrominated Biphenyls* (WHO Press, 2016).
- 38. Chowdhury, J., Mandal, T. K. & Mondal, S. Genotoxic impact of emerging contaminant amoxicillin residue on zebra fish (*Danio rerio*) embryos. *Heliyon* **6**, E05379 (2020).
- 39. Isidori, M., Lavorgna, M., Nardelli, A., Pascarella, L. & Parrella, A. Toxic and genotoxic evaluation of six antibiotics on non-target organisms. *Sci. Total Environ.* **346**, 87–98 (2005).
- 40. Metzler, M., Kulling, S. E., Pfeifer, E. & Jacobs, E. Genotoxicity of estrogens. *Z. Lebensm. Unters. Forsch. A* **206**, 367–373 (1998).
- 41. Tagorti, G. & Kaya, B. Genotoxic efect of microplastics and COVID-19: the hidden threat. *Chemosphere* **286**, 131898 (2022).
- 42. Roursgaard, M. et al. Genotoxicity of particles from grinded plastic items in Caco-2 and HepG2 cells. *Front. Public Health* **10**, 906430 (2022).
- 43. Yang, H.-H., Lai, S.-O., Hsieh, L.-T., Hsueh, H.-J. & Chi, T.-W. Profiles of PAH emission from steel and iron industries. *Chemosphere* **48**, 1061–1074 (2002).
- 44. Hajat, A., Hsia, C. & O'Neill, M. S. Socioeconomic disparities and air pollution exposure: a global review. *Curr. Environ. Health Rep.* **2**, 440–450 (2015).
- 45. Kim, K. et al. Inequalities in urban greenness and epigenetic aging: diferent associations by race and neighborhood socioeconomic status. *Sci. Adv.* **9**, eadf8140 (2023).
- 46. Iafrate, A. J. et al. Detection of large-scale variation in the human genome. *Nat. Genet.* **36**, 949–951 (2004).
- 47. Sebat, J. et al. Large-scale copy number polymorphism in the human genome. *Science* **305**, 525–528 (2004).
- 48. Zhang, F., Gu, W., Hurles, M. E. & Lupski, J. R. Copy number variation in human health, disease, and evolution. *Annu. Rev. Genomics Hum. Genet.* **10**, 451–481 (2009).
- 49. Brown, T. A. *Genomes*, 2nd edn (Wiley-Liss, 2002).
- 50. Chu, D. & Wei, L. Nonsynonymous, synonymous and nonsense mutations in human cancer-related genes undergo stronger purifying selections than expectation. *BMC Cancer* **19**, 359 (2019).
- 51. Scacheri, C. A. & Scacheri, P. C. Mutations in the non-coding genome. *Curr. Opin. Pediatr.* **27**, 659–664 (2015).
- 52. Orr, H. A. Somatic mutation favors the evolution of diploidy. *Genetics* **139**, 1441–1447 (1995).
- 53. Otto, S. P. & Gerstein, A. C. The evolution of haploidy and diploidy. *Curr. Biol.* **18**, R1121–R1124 (2008).
- 54. Anderson, J. B. et al. Clonal evolution and genome stability in a 2500-year-old fungal individual. *Proc. R. Soc. B* **285**, 20182233 (2018).
- 55. Burian, A. Does shoot apical meristem function as the germline in safeguarding against excess of mutations? *Front. Plant Sci.* **12**, 707740 (2021).
- 56. Wang, Y. & Obbard, D. J. Experimental estimates of germline mutation rate in eukaryotes: a phylogenetic meta-analysis. *Evol. Lett.* **7**, 216–226 (2023).
- 57. Otto, S. P. The evolutionary enigma of sex. *Am. Nat.* **174**, S1–S14 (2009).
- 58. Charlesworth, B. The efects of deleterious mutations on evolution at linked sites. *Genetics* **190**, 5–22 (2012).
- 59. Charlesworth, B. Efective population size and patterns of molecular evolution and variation. *Nat. Rev. Genet.* **10**, 195–205 (2009).
- 60. Lanfear, R. Do plants have a segregated germline? *PLoS Biol.* **16**, e2005439 (2018).
- 61. Hecht, S. S. Tobacco smoke carcinogens and lung cancer. *J. Natl Cancer Inst.* **91**, 1194–1210 (1999).
- 62. Foo, J. & Michor, F. Evolution of acquired resistance to anti-cancer therapy. *J. Theor. Biol.* **355**, 10–20 (2014).
- 63. Godschalk, R. W., Yauk, C. L., van Benthem, J., Douglas, G. R. & Marchetti, F. In utero exposure to genotoxicants leading to genetic mosaicism: an overlooked window of susceptibility in genetic toxicology testing? *Environ. Mol. Mutagen.* **61**, 55–65 (2020).
- 64. Whitham, T. G. & Slobodchikof, C. Evolution by individuals, plant– herbivore interactions, and mosaics of genetic variability: the adaptive significance of somatic mutations in plants. *Oecologia* **49**, 287–292 (1981).
- 65. Schumacher, B., Pothof, J., Vijg, J. & Hoeijmakers, J. H. The central role of DNA damage in the ageing process. *Nature* **592**, 695–703 (2021).
- 66. Li, C. & Williams, S. M. Human somatic variation: it's not just for cancer anymore. *Curr. Genet. Med. Rep.* **1**, 212–218 (2013).
- 67. Claxton, L. D., de, A., Umbuzeiro, G. & DeMarini, D. M. The *Salmonella* mutagenicity assay: the stethoscope of genetic toxicology for the 21st century. *Environ. Health Perspect.* **118**, 1515–1522 (2010).
- 68. Claxton, L. D., Matthews, P. P. & Warren, S. H. The genotoxicity of ambient outdoor air, a review: *Salmonella* mutagenicity. *Mutat. Res. Rev. Mutat. Res.* **567**, 347–399 (2004).
- 69. Chen, G. & White, P. A. The mutagenic hazards of aquatic sediments: a review. *Mutat. Res. Rev. Mutat. Res.* **567**, 151–225 (2004).
- 70. Olivier, M., Hussain, S. P., Caron de Fromentel, C., Hainaut, P. & Harris, C. C. TP53 mutation spectra and load: a tool for generating hypotheses on the etiology of cancer. *IARC Sci. Publ.* **157**, 247–270 (2004).
- 71. Alexandrov, L. B. et al. The repertoire of mutational signatures in human cancer. *Nature* **578**, 94–101 (2020).
- 72. Ferreira, M. I., Domingos, M., Gomes, H. de A., Saldiva, P. H. & De Assuncao, J. V. Evaluation of mutagenic potential of contaminated atmosphere at Ibirapuera Park, São Paulo–SP, Brazil, using the *Tradescantia* stamen-hair assay. *Environ. Pollut.* **145**, 219–224 (2007).
- 73. DeMarini, D. M. et al. Lung tumor KRAS and TP53 mutations in nonsmokers reflect exposure to PAH-rich coal combustion emissions. *Cancer Res.* **61**, 6679–6681 (2001).
- 74. Yu, X.-J. et al. Characterization of somatic mutations in air pollution-related lung cancer. *eBioMedicine* **2**, 583–590 (2015).
- 75. Acito, M., Fatigoni, C., Villarini, M. & Moretti, M. Cytogenetic efects in children exposed to air pollutants: a systematic review and meta-analysis. *Int. J. Environ. Res. Public Health* **19**, 6736 (2022).
- 76. León-Mejía, G. et al. Cytotoxic and genotoxic effects in mechanics occupationally exposed to diesel engine exhaust. *Ecotoxicol. Environ. Saf.* **171**, 264–273 (2019).
- 77. Hansen, Å. M. et al. Urinary 1-hydroxypyrene and mutagenicity in bus drivers and mail carriers exposed to urban air pollution in Denmark. *Mutat. Res. Genet. Toxicol. Environ. Mutagen.* **557**, 7–17 (2004).
- 78. Wong, J. Y. et al. Elevated urinary mutagenicity among those exposed to bituminous coal combustion emissions or diesel engine exhaust. *Environ. Mol. Mutagen.* **62**, 458–470 (2021).
- 79. *Principles and Methods for the Risk Assessment of Chemicals in Food* Environmental Health Criteria No. 240 (FAO and WHO, 2020).
- 80. Eberwine, J., Sul, J.-Y., Bartfai, T. & Kim, J. The promise of single-cell sequencing. *Nat. Methods* **11**, 25–27 (2014).
- 81. Kennedy, S. R. et al. Detecting ultralow-frequency mutations by duplex sequencing. *Nat. Protoc.* **9**, 2586–2606 (2014).
- 82. Cho, E. et al. Error-corrected duplex sequencing enables direct detection and quantification of mutations in human TK6 cells with strong inter-laboratory consistency. *Mutat. Res. Genet. Toxicol. Environ. Mutagen.* **889**, 503649 (2023).
- 83. Shendure, J. & Akey, J. M. The origins, determinants, and consequences of human mutations. *Science* **349**, 1478–1483 (2015).
- 84. Ton, N. D. et al. Whole genome sequencing and mutation rate analysis of trios with paternal dioxin exposure. *Hum. Mutat.* **39**, 1384–1392 (2018).
- 85. Dubrova, Y. E. et al. Human minisatellite mutation rate after the Chernobyl accident. *Nature* **380**, 683–686 (1996).
- 86. Kovalchuk, I., Kovalchuk, O., Arkhipov, A. & Hohn, B. Transgenic plants are sensitive bioindicators of nuclear pollution caused by the Chernobyl accident. *Nat. Biotechnol.* **16**, 1054–1059 (1998).
- 87. Ellegren, H., Lindgren, G., Primmer, C. R. & Møller, A. P. Fitness loss and germline mutations in barn swallows breeding in Chernobyl. *Nature* **389**, 593–596 (1997).
- 88. Yeager, M. et al. Lack of transgenerational effects of ionizing radiation exposure from the Chernobyl accident. *Science* **372**, 725–729 (2021).
- 89. Kessler, M. D. et al. De novo mutations across 1,465 diverse genomes reveal mutational insights and reductions in the Amish founder population. *Proc. Natl Acad. Sci. USA* **117**, 2560–2569 (2020).
- 90. King, L., De Solla, S., Small, J., Sverko, E. & Quinn, J. Microsatellite DNA mutations in double-crested cormorants (*Phalacrocorax auritus*) associated with exposure to PAH-containing industrial air pollution. *Environ. Sci. Technol.* **48**, 11637–11645 (2014).
- 91. Somers, C. M., Yauk, C. L., White, P. A., Parfett, C. L. & Quinn, J. S. Air pollution induces heritable DNA mutations. *Proc. Natl Acad. Sci. USA* **99**, 15904–15907 (2002).
- 92. Ely, D. & Hamilton, B. *Trends in Fertility and Mother's Age at First Birth among Rural and Metropolitan Counties: United States, 2007–2017* NCHS Data Brief No. 323 (National Center for Health Statistics, Centers for Disease Control and Prevention, 2018).
- 93. Lerch, M. Fertility decline in urban and rural areas of developing countries. *Popul. Dev. Rev.* **45**, 301–320 (2019).
- 94. Goldmann, J. M. et al. Parent-of-origin-specific signatures of de novo mutations. *Nat. Genet.* **48**, 935–939 (2016).
- 95. Merckx, T. et al. Body-size shifts in aquatic and terrestrial urban communities. *Nature* **558**, 113–116 (2018).
- 96. Eyre-Walker, A. & Keightley, P. D. The distribution of fitness efects of new mutations. *Nat. Rev. Genet.* **8**, 610–618 (2007).
- 97. Schultz, S. T. & Lynch, M. Mutation and extinction: the role of variable mutational efects, synergistic epistasis, beneficial mutations, and degree of outcrossing. *Evolution* **51**, 1363–1371 (1997).
- 98. Sproufske, K., Aguilar-Rodriguez, J., Sniegowski, P. & Wagner, A. High mutation rates limit evolutionary adaptation in *Escherichia coli*. *PLoS Genet.* **14**, e1007324 (2018).
- 99. Keightley, P. D. Rates and fitness consequences of new mutations in humans. *Genetics* **190**, 295–304 (2012).
- 100.Pineda‐Krch, M. & Lehtilä, K. Costs and benefits of genetic heterogeneity within organisms. *J. Evol. Biol.* **17**, 1167–1177 (2004).
- 101. Doonan, J. H. & Sablowski, R. Walls around tumours—why plants do not develop cancer. *Nat. Rev. Cancer* **10**, 794–802 (2010).
- 102. Jiang, X. et al. Impacts of mutation efects and population size on mutation rate in asexual populations: a simulation study. *BMC Evol. Biol.* **10**, 298 (2010).
- 103. Sniegowski, P. D., Gerrish, P. J., Johnson, T. & Shaver, A. The evolution of mutation rates: separating causes from consequences. *Bioessays* **22**, 1057–1066 (2000).
- 104. Wei, W. et al. Rapid evolution of mutation rate and spectrum in response to environmental and population-genetic challenges. *Nat. Commun.* **13**, 4752 (2022).
- 105. Lynch, M. Evolution of the mutation rate. *Trends Genet.* **26**, 345–352 (2010).
- 106. Carlson, S. M., Cunningham, C. J. & Westley, P. A. Evolutionary rescue in a changing world. *Trends Ecol. Evol.* **29**, 521–530 (2014).
- 107. Metzgar, D. & Wills, C. Evidence for the adaptive evolution of mutation rates. *Cell* **101**, 581–584 (2000).
- 108. Thompson, K. A., Rieseberg, L. H. & Schluter, D. Speciation and the city. *Trends Ecol. Evol.* **33**, 815–826 (2018).
- 109. Orr, H. A. & Turelli, M. The evolution of postzygotic isolation: accumulating Dobzhansky–Muller incompatibilities. *Evolution* **55**, 1085–1094 (2001).
- 110. Van Drunen, W. E. & Johnson, M. T. J. Polyploidy in urban environments. *Trends Ecol. Evol.* **37**, 507–516 (2022).
- 111. Guo, H., Chang, Z., Wu, J. & Li, W. Air pollution and lung cancer incidence in China: who are faced with a greater efect? *Environ. Int.* **132**, 105077 (2019).
- 112. Dey, S. et al. Urban–rural diferences in breast cancer incidence in Egypt (1999–2006). *Breast* **19**, 417–423 (2010).
- 113. Ayuso-Álvarez, A. et al. Association between proximity to industrial chemical installations and cancer mortality in Spain. *Environ. Pollut.* **260**, 113869 (2020).
- 114. Giraudeau, M., Sepp, T., Ujvari, B., Ewald, P. W. & Thomas, F. Human activities might influence oncogenic processes in wild animal populations. *Nat. Ecol. Evol.* **2**, 1065–1070 (2018).
- 115. Sepp, T., Ujvari, B., Ewald, P. W., Thomas, F. & Giraudeau, M. Urban environment and cancer in wildlife: available evidence and future research avenues. *Proc. R. Soc. B* **286**, 20182434 (2019).
- 116. Baines, C. et al. Linking pollution and cancer in aquatic environments: a review. *Environ. Int.* **149**, 106391 (2021).
- 117. Mulvihill, J. J. Preconception exposure to mutagens: medical and other exposures to radiation and chemicals. *J. Community Genet.* **3**, 205–211 (2012).
- 118. Wang, L. et al. Association of ultra-processed food consumption with colorectal cancer risk among men and women: results from three prospective US cohort studies. *BMJ* **378**, e068921 (2022).
- 119. Gámez, S. et al. Downtown diet: a global meta-analysis of increased urbanization on the diets of vertebrate predators. *Proc. R. Soc. B* **289**, 20212487 (2022).
- 120. Lyons, J., Mastromonaco, G., Edwards, D. B. & Schulte-Hostedde, A. I. Fat and happy in the city: eastern chipmunks in urban environments. *Behav. Ecol.* **28**, 1464–1471 (2017).
- 121. Schulte-Hostedde, A. I., Mazal, Z., Jardine, C. M. & Gagnon, J. Enhanced access to anthropogenic food waste is related to hyperglycemia in raccoons (*Procyon lotor*). *Conserv. Physiol.* **6**, coy026 (2018).
- 122. Kliemann, N. et al. Ultra-processed foods and cancer risk: from global food systems to individual exposures and mechanisms. *Br. J. Cancer* **127**, 14–20 (2022).
- 123. Winglee, K. et al. Recent urbanization in China is correlated with a Westernized microbiome encoding increased virulence and antibiotic resistance genes. *Microbiome* **5**, 121 (2017).
- 124. Schell, C. J. et al. The ecological and evolutionary consequences of systemic racism in urban environments. *Science* **369**, eaay4497 (2020).
- 125. Des Roches, S. et al. Socio‐eco‐evolutionary dynamics in cities. *Evol. Appl.* **14**, 248–267 (2020).
- 126. Valentine, C. C. III et al. Direct quantification of in vivo mutagenesis and carcinogenesis using duplex sequencing. *Proc. Natl Acad. Sci. USA* **117**, 33414–33425 (2020).
- 127. Marchetti, F. et al. Error-corrected next-generation sequencing to advance nonclinical genotoxicity and carcinogenicity testing. *Nat. Rev. Drug Discov.* **22**, 165–166 (2023).
- 128. Frazer, J. et al. Disease variant prediction with deep generative models of evolutionary data. *Nature* **599**, 91–95 (2021).
- 129. Salk, J. J. & Kennedy, S. R. Next‐generation genotoxicology: using modern sequencing technologies to assess somatic mutagenesis and cancer risk. *Environ. Mol. Mutagen.* **61**, 135–151 (2020).
- 130. Du Four, V., Janssen, C., Brits, E. & Van Larebeke, N. Genotoxic and mutagenic activity of environmental air samples from diferent rural, urban and industrial sites in Flanders, Belgium. *Mutat. Res. Genet. Toxicol. Environ. Mutagen.* **588**, 106–117 (2005).
- 131. Ceretti, E. et al. Monitoring of volatile and non-volatile urban air genotoxins using bacteria, human cells and plants. *Chemosphere* **120**, 221–229 (2015).
- 132. David, E. & Niculescu, V.-C. Volatile organic compounds (VOCs) as environmental pollutants: occurrence and mitigation using nanomaterials. *Int. J. Environ. Res. Public Health* **18**, 13147 (2021).
- 133. Jameson, C. W. in *Tumour Site Concordance and Mechanisms of Carcinogenesis* (eds Baan, R. A. e al.) Ch. 7 (WHO Press, 2021).
- 134. Ravindra, K., Sokhi, R. & Van Grieken, R. Atmospheric polycyclic aromatic hydrocarbons: source attribution, emission factors and regulation. *Atmos. Environ.* **42**, 2895–2921 (2008).
- 135. Abdel-Shafy, H. I. & Mansour, M. S. A review on polycyclic aromatic hydrocarbons: source, environmental impact, efect on human health and remediation. *Egypt. J. Pet.* **25**, 107–123 (2016).
- 136. Levy, R. J. Carbon monoxide pollution and neurodevelopment: a public health concern. *Neurotoxicol. Teratol.* **49**, 31–40 (2015).
- 137. Brook, J. R. et al. Further interpretation of the acute efect of nitrogen dioxide observed in Canadian time-series studies. *J. Expo. Sci. Environ. Epidemiol.* **17**, S36–S44 (2007).
- 138. Zhang, L. et al. Understanding the industrial NOx and $SO₂$ pollutant emissions in China from sector linkage perspective. *Sci. Total Environ.* **770**, 145242 (2021).
- 139. Meftaul, I. M., Venkateswarlu, K., Dharmarajan, R., Annamalai, P. & Megharaj, M. Pesticides in the urban environment: a potential threat that knocks at the door. *Sci. Total Environ.* **711**, 134612 (2020).
- 140. Li, Z., Liang, Y., Zhou, J. & Sun, X. Impacts of de-icing salt pollution on urban road greenspace: a case study of Beijing. *Front. Environ. Sci. Eng.* **8**, 747–756 (2014).
- 141. García-Pérez, J., Gómez-Barroso, D., Tamayo-Uria, I. & Ramis, R. Methodological approaches to the study of cancer risk in the vicinity of pollution sources: the experience of a population-based case–control study of childhood cancer. *Int. J. Health Geogr.* **18**, 12 (2019).
- 142. García-Pérez, J. et al. Childhood leukemia and residential proximity to industrial and urban sites. *Environ. Res.* **140**, 542–553 (2015).
- 143. García-Pérez, J. et al. Association between residential proximity to environmental pollution sources and childhood renal tumors. *Environ. Res.* **147**, 405–414 (2016).
- 144. Chen, X. et al. Long-term exposure to urban air pollution and lung cancer mortality: a 12-year cohort study in Northern China. *Sci. Total Environ.* **571**, 855–861 (2016).
- 145. Beeson, W. L., Abbey, D. E. & Knutsen, S. F. Long-term concentrations of ambient air pollutants and incident lung cancer in California adults: results from the AHSMOG study. *Environ. Health Perspect.* **106**, 813–823 (1998).
- 146. Bai, X. et al. Linking urbanization and the environment: conceptual and empirical advances. *Annu. Rev. Environ. Resour.* **42**, 215–240 (2017).
- 147. Gogna, P. et al. Estimates of the current and future burden of lung cancer attributable to PM2.5 in Canada. *Prev. Med.* **122**, 91–99 (2019).
- 148. Nyberg, F. et al. Urban air pollution and lung cancer in Stockholm. *Epidemiology* **11**, 487–495 (2000).
- 149. Fei, X. et al. The association between heavy metal soil pollution and stomach cancer: a case study in Hangzhou City, China. *Environ. Geochem. Health* **40**, 2481–2490 (2018).
- 150. Cheng, I. et al. Association between ambient air pollution and breast cancer risk: the multiethnic cohort study. *Int. J. Cancer* **146**, 699–711 (2020).
- 151. Ebenstein, A. The consequences of industrialization: evidence from water pollution and digestive cancers in China. *Rev. Econ. Stat.* **94**, 186–201 (2012).
- 152. Wei, J. & Zhanqing, L. GlobalHighPM2.5: big data gapless 1km global ground-level PM2.5 dataset over land [Data set]. *Zenodo* <https://zenodo.org/records/10081359> (2022).
- 153. Wei, J. et al. First close insight into global daily gapless 1 $km PM_{2.5}$ pollution, driving factors, and health impact. *Nat. Commun.* **14**, 8349 (2023).
- 154. Center for International Earth Science Information Network *Annual PM2.5 Concentrations for Countries and Urban Areas, 1998–2016* (Columbia Univ., 2021).
- 155. Wolf, M. J. et al. *Country Trends in Major Air Pollutants, v1 (2003–2018)* (Socioeconomic Data and Applications Center, 2022).
- 156. Wolf, M. J. et al. New insights for tracking global and local trends in exposure to air pollutants. *Environ. Sci. Technol.* **56**, 3984–3996 (2022).
- 157. Figueroa, X. F., Lillo, M. A., Gaete, P. S., Riquelme, M. A. & Sáez, J. C. Difusion of nitric oxide across cell membranes of the vascular wall requires specific connexin-based channels. *Neuropharmacology* **75**, 471–478 (2013).
- 158. Su, R., Jin, X., Li, H., Huang, L. & Li, Z. The mechanisms of $PM_{2.5}$ and its main components penetrate into HUVEC cells and efects on cell organelles. *Chemosphere* **241**, 125127 (2020).
- 159. Yauk, C., Lambert, I., Marchetti, F. & Douglas, G. *Adverse Outcome Pathway on Alkylation of DNA in Male Pre-meiotic Germ Cells Leading to Heritable Mutations* (OECD, 2016).
- 160. Cho, E. et al. AOP report: development of an adverse outcome pathway for oxidative DNA damage leading to mutations and chromosomal aberrations. *Environ. Mol. Mutagen.* **63**, 118–134 (2022).
- 161. Lakey, P. S. et al. Chemical exposure–response relationship between air pollutants and reactive oxygen species in the human respiratory tract. *Sci. Rep.* **6**, 32916 (2016).
- 162. Sasaki, J. C. et al. Application of the adverse outcome pathway framework to genotoxic modes of action. *Environ. Mol. Mutagen.* **61**, 114–134 (2020).
- 163. Fucic, A. et al. Genomic damage in children accidentally exposed to ionizing radiation: a review of the literature. *Mutat. Res. Rev. Mutat. Res.* **658**, 111–123 (2008).
- 164. Chauhan, V., Sherman, S., Said, Z., Yauk, C. L. & Stainforth, R. A case example of a radiation-relevant adverse outcome pathway to lung cancer. *Int. J. Radiat. Biol.* **97**, 68–84 (2021).
- 165. Ignatov, A. V., Bondarenko, K. & Makarova, A. Non-bulky lesions in human DNA: the ways of formation, repair, and replication. *Acta Nat.* **9**, 12–26 (2017).

Acknowledgements

The ideas for this Perspective were developed over several workshops and meetings, including the Urban Eco-Evo Research Coordination Network (NSF DEB-184063), the 'Satellite Workshop on Urban Evolutionary and Ecological "Omics"' funded by the Society of Molecular Biology and Evolution, and the Center for Biological Data Science at Virginia Commonwealth University. M.T.J.J. was supported by an NSERC Steacie Fellowship, a Canada Research Chair and an NSERC Discovery Grant. C.L.Y. was supported by a Canada Research Chair and, along with F.M., a Burroughs Wellcome Fund Innovations in Regulatory Sciences Award. D.N.A. was supported by a Plant Resilience Institute Fellowship from Michigan State University. E.J.C. was funded by NSF DBI-2109587 and the Living Earth Collaborative at Washington University in St. Louis. M.P.-R. (DEB-2332998) and W.B. (DEB-1754394) received funding from the National Science Foundation. C.G.L. was funded by ANID PIA/BASAL FB0002. J.G. was supported by grant PID2020-115874GB-I00 funded by MCIN/ AEI/10.13039/501100011033 and by grant 2021 SGR 00417 funded by Departament de Recerca i Universitats, Generalitat de Catalunya. M.S. was supported by NCN Opus grant 2021/41/B/NZ8/04472.

Author contributions

M.T.J.J. conceived and led the project. All authors (M.T.J.J., I.A., F.M., J.M.-S., R.W.N., M.S., B.C.V., C.L.Y., D.N.A., W.B., A.E.C., E.J.C., A.D., J.G., C.G.L., M.O., M.P.-R., D.J.R., M.S.R. and K.M.W.) contributed to brainstorming the ideas covered in the paper, and the original outline was written by the lead team members (F.M., J.M.-S., R.W.N., M.S., B.C.V. and C.L.Y.). I.A., M.T.J.J., F.M., J.M.-S., R.W.N., M.S., B.C.V., K.M.W. and C.L.Y. led the writing of specific sections and/or the preparation of the figures and tables. All remaining authors (D.N.A., W.B., A.E.C., E.J.C., A.D., J.G., C.G.L., M.O., M.P.-R., D.J.R., M.S.R. and K.M.W.) contributed to one or more sections, and all authors (M.T.J.J., I.A., F.M., J.M.-S., R.W.N., M.S., B.C.V., C.L.Y., D.N.A., W.B., A.E.C., E.J.C., A.D., J.G., C.G.L., M.O., M.P.-R., D.J.R., M.S.R. and K.M.W.) edited the final drafts of the paper.

Competing interests

The authors declare no competing interests.

Additional information

Correspondence should be addressed to Marc T. J. Johnson.

Peer review information *Nature Ecology & Evolution* thanks Matthew Nielsen and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Springer Nature or its licensor (e.g. a society or other partner) holds exclusive rights to this article under a publishing agreement with the author(s) or other rightsholder(s); author self-archiving of the accepted manuscript version of this article is solely governed by the terms of such publishing agreement and applicable law.

© Springer Nature Limited 2024

¹Centre for Urban Environments, University of Toronto Mississauga, Mississauga, Ontario, Canada. ²Department of Biology, University of Toronto Mississauga, Mississauga, Ontario, Canada. ³Environmental Health Science and Research Bureau, Health Canada, Ottawa, Ontario, Canada. ⁴Department of Biology and Louis Calder Center, Fordham University, Armonk, NY, USA. ⁵Institute of Evolutionary Biology, Faculty of Biology, Biological and Chemical Research Centre, University of Warsaw, Warsaw, Poland. ⁶Center for Biological Data Science, Virginia Commonwealth University, Richmond, VA, USA. ⁷Department of Biology, University of Ottawa, Ottawa, Ontario, Canada. ⁸Department of Plant Biology, Department of Entomology, Plant Resilience

Institute, Michigan State University, East Lansing, MI, USA. ⁹Department of Entomology, Virginia Polytechnic Institute and State University, Blacksburg, VA, USA. ¹⁰Living Earth Collaborative, Washington University in St. Louis, St. Louis, MO, USA. ¹¹Department of Ecology and Evolutionary Biology, University of Arizona, Tucson, AZ, USA. ¹²Institute of Evolutionary Biology, CSIC, UPF, Barcelona, Spain. ¹³Departamento de Ciencias, Facultad de Artes Liberales, Universidad Adolfo Ibáñez, Santiago, Chile. ¹⁴Center of Applied Ecology and Sustainability (CAPES), Santiago, Chile. ¹⁵Department of Biology, Drexel University, Philadelphia, PA, USA. ¹⁶School of Biological Sciences, University of California, San Diego, La Jolla, CA, USA. ¹⁷Department of Biology, New York University, New York, NY, USA. \boxtimes e-mail: marc.johnson@utoronto.ca