- A mass budget and box model of global plastics cycling, degradation and dispersal in the land-ocean atmosphere system
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18 List of Abbreviations

- 20 Tg, teragrams
- 21 P, macroplastics
- 22 LMP, large microplastics
- 23 SMP, small microplastics
- 24 y, year
- 25 F, flux (in Tg y^{-1})
- 26 M, mass (in Tg)
- 27 1σ , one (sigma) standard deviation
- 28 BAU, business as usual
- 29 SCS, systems change scenario
- 30

31 Abstract

- Since 1950 humans have introduced 8300 teragrams (Tg, 10¹² grams, millions of metric tons) of plastic 32 polymers into the Earth's surface environment. Accounting for the dispersal and fate of produced plastics and 33 fragmented microplastics in the environment has been challenging. Recent studies have fueled debate on the 34 global river budget for plastic transport to oceans, the sinking and beaching of marine plastics and the emission 35 and deposition of atmospheric microplastics. Here we define a global plastics cycle and budget, and develop 36 a box model of plastic cycling, including the fragmentation and transport of large and small microplastics 37 (LMP, SMP) within coupled terrestrial, oceanic and atmospheric reservoirs. We force the model with 38 historical plastics production and waste data, and explore how macroplastics, LMP and SMP propagate 39 through the reservoirs from 1950 to 2015 and beyond. We find that considerable amounts of plastics reside 40 most likely in the deep ocean (82 Tg), in shelf sediments (116 Tg), on beaches (1.8 Tg) and, as a result of 41 marine emissions, in the remote terrestrial surface pool (28 Tg). Business as usual or maximum feasible 42 reduction and discard scenarios show similar, 4-fold increases in atmospheric and aquatic ecosystem SMP 43 exposure by 2050, because future plastics mobilization is controlled by releases from the large terrestrial 44 45 discarded plastics reservoir (3500 Tg). Zero-release from 2025 onwards illustrates recovery of P and LMP reservoirs on centennial time scales, while SMP continue to cycle in air, soil, and surface ocean for millennia. 46 Limiting dramatic future dispersal of plastics requires, in addition to reducing use and waste, remediation of 47 the large terrestrial legacy plastics pool. 48
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50 Introduction

51 A characteristic feature of the Anthropocene is the widespread dispersal of plastic polymers across Earth's

- 52 surface since the 1950s (1). Of the 1.5 trillion barrels of oil (200,000 Tg) produced since the 1950s (2) about
- 53 4% (8300 Tg) has been transformed into non-biodegradable polymers, and used in predominantly single-use
- 54 packaging or short-lived (1-25y) technological applications (3). Produced plastics have been abundantly

(60%) discarded into the technosphere, the part of the environment that has been made or modified by humans: 55 urban, sub-urban, agricultural, and industrial areas, including landfills (3,4). The pool of discarded managed 56 and mismanaged plastic waste has been slowly mobilized by wind, runoff, rivers and ocean currents to all 57 remote corners of planet Earth, including the poles and the deep ocean (5–8). Large plastic debris tend to 58 fragment to micro- and nano-sized particles, which due to their increased surface area can absorb, adsorb or 59 60 release a range of secondary natural and man-made chemical compounds in the environment (9). Assessing the possible impact of plastics on ecosystem and human health, and mitigating this impact, requires a solid 61 understanding of where and when discarded plastics end-up, and to which size range they evolve. 62

Over the past decades important efforts have been made to chart the abundance, size properties, and 63 64 bulk polymer composition of plastics in the surface ocean, soils, rivers, wetlands, biota and atmosphere. A perceived mismatch between the relatively small quantity of plastics in the surface ocean (0.3 Tg) (10) and 65 the large quantity delivered by global rivers $(4.8 - 12.7 \text{ Tg y}^{-1})$ (11) has fueled a 'missing marine plastics' 66 paradox (12). Solutions to this issue have been proposed in the transfer of marine plastics to the deep ocean 67 (13,14), to coastal environments, via beaching (15,16), to the subsurface ocean and marine sediments by 68 sinking (17,18), and to marine emission of microplastics to the atmosphere (19,20). Recently, a 1000-fold 69 lower global river flux of 0.0064 Tg y⁻¹ was suggested, based on an alternative median plastics mass (21). 70 Such a low river flux would imply a marine residence time of several years, and possibly removes the need 71 for a missing marine plastics sink. In parallel to rivers and ocean currents, the atmosphere has been identified 72 as a global vector of MP, in both urban (22,23) and remote environments (5,24,25), including MP emission 73 from land (19,26) and sea (19,20). In this study we use the best available estimates of both plastics abundance 74 and fluxes to construct a global plastics mass budget. This budget is implemented in a global box model of 75 plastics cycling between land, atmosphere and ocean from 1950 to 2015. We then use the model to explore 76 how plastics disperse through Earth's surface environment over times scale ranging from decades, focusing 77 on policy scenarios, to millennia, addressing the fate and potential burden of global plastics contamination. 78 79

80 Plastics cycling box model

In order to construct a global plastics mass and mass transfer budget (Figure 1), we use plastics observations 81 from the literature and a box modeling approach (see Methods for details). We subdivide macroplastics 82 83 (P, >5mm), large microplastics (LMP, >0.3mm and <5mm) and small microplastics (SMP, <0.3mm), and define 'MP' as the sum of LMP plus SMP. The GBM-Plastics model (version 1.0) is a coupled 15-reservoir 84 numerical box model that simulates how produced P and MP propagate through the terrestrial, marine, and 85 atmospheric environments upon release or emission. P fragment to LMP, and LMP fragment to SMP, and 86 only SMP become airborne, emitted from and deposited to oceans and land. Terminal P, LMP and SMP sinks 87 are marine sediments, whereas remote terrestrial (soils, barren rock, ice sheets) and deep ocean pools act as 88 long-term temporary reservoirs. The mass flux, F_{ab} (Tg y⁻¹) between two reservoirs a and b is $F_{ab} = k_{ab} \times M_a$, 89 where M_a is the mass of plastics in reservoir a (Tg), and k_{ab} is a first-order mass transfer (rate) coefficient (y⁻ 90 ¹). In such a first-order mass transfer model, an increase in mass of reservoir 'a' leads to a proportional (by 91 'k_{ab}') increase of the flux, F_{ab}, to reservoir b, and a consequent proportional increase in the mass of reservoir 92 'b'. In order to derive the k's one must know, for a given year or reasonably short period of observation, the 93 magnitudes of F and M. Alternatively, some k's can be derived experimentally, such as plastics sedimentation 94 95 rates, with units y^{-1} , or from theory. Therefore, in a first step, we calculate most k values from published, recent, 2005-2022 observations (see Methods) and from model estimates of atmospheric SMP fluxes (19). 96 The model is then run from 1950 to 2015, with only the k_{ab} transfer coefficients and plastics production and 97 waste generation as external forcing. In the following we discuss whether the simulated modern plastics 98 distribution for 2015 corresponds to observations, which k values (and therefore fluxes) need to be adjusted, 99 and what the model implications are for our understanding of plastic cycling. With the addition of atmospheric 100 transport of plastics, the term 'emission' refers here exclusively to the suspension of terrestrial and marine 101 102 SMP in air. 'Release' is used as the generic term for plastics discharge and mobilization to the technosphere and in-land aquatic and marine environments. Conversion of plastics number concentration to mass 103 concentration is detailed in the Methods. All uncertainties reported are 1^o standard deviation (or 16th and 84th 104 percentiles, corresponding to a 1σ uncertainty; see Methods). 105

We start by detailing the 'base case' plastics cycling model, based on best known modern observations
of reservoir sizes and fluxes between reservoirs (see Methods for details). We include plastics production
(8300 Tg since 1950), waste generation and waste disposal from Geyer et al. (3) who estimated 2600 Tg of

plastics to be in use in 2015, 4900 Tg discarded (split into 4200 Tg of P, and 700 of primary LMP following 109 Lau et al. (4) and 800 Tg incinerated. In the base case we use the mid-point of the river plastics flux estimate 110 by Jambeck et al., of 8.5 Tg $y^{-1}(11)$, containing equal fractions of P and LMP. We adopt surface ocean mixed 111 layer buoyant P and LMP inventories of 0.23 and 0.04 Tg (10), and a surface mixed layer SMP inventory of 112 0.003 Tg (27). We make an order of magnitude estimate of beached LMP of 0.5 ± 0.4 Tg, based on the global 113 surface of sandy beaches (2.63 10⁵ km²; (28)), a median global beach sand LMP abundance of 2450 MP km⁻ 114 2 , and median beached LMP size of 2.0 mm (29). We estimate beached P, and shelf sediment P pools from a 115 review study (30) that estimates mean beached P and sea floor P concentrations of 2 and 5 Mg km⁻² 116 respectively (uncertainty not estimated). Multiplying by beach and continental shelf surfaces of 2.63 10⁵ and 117 118 2.89 10⁷ km² results in beached and shelf sediment P pools of 1.3 and 51 Tg. An estimate for the global deep ocean sediment MP pool of 1.5 Tg is based on observed mean deep sediment MP concentrations of 0.72 MP 119 g⁻¹ (see Methods) (31). A shelf sediment MP pool of 65 Tg (1σ , 21 to 78Tg) is estimated from subtidal 120 sediment median MP concentrations of 100 MP kg⁻¹ (see Methods) (29). Rate coefficients for P and LMP 121 beaching (the transfer from ocean to beach), k_{beaching} of 0.15 y⁻¹ are approximated based on Onink et al. (15). 122 Surface mixed layer to deep subsurface ocean sinking rates of P, LMP, SMP lack in situ observations; we 123 estimate model sinking rate coefficients, k_{P,sinking} of 1367 y⁻¹, k_{LMP,sinking} of 196 y⁻¹ and k_{SMP,sinking} of 33 y⁻¹ for 124 the 100 m deep surface ocean mixed layer, based on the empirical results of a sinking tank study of mixed 125 phytoplankton aggregates with MP(17). We include the sinking and sedimentation of non-buoyant P over the 126 shelf, but not from open ocean waters, assuming that only buoyant P dominate open ocean P. Macroplastics, 127 P, are beached as described above, and fragmented in surface ocean waters to LMP at a rate $k_{oceP \rightarrow LMP}$ of 0.03 128 y^{-1} (16), supported by observations (32). A recent review of plastics degradation rates highlights the 129 complexity and variability of plastics degradation rates as a function of polymer type, sunlight, and physical 130 environment (33). The authors use an observed median HDPE degradation rate of 4.3 μ m y⁻¹ in the marine 131 environment, and a theoretical degradation framework to illustrate how a typical HDEP bag (film), fiber (2 132 mm diameter, 230mm long) or bead (8.8 mm diameter) would degrade at relative mass loss rates of 0.5, 0.005 133 and 0.0014 y⁻¹. The rate of 0.03 y⁻¹ (1 σ uncertainty: 0.006 to 0.06 y⁻¹) we adopt lies within this estimated 134 variability. We consider that for the purpose of our study, it is too early at present to try and incorporate more 135 detailed plastics fragmentation or degradation parameterizations. We agree with Chamas et al. (33) that more 136 137 robust degradation observations are needed, and we suggest that a follow-up box model that incorporates variable polymer types would be a more appropriate occasion. In the absence of fragmentation rates for LMP 138 to SMP in surface, subsurface waters, beach zone, and discarded pool, and for P to LMP in subsurface water, 139 beach zone and discarded pool we adopt, in the base case, the same rate $k_{oceLMP \rightarrow SMP}$ of 0.03 y⁻¹ for all these 140 fragmentation sites. 141

The subsurface ocean pool of LMP and SMP, below the surface mixed layer, is of importance to complete the marine plastics budget and to parameterize model settling and sedimentation of plastics. Table 144 1 and Figure 2 summarize recent observations of subsurface marine MP. We estimate a global deep ocean MP 145 inventory of 82 ± 47 Tg based on mean N-Pacific pelagic concentrations of $131 \pm 44 \ \mu g \ m^{-3}$ (6,34), mean N 146 and S-Atlantic concentrations of $91 \pm 46 \ \mu g \ m^{-3}$ (35–37), and extrapolated estimates for the Indian, Southern, 147 and S-Pacific Oceans (Table 2, Methods).

Recent studies on atmospheric MP cycling show fragment and fiber size distributions to be in the SMP 148 range <300 µm. While LMP emission and deposition occurs, these tend to deposit more rapidly back to the 149 same reservoir (e.g., marine emission followed by marine deposition) and are therefore ignored in the box 150 model. Table 3 summarizes SMP observations in the boundary layer and free troposphere, yielding a total 151 tropospheric SMP mass of 0.031 ± 0.027 Tg. This observed stock is 10x higher, though within uncertainty, of 152 a model estimate of 0.0036 Tg (19). Our estimate of 0.031 Tg is very sensitive to the assumed median SMP 153 size of 70 µm, which is where the atmospheric SMP model allocates most SMP mass (19). We adopt global 154 SMP emissions from the same model study (19): emissions from roads, 0.1 Tg y⁻¹, agricultural dust, 0.07 Tg 155 y⁻¹, population dust, 0.02 Tg y⁻¹, and oceans, 8.6 Tg y⁻¹. We use SMP deposition observations over land 156 (5,24,38) in combination with population density data for 2015 (39) to estimate global SMP deposition over 157 land of 1.1 ± 0.3 Tg y⁻¹ and an accumulated remote terrestrial SMP pool of 28 ± 10 Tg (see Methods). We 158 assume that global SMP emissions (8.6 Tg y⁻¹; (19)) equal deposition, and estimate SMP deposition over 159 oceans as the difference between total deposition and deposition over land: 7.6 Tg y⁻¹. This large re-deposition 160 of SMP over the ocean is coherent with the short, 2.4 h, lifetime of the coarsest, 70 µm SMP size fraction that 161 represents 85% or marine SMP emission in the model of ref (40). 162

164 **Results and Discussion**

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The box model 'base case' is run from 1950 to 2015 and results, in terms of plastics reservoir sizes and fluxes 166 for the year 2015, are shown in Table 4 in comparison to the above-mentioned observations. The base case 167 reproduces observed amounts of in-use P, discarded P, LMP, SMP and terrestrial SMP to within 40%. Using 168 the mid-point river plastics flux of 8.5 Tg y⁻¹ (11) the base case also reproduces well the observed downstream 169 plastics mass in marine and remote terrestrial systems (surface and deep ocean, sediments, beach, remote 170 terrestrial surfaces) of 201 Tg (1σ , 120 to 630 Tg). Remote terrestrial surfaces are included in the downstream 171 172 environment, because its accumulated SMP mass is for 96% derived from the SMP river flux, surface ocean LMP degradation, and the important marine emission of SMP to the atmosphere where it leads to global 173 dispersal and deposition to remote terrestrial surfaces (soil, rock, deserts, ice). We note that using the 2000-174 fold lower river plastics flux of 0.0064 Tg y⁻¹ by Weiss et al. (21) would lead to important low bias in the 175 marine and remote terrestrial reservoirs. A model river plastics flux of 13 Tg y⁻¹ (1 σ , 9 to 51 Tg) balances the 176 overall marine plastics budget, and gives satisfactory (within a factor 10x) reproduction of surface ocean P. 177 LMP and SMP, shelf sediment P, LMP and SMP, and beached LMP reservoirs. Within the marine system, the 178 modeled deep sediment MP pool is however biased high 90-fold, and beached P biased low 26-fold, exceeding 179 the 10x uncertainty we apply to the observed pools. We therefore optimize and lower subsurface ocean specific 180 $k_{\text{LMP,sinking}}$ from 4.9 y⁻¹ to 0.0012 y⁻¹ and $k_{\text{SMP,sinking}}$ from 0.8 y⁻¹ to 0.0002 y⁻¹, and increase k_{beaching} from 0.15 181 y^{-1} to 4.0 y^{-1} . We argue that the base case sinking rates and k estimates for experimental biofouled LMP and 182 SMP are inappropriate for deep ocean sedimentation because remineralization of biofilm during sinking 183 increases buoyancy, halts sinking and lowers the effective sinking rate (14). The base case k_{beaching} was derived 184 for the coastal ocean (15), which we do not explicitly separate and simulate here, likely leading to its 185 underestimation relative to whole surface ocean P cycling. It is important to note that out of 23 mass transfer 186 coefficients (k's) only 2 needed fitting in the 'base case'. This indicates that current understanding of P, LMP 187 and SMP stocks and fluxes, which determine k's, is sufficiently accurate to formulate and use the box model. 188

Figure 1 presents our best estimate of the global plastics cycle for the year 2015, based on observed inventories and fluxes (black), modeled inventories and fluxes (red), including the modeled river plastics flux of 13 Tg y⁻¹ to the ocean (see Table 4 for uncertainties). Key properties of the global plastics cycle are:

The large mass, 1200 Tg of discarded LMP (of which 840 Tg primary LMP) and on the order of 540 Tg of
 discarded SMP in the technosphere, which are potentially mobilizable to wetlands, oceans, groundwater,
 atmosphere and remote terrestrial surfaces.

2. The substantial mass of plastics, 201 Tg, representing 3% of all plastics produced since 1950, that has been
released from the technosphere to pristine terrestrial and marine ecosystems.

197 3. The 65-fold larger river plastics flux (13 Tg y^{-1}) compared to the total terrestrial atmospheric SMP emission 198 flux (0.2 Tg y^{-1}) .

4. The importance of marine SMP emissions on further distributing microplastics to remote ocean waters and
to remote terrestrial surfaces (96% of the 28 Tg on remote land originates from marine emissions, and only
4% from terrestrial emissions).

5. The potentially large subsurface oceanic LMP and SMP (82 ± 47 Tg), and shelf sediment P and LMP (116 Tg) reservoirs, compared to beached P and LMP (1.8 Tg), and compared to surface ocean plastics (0.27 Tg).

The uncertainties associated with the global plastics cycle (Table 4) are large, due to an overall lack of observations and underlying plastics quantification challenges. In particular, observations of SMP number and mass in the terrestrial discarded and remote terrestrial pools, and in terrestrial and marine emissions and deposition are needed.

We use the box model to simulate and illustrate at what timescales P, LMP and SMP propagate through 208 Earth surface reservoirs if we were to halt plastics production and waste generation in 2025. While such a 209 210 scenario is not realistic, it serves to understand the timescales involved in plastics dispersal and degradation across the Earth's surface. Figure 3 shows P, LMP and SMP dispersal from 1950 to the year 3000 (see also SI 211 2 for model data output): The discarded terrestrial P pool decreases rapidly, by 90% in 2100, due to 212 fragmentation to LMP, which in turn decreases by 90% in 2150 due to further fragmentation to SMP. LMP 213 and SMP transport by rivers and air leads to rapid increases of LMP and SMP in the marine pools and of SMP 214 in the remote terrestrial pool. The discarded SMP pool takes longer, 90% by 2500, to mobilize to the surface 215 ocean, and from there via marine emission back to the remote terrestrial pool. The cyclical behavior that 216

develops, cycles SMP for millennia back and forth between surface ocean and continents, before gradual escape of SMP to the deep ocean marine sediments (Figure 4). This scenario illustrates that even if we would entirely replace plastics by alternative materials, the legacy of historical plastics mismanagement could result in prolonged plastics dispersal over centuries (LMP) or millennia (SMP), unless we better manage present and future discarded plastics pools on land. I should be noted that the relevance of persistent SMP cycling over millennial timescales will depend on their degradation to nanoplastics, and eventually to dissolved monomers that serve as carbon substrates to biological organisms.

Next, we explore in detail how two more realistic production and waste management scenarios affect 224 plastics cycling over the period 2015 to 2050 (Figure 5 and SI 2): 1. The business as usual (BAU) scenario 225 from Geyer et al. (3) reaching 30,000 Tg of produced plastics in 2050, and with discard below 10% and 226 recycling and incineration of 43% and 49% in 2050, 2. The systems change scenario (SCS) from Lau et al. 227 (4) which proposes ambitious, but realistic measures to reduce, substitute, recycle, and dispose of plastics (see 228 Methods for details, and SI 2). Figure 5A illustrates how a 2-fold drop in plastics production from 550 Tg y⁻¹ 229 to 250 Tg y⁻¹ in 2040 under the SCS scenario significantly limits further plastics accumulation in the 230 technosphere compared to BAU. Yet, despite the projected strong decrease of mismanaged waste, and increase 231 in safe disposal and recycling, the SCS does not lead to measurable changes in key metrics, such as beached 232 P, total river plastics flux (P+LMP+SMP) or atmospheric SMP deposition to remote terrestrial surfaces by 233 2050 (Figure 5 B, C, D). The reason for this is the persistent mobility of legacy plastic waste in the large 234 terrestrial discarded P, LMP and SMP reservoirs. To render SCS policy effective, it will have to be supported 235 by immobilization or remediation of the terrestrial discarded plastics pool. We explore the potential impact of 236 remediation of the discarded P pool from 2025 onwards at a rate of 3% P isolation and safe disposal per year 237 (Figure 5 B, C, D). Discarded P remediation halts beached P dispersal by 2040, curbs total river plastics 238 discharge to some extent but does not impact atmospheric SMP deposition to land. Although technically more 239 challenging, remediation of discarded LMP and SMP pools on land at an identical 3% per year rate is needed 240 to also inverse dispersal of river and atmospheric plastics (Figure 5 B, C, D) and to truly limit future planetary 241 dispersal of plastics. The current clean-up initiative of surface ocean plastics does not sufficiently address the 242 long-term mobilization of the legacy plastics pool on land. The fragmentation of SMP to nanoplastics and 243 ultimately to dissolved and colloidal polymers, that are energy sources to microbes needs further study, in 244 particular their rates of production, before they can be included in the box model. Engineered LMP and SMP 245 biodegradation could be a solution to the suggested need for remediation of these legacy pools on land. 246

248 Conclusions

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In this study we define a global plastics cycling budget for the year 2015, and develop a box model of plastic 249 cycling, including the transport and fragmentation of macroplastics (P) to large (LMP) and to small 250 microplastics (SMP) within coupled terrestrial, oceanic and atmospheric reservoirs. We drive the model with 251 historical plastics production and waste data, and investigate how macroplastics (P), LMP and SMP propagate 252 through Earth surface reservoirs from 1950 to 2015 and beyond, to 2050 and to the year 3000. Based on 253 published plastics observations we estimate that important amounts of plastics are present in the deep ocean 254 (82 Tg), in shelf sediments (116 Tg), on beaches (1.8 Tg) and in the remote terrestrial surface pool (28 Tg). 255 The box model suggests that plastics in the remote terrestrial surface pool originate predominantly from 256 marine SMP emissions that are transported via the atmosphere and deposited over land. Simulated zero-release 257 of plastics to land, water and air from 2025 onwards illustrates how P and LMP reservoirs recover on 258 centennial time scales, while SMP continue to cycle in air, soil, and surface ocean for millennia. Business as 259 usual or maximum feasible reduction and discard scenarios show similar, 4-fold increases in atmospheric and 260 aquatic ecosystem SMP exposure by 2050, because future plastics mobilization is controlled by releases from 261 the large terrestrial discarded plastics reservoir. We conclude that in order to limit future dispersal of plastics 262 we should, in addition to reducing plastics use and waste, anticipate remediation of the large terrestrial legacy 263 264 plastics pool.

266 Methods

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The GBM-Plastics-v1.0 model (global box model for plastics, version 1.0) code is included in SI 3 as Python 267 scripts, and in SI 2 in a Microsoft© Excel© version. It is also available via 268 https://github.com/AlkuinKoenig/GBM-Plastics. Definitions of plastics size categories are continuously 269 debated; here we use operational definitions of macroplastics (P, >5mm), large microplastics (LMP, >0.3mm) 270

and <5mm) and small microplastics (SMP, <0.3mm). The 0.3mm distinction is based on the frequently used 271 plankton net mesh size of approximately 0.3 mm. The 0.3mm cut-off is also a reasonable starting point for the 272 simulation of atmospheric cycling of SMP, with nearly all remote airborne SMP particles, films and 50% of 273 fibers falling in the 1-300 µm range (5,24). All P, LMP, SMP reservoir sizes (i.e., inventory) and fluxes are 274 expressed in teragrams (Tg = 10^{12} grams) and Tg y⁻¹. For some reservoirs, studies do not discern LMP or 275 SMP, in which case we retain the generic 'MP' abbreviation. 276

LMP and SMP observations are typically expressed as MP counts per unit volume or per unit area. To 277 estimate mass concentrations, we use, whenever reported, the full MP size distribution reported, a uniform 278 density of 1 x 10⁻⁶ μ g μ m⁻³ (41), and the MP volume approximation, V = L³ x 0.1, where L are the reported 279 length values of the size distribution. 280

We use global plastics production, 8300 Tg (teragrams or millions of metric tons), and waste 281 generation (discarded, recycled or incinerated) from Geyer et al. (3). Produced plastics enter the 'in-use' pool, 282 where they are mostly discarded within a single year due to the dominant use of single-use packaging. In 2015, 283 55% of non-fiber plastics are still discarded within a year, 25% incinerated and 20% recycled (3). We assume 284 fiber plastics to undergo similar relative discarding and incineration fates, leading to a 'discarded P+MP' 285 reservoir of 4900 Tg, an incinerated pool of 800 Tg (atmospheric CO₂) and an in-use pool of 2600 Tg in 2015 286 as described by Geyer et al.(3). Lau et al. (4) estimated the proportion of municipal solid waste that enters 287 aquatic and terrestrial environments as primary LMP to be 14 ± 4 % in 2016, which we apply here to all 288 discarded plastics (4). We therefore apply a primary f_{LMP} fraction of 0.14 and primary f_P fraction of 0.86 to 289 estimate transfer from the in-use to discarded reservoir for the period 2050-2015. The following mass balance 290 equations are defined for in-use and discarded pools: 291

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$$\frac{d(P_{use})}{dt} = P_{prod} - f_{disc} \times P_{waste} - f_{inc} \times P_{waste} \quad (\text{Eq.1})$$
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Where P_{use} is the mass of total plastic (P + LMP) in use, P_{prod} the mass of total plastics produced (Tg y⁻¹), 295 Pwaste the mass of total plastic waste, and fdisc, and finc are the fractions of Puse that are discarded, incinerated 296 and recycled. 297

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$$\frac{d(P_{disc})}{dt} = f_{disc} \times P_{waste} \times f_P - k_{P-river} \times P_{disc} - k_{discP \to LMP} \times P_{disc} \quad (Eq.2)$$
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Where P_{disc} is the mass of P discarded, f_P is the fraction of total plastic waste that are macroplastics, k_{P-river} is 301 the transfer coefficient for P to the ocean, via river runoff. 302

$$304 \quad \frac{d(LMP_{disc})}{dt} = f_{disc} \times P_{waste} \times f_{LMP} + k_{discP \to LMP} \times P_{disc} - k_{LMP-river} \times LMP_{disc} - k_{discLMP \to sMP} \times 305 \quad LMP_{disc} \text{ (Eq.3)}$$

Where LMP_{disc} is the mass of LMP discarded, f_{LMP} is the fraction of total plastics waste that are primary 307 microplastics (pellets, synthetic textiles, personal care products, etc), k_{LMP-river} is the transfer coefficient for 308 LMP to the ocean, via river runoff, and $k_{LMP \rightarrow SMP}$ is the transfer coefficient for LMP degradation to SMP 309 within the terrestrial 'discarded' pool. 310

$$\frac{d(SMP_{disc})}{dt} = k_{discLMP \to SMP} \times SMP_{disc} - k_{SMP-river} \times SMP_{disc} - k_{disc-atm} \times SMP_{disc}$$
(Eq.4)

Where SMP_{disc} is the mass of SMP discarded, k_{SMP-river} is the transfer coefficient for SMP to the ocean, via 314 river runoff, and k_{SMP-atm} is the transfer coefficient for SMP emission to the atmosphere from the terrestrial 315 'discarded' pool, including tire wear particles (TWP). 316

Transfer coefficients k_{P-river}, k_{LMP-river}, and k_{SMP-river} are calculated from 2015 plastic fluxes and 317 inventories, e.g. k_{P-river}=P_{disc}/F_{P-river} where F stands for flux (SI 1, Table S1). The mid-point estimate for F_{P-river} 318 of 8.8 Tg y⁻¹ ((11,42)) is used here, and subdivided into 50% P and 50% LMP (21). The 'discarded pool to 319 atmosphere' transfer coefficient, k_{disc-atm}, which theoretically equals SMP_{disc}/F_{SMP disc-atm} is unconstrained, 320 because the SMP_{disc} pool size, in Tg, is unknown (F_{SMP_disc-atm} is 0.18 Tg y⁻¹, based on Brahney et al. (19), and 321 was therefore fitted at 0.00037 y^{-1} as described in the text. 322

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324 The global ocean. Two previous box models have examined the plastics budget of the marine environment (13,16). In addition, a number of Lagrangian oceanic or atmospheric transport models have provided insight 325 in marine plastics dispersal and surface ocean plastics mass balance (15,19,43). Koelmans et al. (13) used a 326 plastics mass budget for the surface ocean to fit a marine P to LMP fragmentation rate, and a LMP 327 sedimentation rate, under the assumption of 100% buoyant P (no settling to deep waters). To accommodate 328 the high river plastic inputs, rapid plastic fragmentation to LMP (>90% per year), and rapid LMP settling rates 329 were fitted, and suggested a short plastics and LMP residence time for the surface ocean (<3 yrs). Subsequent 330 modeling work has investigated P and LMP beaching, resuspension in coastal waters (15,16), marine SMP 331 emissions(19), and P sedimentation due to loss of buyuancy(16). Lebreton et al. (16), in their marine box 332 model study(16), argued that observations of old plastics in the surface ocean disagree with rapid 333 fragmentation and settling and fitted a plastics to LMP degradation rate of 3% per year, which we adopt here 334 for the surface mixed layer ($k_{Poce \rightarrow LMP} = 0.03 \text{ y}^{-1}$). 335

Lebreton et al. (16) fitted important beaching of coastal plastics (97% per year). In the absence of a robust estimate for global beached macroplastics (44), Onink et al. (15) recently analyzed model beaching and resuspension scenarios finding at least 77% of net beaching for positively buoyant plastic debris over 5 years (15), which we adopt here in the base case as $k_{P,beaching} = 0.15 \text{ y}^{-1}$. Surface ocean P, LMP, and SMP equations are:

$$\frac{d(P_{surf-oce})}{dt} = k_{P-river} \times P_{disc} - k_{Psurf-oce-beach} \times P_{oce} - k_{Psurf-oce \rightarrow LMP} \times P_{surf-oce} - k_{Psurf-oce \rightarrow sed} \times P_{surf-oce} \times f_{shelf}$$
(Eq.5)
344

$$\frac{d(LMP_{surf-oce})}{dt} = k_{LMP-river} \times LMP_{disc} + k_{Psurf-oce \rightarrow LMP} \times P_{oce} - k_{LMPsurf-oce \rightarrow beach} \times LMP_{surf-oce} - k_{LMPsurf-oce \rightarrow shelfsed} \times LMP_{surf-oce} \times f_{shelf} - k_{LMP-sink} \times LMP_{surf-oce} \times f_{pelagic} - k_{LMPsurf-oce \rightarrow SMP} \times LMP_{surf-oce} \quad (Eq.6)$$

$$349 \quad \frac{d(SMP_{surf-oce})}{dt} = k_{SMP-river} \times SMP_{disc} + k_{atm \to oce} \times SMP_{atm} + k_{terr \to oce} \times SMP_{terr} +
350 \quad k_{LMPsurf-oce \to SMP} \times LMP_{surf-oce} - k_{oce \to atm} \times SMP_{surf-oce} - k_{SMPsurf-oce \to sed} \times SMP_{surf-oce} \times
351 \quad f_{shelf} - k_{SMP-sink} \times SMP_{surf-oce} \times f_{pelagic} (Eq.7)
352
353 \quad \frac{d(P_{shelf-sed})}{dt} = k_{Psurf-oce \to sed} \times P_{surf-oce} \times f_{shelf} (Eq.8)$$

354
355
$$\frac{d(LMP_{shelf-sed})}{dt} = k_{LMPsurf-oce \rightarrow sed} \times LMP_{surf-oce} \times f_{shelf} \text{ (Eq.9)}$$
356

357
$$\frac{d(SMP_{shelf-sed})}{dt} = k_{SMPsurf-oce \rightarrow sed} \times SMP_{surf-oce} \times f_{shelf}$$
(Eq.10)

358 359

362

Where $f_{shelf} = 0.08$, is the fraction of global continental shelf surface area, and $f_{pelagic}$ is the fraction of open ocean surface area. Subsurface ocean equations are:

$$\begin{array}{l} 363 \quad \frac{d(LMP_{deep-oce})}{dt} = k_{LMP-sink} \times LMP_{surf-oce} \times f_{pelagic} - k_{LMP \rightarrow SMP} \times LMP_{deep-oce} - k_{LMPdeep \rightarrow deepsed} \times \\ 364 \quad LMP_{deep-oce} \quad (Eq.11) \\ 365 \\ 366 \quad \frac{d(SMP_{deep-oce})}{dt} = k_{SMP-sink} \times SMP_{surf-oce} \times f_{pelagic} + k_{LMP \rightarrow SMP} \times LMP_{deep-oce} - k_{SMPdeep \rightarrow deepsed} \times \\ 367 \quad SMP_{deep-oce} \quad (Eq.12) \\ 368 \\ 369 \quad \frac{d(P_{beach})}{dt} = k_{P-beach} \times P_{surf-oce} - k_{P \rightarrow LMP} \times P_{beach} \quad (Eq.13) \\ 370 \end{array}$$

$$\frac{d(LMP_{beach})}{dt} = k_{LMP-beach} \times LMP_{surf-oce} + k_{P \to LMP} \times P_{beach} \quad (Eq.14)$$

$$\frac{d(LMP_{deep-sed})}{dt} = k_{LMP-sed} \times LMP_{surf-oce} \times f_{pelagic} \quad (Eq.15)$$

$$\frac{d(SMP_{deep-sed})}{dt} = k_{SMP-sed} \times SMP_{surf-oce} \times f_{pelagic} \quad (Eq.16)$$

$$\frac{d(SMP_{deep-sed})}{dt} = k_{SMP-sed} \times SMP_{surf-oce} \times f_{pelagic} \quad (Eq.16)$$

$$\frac{d(SMP_{deep-sed})}{dt} = k_{SMP-sed} \times SMP_{surf-oce} \times f_{pelagic} \quad (Eq.16)$$

based on reviews of literature data reporting MP counts per surface area and particle size statistics, is relatively straightforward. The beached MP 378 pool is estimated at 0.5 Tg, based on the global surface of sandy beaches $(2.63 \cdot 10^5 \text{ km}^2; (28))$, a median global 379 beach sand MP abundance of 2450 MP km⁻² (IQR, 613 - 2700), and median MP size of 2.0 mm (IQR, 1.1 - 2700) 380 3.8) (29). Reviews of deep ocean MP and shelf sediment MP pools report numbers of MP counts per mass 381 unit, which leads to more intricate pool mass estimates: Barrett et al. (31) reported mean deep sediment MP 382 concentrations of 0.72 MP g⁻¹ for cored and grab sediment samples of 9cm depth. Deep sea sedimentation 383 rates are typically on the order of 0.1-1 cm per 1000 years, suggesting that the majority of such composite 384 sediment samples pre-date the plastics mass production period <1950. Yet, the measurement (0.72 MP g⁻¹) is 385 expressed relative to the bulk of the composite sample mass, representing on average 9 cm of deep sea 386 sediment (31). In this case we used the following data to estimate the global deep sea MP pool mass: depth in 387 cm, dry sediment bulk density of 1.37 g cm⁻³, a water to sediment mass ratio of 3.0, the mean MP size of 0.1 388 mm reported (31), a MP density of 1 x 10^{-6} µg µm⁻³, and an open ocean seafloor surface area of 3.36 x 10^{8} 389 km². Similarly; the shelf sediment MP pool is estimated from subtidal sediment median MP concentrations of 390 100 MP kg⁻¹ (IQR, 32-120), reviewed and reported by Shim et al. (29), a corresponding median MP size of 391 392 2.0 mm (IQR, 1.1 - 3.8), a dry sediment bulk density of 1.37 g cm⁻³, a typical shelf sedimentation rate of 1 mm y^{-1} , 65 years of MP accumulation (1950 – 2015), a water to sediment mass ratio of 3.0, and a shelf seafloor 393 surface area of 3.53×10^7 km². The final estimates for the deep ocean and shelf sediment MP pools are 1.5 Tg 394 and 65 Tg (1σ , 21 to 78Tg) respectively. We acknowledge that plastic litter concentrates in given areas of the 395 396 seafloor, and therefore, sediment sampling data could be biased depending on the sampling site. This is ultimately reflected in the large budget uncertainties. 397

The global atmosphere. Brahney et al. (19,24) estimated the global atmosphere to contain 0.0036 Tg of SMP. 399 They also estimated global emissions from roads, 0.096 Tg y⁻¹, agricultural dust, 0.069 Tg y⁻¹, population dust, 400 0.018 Tg y⁻¹, and oceans, 8.6 Tg y⁻¹, which we adopt here. Atmospheric SMP deposition to remote terrestrial 401 surfaces has been investigated by Allen et al. (5) in France, finding a median SMP deposition of 0.011 Mg 402 km⁻² y⁻¹, and by Brahney et al. (24). who observed a median of 0.0012 Mg km⁻² y⁻¹ in the western USA. 403 Similar sampling and analysis techniques were used, and similar SMP particle and fiber size distributions 404 405 found, suggesting that the 9x difference reflects the difference in population density of both areas, 100 inhabitants per km² in SW Europe vs. 16 per km² in the western USA. In (sub-)urban environments in 406 Hamburg (Germany, 240 inhabitants per km²) mean SMP deposition of 0.016 \pm 0.006 Tg km⁻² y⁻¹ was 407 observed (38). Precursor studies on atmospheric plastics observed mostly the LMP fiber fraction (0.3 to 5mm) 408 with for example 0.014 Tg LMP km⁻² y⁻¹ in Dongguan (China) (23), but only 0.002 Tg km⁻² y⁻¹ in Paris 409 (France) (22). For simplicity we do not include LMP emission to the atmosphere in the box model, since the 410 short residence time of LMP likely leads to immediate deposition back to the broad terrestrial discarded LMP 411 reservoir. We regress SMP deposition over land, from the three detailed recent studies mentioned above, as a 412 function of population density (SI 1, Figure S1). We then extrapolate the observed relationship globally using 413 population density and surface area data per country for the year 2015 (45), capping SMP deposition at 0.016 414 Tg km⁻² y⁻¹ based on the Hamburg observations. Doing so leads to a global SMP deposition estimate over land 415 of 1.1 ± 0.5 Tg y⁻¹. SMP deposition over oceans is unconstrained by observations. We assume that global SMP 416 emissions (8.6 Tg y⁻¹; (19)) equal deposition, and estimate SMP deposition over oceans to be 7.5 Tg y⁻¹ (total 417 deposition of 8.6– 1.1 Tg y^{-1} deposition over land). 418

The mass inventory, emission and deposition flux estimates for 2015 serve to approximate the mass transfer coefficients associated with emission, $k_{oce \rightarrow atm}$ and deposition, $k_{atm \rightarrow oce}$, $k_{atm \rightarrow terr}$, in the following mass balance equation:

422

423 $\frac{d(SMP_{atm})}{dt} = k_{terr \to atm} \times SMP_{terr} + k_{disc \to atm} \times SMP_{disc} + k_{oce \to atm} \times SMP_{surf-oce} - k_{atm \to terr} \times$ 424 $SMP_{atm} - k_{atm \to oce} \times SMP_{atm}$ (Eq.17)

425

We assume $k_{terr \rightarrow atm}$ to be equal to $k_{disc \rightarrow atm}$ which was derived from the modeled discarded SMP pool and the anthropogenic SMP emission flux of 0.18 Tg y⁻¹ (sum of road, population and agricultural SMP emission) derived from the 3D global aerosol model for SMP dispersal by Brahney et al. (19).

429 **Remote terrestrial pool.** In the box model, agricultural and urban soils are included in the discarded plastics 430 pool. We use a separate box for remote terrestrial surfaces, outside of the technosphere, that are solely supplied 431 by atmospheric SMP. These include pristine soils, barren rock and land, ice sheets and remote inland waters. 432 We estimate the approximate amount of SMP in the remote terrestrial pool by making use of the quasi-linear 433 increase in global plastics production, discard and dispersal fluxes: global SMP deposition of 1.15 Tg y^{-1} in 434 2015 suggests a mean SMP deposition flux that is about half, 0.58 Tg y^{-1} since 1965, which multiplied by a 435 land surface area of 1.49 10⁸ km² amounts to 28 Tg of remote terrestrial SMP. SMP in this pool is mobilized 436 by rainfall to river runoff to the surface ocean, with the same k_{SMP→river-oce} that we derived for SMP runoff 437 from the discarded SMP pool. The remote terrestrial pool mass balance is: 438

439

$$\frac{d(SMP_{terr})}{dt} = k_{atm \to terr} \times SMP_{atm} - k_{terr \to atm} \times SMP_{terr} - k_{SMP \to river \to oce} \times SMP_{terr}$$
(Eq. 18)
441

BAU and SCS model scenarios. Both future, 2015 – 2050, model scenarios, business as usual (BAU), and 442 systems change scenario (SCS, from Lau et al. (4)), use the same mass transfer coefficients, k, but different 443 production, and waste management strategies summarized in the SI 2. BAU uses exponentially increasing 444 production, and quasi-linearly increasing incineration and recycling, and decreasing discard from Geyer et al. 445 (3). Lau et al. (4) developed a detailed model of plastics stocks and flows from municipal solid waste (MSW) 446 447 and four sources of LMP. Their CSC scenario presents the most complete, yet feasible plastics management strategy over the period 2016 – 2040 for MSW, including a decrease in plastics production by 2040 to 448 220 Tg y^{-1} . We digitized their disposal (incineration + safe landfilling), recycling and discard model output 449 (Tg y^{-1}), expressed these as fractions of MSW production, and extrapolated these to the year 2050 to compare 450 to BAU. To do so, we anchored (by normalization) the SCS disposal fractions for the period 2015 - 2050 to 451 the disposal fraction for 2050 - 2015 by Geyer et al.(3), in order to maintain a relatively smooth transition. 452 We acknowledge that the SCS waste disposal estimates deviate to some extent from the original (4) estimates, 453 but the overall trends are preserved: SCS disposal and recycling towards 2050 increase to 24 and 66 %, while 454 discard declines to 10%. Extrapolation of current waste disposal trends under the BAU scenario leads to 455 surprisingly similar numbers as SCS, though the real difference lies in the plastics production numbers that 456 reach 991 Tg y^{-1} under BAU, and drop to 168 Tg y^{-1} in the SCS by 2050 (SI 2). 457 458

Budget and model uncertainty. The model assumes no temporal evolution of the mass transfer coefficients, k, implying that fragmentation, sedimentation, emission, deposition and release dynamics are considered timeinvariant. While we argue that to first order these processes have remained similar through time, we acknowledge that reality is more complex. As more observational and mechanistic studies become available over the next decade, more appropriate parameterizations for plastics cycling can be tested, including the fragmentation of SMP to nanoplastics and ultimately dissolved and colloidal polymers with potential biological breakdown, i.e., as an energy source to biota.

Plastics data in the literature are predominantly reported as 'items per mass, volume, or surface area'. 466 We converted these data to mass numbers by taking into account, where possible, the reported particle size 467 distribution, or the reported median (or mean) particle size. In the case of fibers, reported length and diameter 468 were used. Studies that did not report particle size properties were not included in the budget estimates. 469 Particles were assumed to be flake shaped (46), with volume V defined as $V = L^{3*}0.1$, where L is the observed 470 effective diameter, and have a mean density of 1 g cm⁻³. In summary, for each particle size class, reported L 471 was used to compute flake volumes, then multiplied by particle/fiber number, and multiplied by density to 472 obtain particle/fiber mass. The obtained masses were summed to obtain total P, LMP or SMP mass in a sample. 473 Table 4 summarizes 1_o (one relative standard deviation, in %) expanded uncertainties of observed P,

Table 4 summarizes 1σ (one relative standard deviation, in %) expanded uncertainties of observed P, LMP and SMP pools (Tg) and fluxes (Tg y⁻¹), based on reported data, or conservatively approximated as

500%. The latter corresponds to a 2σ uncertainty of 1000%, which amounts to a factor 10. In other words, we
consider that a large number of plastics pools and fluxes are at the moment only known to within a factor of
10. In the future, as more observations on plastics pools, fluxes and degradation become available, we will
develop a formal Monte Carlo uncertainty analysis for the model.
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490 **Author contributions**

491 JES designed the study. JES, AK and JLT developed the model. All authors reviewed literature data, and 492 contributed to model data interpretation and writing.

494 Availability of data and material

The authors declare that the data supporting the findings of this study are available within the paper and its supplementary information files.

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498 Ethics approval and consent to participate

499

500

501 **Consent for publication**

502 The authors provide consent for publication

503 **Competing interests**

- 504 The authors declare no competing financial or other interests.
- 505 506

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- 627
- **Table 1. Subsurface ocean microplastics (MS) observations.** MS include fragments and fibers in the 0.3 5 mm (LMP) and <0.3 mm (SMP) range. Reported data in # m⁻³ were converted to mass concentrations, taking 628
- into account the full particle/fiber size distribution (see Methods). 629

Ocean basin	Location	depth	LMP+SMP	Reference
		m	μg m ⁻³	
N-Pacific	Korean East Sea	206	125	(34)
N-Pacific	Korean East Sea	2100	177	(34)
N-Pacific	Mariana Trench	2673	90	(6)
Mean			131	
1σ			44	
N- and S-	-53° S to 47° N	160	134	(36)
Atlantic				
N-Atlantic	Rockall Trough	2200	97	(35)
S-Atlantic	Gyre		43	(37)
Mean			91	
1σ			46	
Arctic Ocean	Central basin	5 to 1000	6	(47)
Arctic Ocean	Central basin	1769	66	(48)
Arctic Ocean	Fram Strait	300 to 5570	0.2	(49)
Mean			24	
1 σ			36	

- Table 2. Global subsurface ocean microplastics budget. Atlantic, N-Pacific and Arctic Ocean data from 631
- Table 1. Microplastics (MP) include fragments and fibers in the 0.3 5 mm (LMP) and <0.3 mm (SMP) range. 632
- Data for the S-Pacific and Southern Ocean are extrapolated based on surface Ocean data from Shim et al.(29) 633
- with uncertainties set to 10x. No data exists for the Indian Ocean, where concentrations were assumed equal 634 to the S-Atlantic observations by Eo et al.(34)(Table 1). Subsurface oceanic budgets in Tg include do not 635 636
- include the mixed layer (upper 0.1 km).

Ocean basin	Area	Volume	MP	MP	1σ
	km ²	km ³	μg m ⁻³	Tg	Tg
Arctic Ocean	15558000	18750000	24	0.4	0.6
North Atlantic	41490000	146000000	91	13.0	3.0
South Atlantic	40270000	16000000	91	14.3	3.3
Indian Ocean	70560000	264000000	43	11.0	11.0
North Pacific	77010000	331000000	131	42.2	14.1
South Pacific	84750000	329000000	4	1.2	12.0
Southern Ocean	21960000	71800000	4	0.3	3.0
Total				82	47

637

Table 3. Atmospheric small microplastics (SMP) budget. Mean $\pm 1\sigma$ SMP concentrations in the BL 639 640

641

(boundary layer) (144 \pm 124 ng m⁻³ for outdoors locations) and FT (free troposphere) (0.3 \pm 0.2 ng m⁻³) are from Allen et al. (25), assuming a mean SMP size of 70 µm for SMP in the BL, based on Brahney et al. (19).

	Mean global BL height	Mean global FT height	Area	BL SMP	FT SMP
	km	km	km ²	Tg	Tg
ocean	0.25	13	3.62 10 ⁸	0.013	0.0014
land	0.75	13	$1.48 \ 10^8$	0.016	0.0005
Total				0.031	

642

Table 4. Comparison of observed and modeled plastics mass (M, in Tg) and fluxes (F, in Tg y⁻¹) for the year 2015. Plastics are divided in macroplastics, P (>5 mm), large microplastics, MP (0.3 - 5mm), and small microplastics, SMP (<0.3 mm). Uncertainties (1 σ) on observations are based on the literature, except when not reported, in which case we assigned an uncertainty, denoted by '*'. Uncertainties (1 σ) on model estimated pools and fluxes are conservatively estimated to be 500% (denoted by '**'), corresponding to a 2 σ uncertainty of a factor of 10 (see Methods). The first column with M and F abbreviations correspond to parameter nomenclature used in mass balance equations 1-18.

Reservoir mass (M) or flux (F)	Abbreviation	Observed	Uncertainty	Box model
			1σ	
M P produced	P _{prod}	8300	10%*	8300
M P in-use	Puse	2600	10%*	3320
M P discarded	P _{disc}	4214	22%	2382
M LMP discarded	LMP _{disc}	686	22%	1222
M SMP discarded	SMP _{disc}			540**
M P Surface Ocean	P _{surf-oce}	0.23	75%*	0.021
M LMP Surface Ocean	LMP _{surf-oce}	0.031	75%*	0.038
M SMP Surface Ocean	SMP _{surf-oce}	0.0028	196%	0.009
M LMP Deep Ocean	LMP _{deep-oce}		F 70/	77
M SMP Deep Ocean	SMP _{deep-oce}	82	57%	33
M SMP atmosphere	SMP _{atm}	0.03	500%	0.011
M SMP remote terrestrial	SMP _{terr}	28	37%	41
M P beach	Pbeach	1.3	500%*	1.1
M LMP beach		0.53	100%	2.9
M P shelf sediment	P _{shelf-sed}	51	500%*	43
M LMP shelf sediment	LMP _{shelf-sed}	65	100%	9.5
M SMP shelf sediment	SMP _{shelf-sed}			0.3**
M LMP deep ocean sediment	LMP _{deep-sed}			1.2
M SMP deep ocean sediment	SMP _{deep-sed}	1.0	500%*	0.1
M P Incinerated	Pincin	800	20%*	626
M P Recycled	Precyc	750	20%*	554
F Puse to LMPdisc		42	22%	41
F Puse to Pdisc		118	22%	130
F Puse to Pincin		74	20%*	63
F Puse to Precyc		56	20%*	57
F P _{disc} to LMP _{disc}				71**
F LMP _{disc} to SMP _{disc}				36**
F P river				2.4**
F LMP river				7.5**
F SMP river (from SMP _{disc})				3.3**
F SMP river (from P _{terr})				0.3**
F river total		0.006-13		13
F Surface Ocean P to LMP		0.000 10		0.001**
F Surface Ocean LMP to SMP				0.001**
F Deep Ocean LMP to SMP				2.3**
F SMP Ocean to atmosphere		8.6	500%*	27
F SMP Atmosphere to ocean		7.6	500%*	23
F P beaching				0.1**
F LMP beaching				0.2**
F beach P to LMP				0.03**
F LMP surface to deep ocean				6.8**
F SMP surface to deep ocean	<u> </u>			0.3**
F P surface to shelf sediments				2.3**
F LMP surface to shelf sediments	<u> </u>			0.6**
F SMP surface to shelf sediments				0.02**
F LMP deep ocean to sediments				0.02
F SMP deep ocean to sediments				0.01**
F SMP terrestrial to atmosphere				0.01**
i Sivir terrestrial to atmosphere				0.01

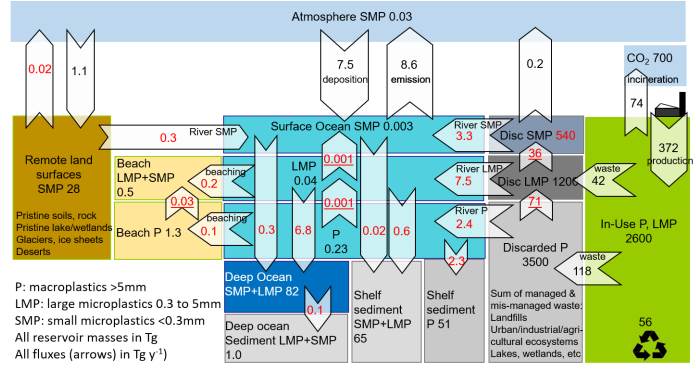
Non-peer-reviewed EarthArXiv Preprint

F SMP atm to terrestrial pool	1.1	37%	3.5
F SMP Discard to atmosphere	0.183	500%*	0.2

651

652 Figures

GLOBAL PLASTICS CYCLE FOR THE YEAR 2015



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Figure 1. Global plastics budget and cycle for the year 2015 based on best-available observations and 654 **model estimates.** Reservoir sizes are shown in teragrams (Tg), and fluxes in Tg y^{-1} (arrows). Three plastics 655 size classes are considered: macroplastics > 5mm (P), microplastics from 0.3 to 5mm (LMP), and small 656 microplastics <0.3mm (SMP) that can become airborne. The discarded (Disc) plastic pools represent the 657 terrestrial technosphere, where managed and mismanaged waste has accumulated in urban-industrial areas, 658 landfills, agricultural soils impacted by mulching or waste disposal, wetlands, lakes and other ecosystems. 659 The remote terrestrial reservoir lies outside the technosphere and is only impacted by airborne SMP deposition, 660 re-emission and runoff. Numbers in black are based on observations, and numbers in red on the box model 661 simulation. <u>Underlined</u> red fluxes indicate P and LMP degradation at a rate of 3% per year. Uncertainties are 662 provided in Table 4. Note that such the 2015 global budget is not at steady-state and fluxes and pool sizes 663 continue to gradually increase today and in the future. 664

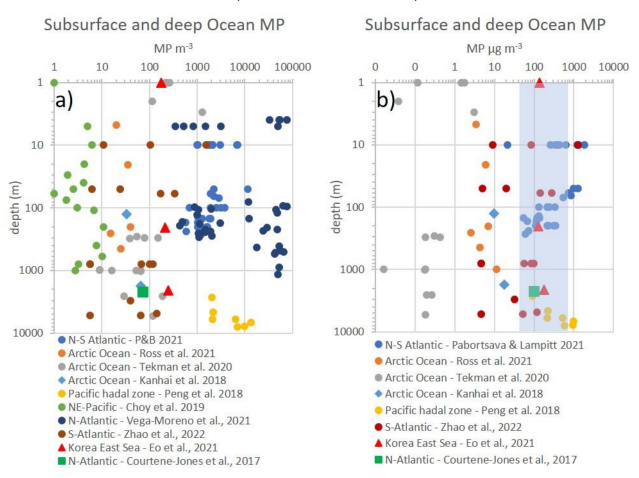
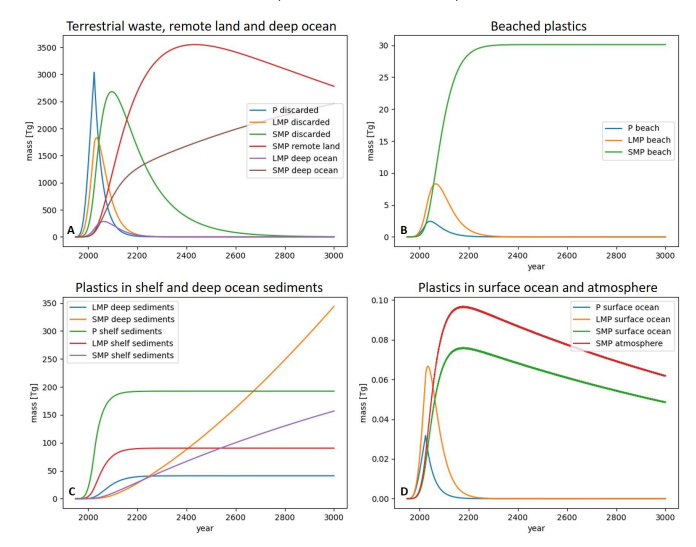


Figure 2. Subsurface Ocean microplastics (MP=LMP+SMP) observations. (A) MP number concentrations per m³ of sea water. (B) MP mass concentrations for datasets where particle/fiber size distribution was reported (see Methods). The shaded vertical bar indicates the range of mean $\pm 1\sigma$ mass concentrations estimated for the Pacific and Atlantic Oceans (45 to 175 µg m⁻³, Table 1).

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Figure 3. Plastics dispersal through Earth surface reservoirs from 1950 to the year 3000, following a 672 halt on production and discard in 2025. This unrealistic model scenario illustrates over what timescales 673 discarded microplastic (P, >5mm), large microplastic (LMP) and small microplastic (SMP, <0.3mm) 674 potentially disperse via rivers and air into oceans, remote terrestrial surfaces, beach and marine sediments. (A) 675 P and LMP disappear in all transitory reservoirs within 100 and 200 years due to fragmentation at an annual 676 rate of 3%. The prolonged dispersal of SMP in all reservoirs is driven by cyclical marine emissions to air, 677 deposition to terrestrial surfaces, runoff to surface oceans, and re-emission to air. Only a small fraction of 678 SMP sinks to shelf sediments and to the deep ocean, followed by slow sedimentation to deep ocean sediments. 679 SMP mass, and concentrations, in the surface ocean and atmosphere, where human SMP exposure is relevant, 680 only return to 2025 levels towards the year 5000 (Figure 4). 681

