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Special Section:

REgional Carbon Cycle Assessment and Processes - 2

Key Points:

- We explore the state-of-the-art in inland water greenhouse gas emissions, discussing existing estimates and underlying methodologies
- Development of models increasingly allows for assessment of spatial and temporal variability of emission fluxes
- There is a persisting need for observations that capture hot-spots and hot-moments in emissions, including from small water bodies

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










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Inland Water Greenhouse Gas Budgets for RECCAP2: 1. State-Of-The-Art of Global Scale Assessments

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Abstract Inland waters are important emitters of the greenhouse gasses (GHGs) carbon dioxide (CO₂), methane (CH₄), and nitrous oxide (N₂O) to the atmosphere. In the framework of the 2nd phase of the REgional Carbon Cycle Assessment and Processes (RECCAP-2) initiative, we review the state of the art in estimating inland water GHG budgets at global scale, which has substantially advanced since the first phase of RECCAP nearly 10 years ago. The development of increasingly sophisticated upscaling techniques, including statistical prediction and process-based models, allows for spatially explicit estimates that are needed for regionalized assessments of continental GHG budgets such as those established for RECCAP. A few recent estimates also resolve the seasonal and/or interannual variability in inland water GHG emissions. Nonetheless, the global-scale assessment of inland water emissions remains challenging because of limited spatial and temporal coverage of observations and persisting uncertainties in the abundance and distribution of inland water surface areas. To decrease these uncertainties, more empirical work on the contributions of hot-spots and hot-moments to overall inland water GHG emissions is particularly needed.

1. Introduction

Inland waters (streams, rivers, lakes and reservoirs) are net-sources of greenhouse gasses (GHGs) to the atmosphere. They receive considerable amounts of reactive organic matter from terrestrial ecosystems, promoting the production of GHGs like carbon dioxide (CO₂), methane (CH₄), and nitrous oxide (N₂O). Inland waters are usually net-heterotrophic, meaning CO₂ production through respiration exceeds CO₂ consumption by aquatic production (Battin et al., 2023). An additional source of inland water GHG emission comes from terrestrial and wetland runoff and drainage that can be oversaturated in dissolved CO₂ produced by microbial and root respiration (Abril & Borges, 2019). Once this supersaturated aqueous solution enters surface waters, it can release gas into the atmosphere and contribute to inland water CO₂ emissions. Similarly, inland waters receive dissolved CH₄ and N₂O from oversaturated soils and groundwater (Jurado et al., 2017; Rasilo et al., 2017). In addition, the sharp fronts between reducing and oxidizing conditions within the water column or at the interface between surface and subsurface environments (e.g., benthic and hyporheic zones) promotes the production and emissions of N₂O (Marzadri et al., 2017, 2021). Moreover, autochthonous aquatic production may enhance nitrification in the water column through increased oxygen levels, while it may stimulate denitrification and methanogenesis in reducing,

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benthic sediments through delivery of labile organic matter. These processes play an important role in the N₂O and CH₄ budgets of eutrophic lakes and reservoirs (DelSontro et al., 2018; Zhou et al., 2021).

While the processes driving GHG production have been known to limnologists for some time, large-scale quantification of inland water GHG emissions is still difficult and estimates are afflicted by large uncertainties. In their 5th Assessment Report (AR5), the IPCC (2013) acknowledged for the first time that inland waters are a significant contributor to the global GHG budget. At the same time, however, it was recognized that GHG fluxes from these ecosystems remain poorly constrained at the global scale. High uncertainties in inland water GHG emission estimates arise due to a poor spatial and temporal coverage of direct observations (Bastviken et al., 2011; Deemer et al., 2016; Regnier et al., 2013, 2022; Soued et al., 2016) and are reflected in the large range of estimated inland water GHG fluxes reported in AR5: 0.8–1.2 Pg C yr⁻¹ for CO₂, 8–73 Tg CH₄ yr⁻¹ for CH₄, and 0.1–2.9 Tg N yr⁻¹ for N₂O. In their 6th Assessment Report (AR6), the IPCC (2021) provides updated ranges for N₂O (0.5–1.1 Tg N yr⁻¹) and CH₄ (112–217 Tg CH₄ yr⁻¹) emissions which are narrower, but still reflect significant uncertainties in estimating inland water GHG emissions. Note that inland water emissions proportionally remain the largest source of uncertainty in the global land CH₄ budget (Canadell et al., 2011).

The REgional Carbon Cycle Assessment and Processes (RECCAP) initiative aims to establish the GHG budgets of large regions covering the entire globe at the scale of continents (or large regions) and large ocean basins, which are then synthesized at global scale. While the first phase of this initiative (RECCAP1; Canadell et al., 2011) focused on CO₂ only, now the second phase (RECCAP2) accounts for the three GHGs CO₂, CH₄, and N₂O. As part of RECCAP1, Raymond et al. (2013) re-estimated global inland water CO₂ evasion suggesting that the total flux could be as high as 2.1 Pg C yr⁻¹, which is about twice the estimates synthesized in AR5. This much higher estimate was due to a re-estimation of stream surface areas including small headwater streams which contribute disproportionately to the total water surface area and CO₂ emission, but which were neglected in earlier assessments that used data sets representing only larger global rivers (e.g., Cole et al., 2007). More importantly, Raymond et al. (2013) provided the first global maps of inland water CO₂ emissions, which allowed for the use of these estimates in regionalized, global C budgets (Bastos et al., 2020; Ciais et al., 2021; Zscheischler et al., 2017).

Since RECCAP1, a growing number of global estimates of inland water GHG emissions have been published, not only for CO₂ emissions (e.g., Holgerson & Raymond, 2016; Horgby et al., 2019; Lauerwald et al., 2015; Liu et al., 2022), but also for CH₄ (e.g., Holgerson & Raymond, 2016; Rosentreter et al., 2021; Stanley et al., 2016) and N₂O (e.g., Hu et al., 2016; Lauerwald et al., 2019; Maavara et al., 2019; Marzadri et al., 2021; Soued et al., 2016; Yao et al., 2020), or for all three GHGs combined (e.g., Deemer et al., 2016; DelSontro et al., 2018). The limited availability and quality (e.g., length and frequency of time-series), and uneven global coverage of observed emission rates (see e.g., Deemer et al., 2016) still represent a large source of uncertainty. While many studies build largely on the same data that was produced over the past decades, the amount and quality of empirical data is steadily increasing. In addition, global emission estimates profited from the appearance of new, improved data sets of inland water surface areas (Allen & Pavelsky, 2018; Lehner et al., 2011; Messenger et al., 2016; Verpoorter et al., 2014). Finally, global scale estimation of inland water GHG budgets have been improved through novel upscaling techniques based on statistical (e.g., DelSontro et al., 2018; Lauerwald et al., 2015) and process-based models of varying complexity (e.g., Maavara et al., 2019; Yao et al., 2020).

In the framework of RECCAP2, we present a review of existing global estimates of inland water GHG emissions. We start with a general overview of methods to achieve global scale estimates, starting from methods to measure flux rates in the field, followed by methods used to upscale flux rates to the global scale and which comprise a large range of approaches including simple upscaling based on average observed flux rates, statistical prediction and the use of process-based models (Section 2). Then, in three subsections respectively dedicated to estimates of emissions of CO₂ (Section 3), CH₄ (Section 4) and N₂O (Section 5), we discuss the state of the art for each of these GHGs in more detail, review all existing global estimates, and explore differences between flux assessments and their underlying methods. In addition, we highlight for each GHG persisting shortcomings and challenges for future research. The companion paper in the same issue (Lauerwald et al., 2023) then builds on the present review to derive a regionalized assessment for the 10 regions used in RECCAP2. In this companion paper, each previously published global estimate reported here was rescaled using the same new global assessment of inland water surface area, allowing for better consistency and homogeneity across all previously published values.

2. Overview of Upscaling Strategies and Surface Area Estimates Used in Global Studies of Inland Water GHG Emissions

This subsection gives a brief overview of different methods that are used to obtain global scale estimates of inland water GHG emissions. These methods are classified into three main approaches, namely direct upscaling based on observations Section 2.1, statistical upscaling based on functional relationships between emissions and environmental drivers Section 2.2, and process-based models Section 2.3. We also briefly review progress in the global scale assessment of inland water surface areas Section 2.4, which is of vital importance for global upscaling of inland water GHG emissions.

2.1. Upscaling Based on Observations

Large-scale estimates of inland water GHG emission fluxes F_{GHG} are usually calculated as the product of an average flux rate f_{GHG} , which can be expressed in units of mass per area and time, as derived from a set of field observations, and an estimate of the inland water surface area A_{IW} for which this flux rate is assumed to be representative (Equation 1).

$$F_{\text{GHG}} = f_{\text{GHG}} * A_{\text{IW}} \quad (1)$$

Many estimates have applied this simple upscaling technique directly at the global scale using an average f_{GHG} multiplied by the total A_{IW} of one specific type of inland waters. For instance, Deemer et al. (2016) calculated the average rates of GHG emissions from reservoirs, using observations from empirical studies around the world and multiplied those average rates by the estimated total area of reservoirs after Lehner et al. (2011). Others have first broken down the total of inland waters of one type into different subgroups, for example, based on size of water body or stream order (Holgerson & Raymond, 2016; Humborg et al., 2010), geographic region (e.g., Aufdenkampe et al., 2011; Bastviken et al., 2011; Johnson et al., 2021; Soued et al., 2016) or both (Raymond et al., 2013; Rosentreter et al., 2021). An area-integrated flux from each subgroup was then calculated following the same Equation 1, before summing those up to a global flux.

Methods and challenges to obtain estimates of A_{IW} are presented in detail in Section 2.4. In what follows, we will first focus on uncertainties associated with measuring and calculating f_{GHG} . Flux rates can either be obtained from GHG emission rates directly measured in the field (Section 2.1.1), or from measurements or calculation of GHG concentration gradients and concomitant measurements or models of gas transfer velocities (Section 2.1.2). Note that this study does not aim to provide a detailed review of field methods. These aspects are thus only briefly discussed, with a focus on methodological uncertainties.

2.1.1. Directly Observed Flux Rates

A common method to measure aquatic GHG emission rates is the use of floating chambers, which resemble inverted plastic buckets put onto the water surface. The emission rates are then calculated based on the accumulation rate of GHGs within the floating chamber headspace. This method detects the emission rate from the small surface area of the floating chamber across the larger area over which the chamber may be moving during the deployment. Chambers may drift a few meters if tethered or over longer distances if drifting freely during the deployment (Lorke et al., 2015). Such a well-defined footprint is advantageous for studies of local flux regulation and for distinguishing variability in space versus time. Concurrently, the small size of the footprint leads to potentially high uncertainties in the extrapolation of flux chamber measurements to large areas, without numerous representative measurements (Colas et al., 2020). Eddy covariance towers, though less common and only applicable in standing water bodies of a certain size have the advantage of generating net fluxes (i.e., emission or uptake) from a larger surface area (depending on height, surface roughness and wind speeds, eddy covariance towers can have a footprint of up to 3-km radius (Chu et al., 2021)), thus delivering a more representative emission rate (Podgrajsek et al., 2014). In contrast to the floating chamber method, the eddy covariance technique also allows for continuous measurements which provide better temporal resolution in emission rates. However, the flux footprint is constantly moving with wind speed and direction, making variability in time and space challenging to distinguish (Eugster et al., 2011). Fluxes cannot be measured at all when there is no wind (e.g., typical during night time) and complications associated with rainfall and lateral advective gas flux make accurate flux measurements challenging (Podgrajsek et al., 2014; Vesala et al., 2006). Eddy covariance also relies on the performance of advanced equipment and a high level of operator expertise for adequate data filtering and

QA/QC. Above all, limited eddy covariance measurements mean that global upscaling based only on this method is not yet possible, and inherent limitations due to its vulnerability toward unfavorable meteorological conditions and interactions with other nearby ecosystems make eddy covariance suboptimal for key inland water emission measurements such as fluxes from streams and along lake shores.

While the majority of CO₂ and N₂O emissions occur through diffusive flux across the air-water interface, a significant but variable fraction of aquatic CH₄ flux occurs as bubbles (i.e., ebullition) (Bastviken et al., 2011). Ebullition occurs when CH₄ produced in aquatic sediments forms gas bubbles that at a certain size, due to buoyancy, evade the sediment layer and ascend through the water column. Existing emission estimates from floating chambers sometimes intentionally exclude ebullition (e.g., Yang et al., 2021). Other floating chamber methods include both diffusive and ebullitive emissions (e.g., Barbosa et al., 2021). Also, eddy covariance towers measure the sum of both emission pathways (Eugster et al., 2011). There are nonetheless various methods to directly quantify ebullition. However, these methods detect bubbles rather than CH₄ and need supplementary measurements of CH₄ concentration within the bubble gas, usually from manually taken samples, to allow flux estimation (e.g., Linkhorst et al., 2020). This point is critical as CH₄ concentration in bubbles can vary widely, from less than 1% to >80% (Boereboom et al., 2012). The most common methods for directly quantifying ebullition rates is the bubble trap, an inverted funnel that collects ascending bubbles (Maeck et al., 2013) and is sometimes connected to a hydrostatic pressure sensor (Varadharajan et al., 2010) or specialized bubble size sensors (Delwiche & Hemond, 2017) to measure the timing and size distribution of ascending bubbles. Ebullition measurements based on point measurements in space and time are currently very labor intensive given the high spatiotemporal variability of ebullitive fluxes (Linkhorst et al., 2020). Echo sounders (Ostrovsky et al., 2008), robotic boats connected to optical methane detectors (Grinham et al., 2011) and under-ice surveys (Wik et al., 2011) have also been used to quantify ebullition rates. In addition, radar remote sensing approaches are currently being developed that could integrate over space and time for more representative measurements (Engram et al., 2020).

2.1.2. Estimating Diffusive Fluxes Based on Concentration Gradients

The methods for directly measuring emission rates can easily be applied in deeper, slower-moving waters (floating chambers and funnel traps) or in larger water bodies (eddy covariance). However, these methods are often not feasible for smaller streams. Instead, emission flux rates can be calculated from a gradient in concentrations of a specific GHG (ΔC_{GHG}) in the water close to the surface and in the overlying atmosphere and a gas exchange velocity k_{GHG} (Equation 2). Note that this method can be applied to flowing and standing water bodies of any size, but only allows estimation of diffusive emissions, and is not applicable for ebullition. The gradient ΔC_{GHG} can be calculated based on direct field measurements using headspace equilibration methods (e.g., Müller et al., 2015), or using measured headspace partial pressures and solubility constants that depend on salinity and water temperature (Weiss, 1970).

$$f_{\text{GHG}} = \Delta C_{\text{GHG}} * k_{\text{GHG}} \quad (2)$$

The headspace equilibration method consists of equilibrating a known volume of sampled water and a known volume of air or gas, with a known initial partial pressure of the GHG to be analyzed. After full equilibration, a sample of the headspace is analyzed by for example, gas chromatography (Natchimuthu, Wallin, et al., 2017), optical gas analyzers (Grilli et al., 2020), or other gas analysis methods, to measure the corresponding GHG partial pressure from which the C_{GHG} in the sampled surface water can be calculated. This concentration is compared with the theoretical concentration in equilibrium with the background air partial pressure of the GHG in focus to yield ΔC_{GHG} . In the case of CO₂, concentrations can also be calculated from observations of alkalinity and pH based on chemical equilibria and the assumption that non-carbonate contributions to alkalinity are negligible, which can be questioned in some common aquatic systems (Abril et al., 2015) (see Section 3.2 for more discussion).

Gas exchange velocity can be assessed through direct tracer studies in which a specific tracer gas is injected into the stream, and its loss is measured over a defined length. As this method is too cumbersome and costly to be applied everywhere, empirical equations have been established that relate k_{GHG} to the rate of energy dissipation at the water-air interface. Energy dissipation causes the turbulent mixing of the upper water column and thus determines the depth of the water column which interacts with the atmosphere through the process of diffusion. In fact, k_{GHG} in units of length per time (e.g., m day⁻¹) represents the depth of the upper water column that will equilibrate with the overlying atmosphere over that specific amount of time. For streams and rivers, this

energy dissipation rate can be estimated from stream flow velocity and stream channel geometry, in particular the slope of the stream channel (Natchimuthu, Sundgren, et al., 2017; Natchimuthu, Wallin, et al., 2017; O'Connor & Dobbins, 1958; Raymond et al., 2012). More recent work has however noted a breakpoint in the energy dissipation rate at which air entrainment and bubble formation cause k_{GHG} to increase more rapidly with energy dissipation (Ulseth et al., 2019). This suggests that assuming only diffusive water-air gas exchange, as it is assumed in most studies of inland water CO_2 emissions, may lead to underestimated gas transfer velocities in systems with very high hydrological energy. For lakes and reservoirs, empirical equations relate k_{GHG} to wind speed (Cole & Caraco, 1998), lake surface area (Read et al., 2012), or both (Vachon & Prairie, 2013), as the degree to which wind shear versus convective mixing dominate gas transfer dynamics generally changes as a function of waterbody size. More sophisticated modeling of k_{GHG} from lake hydrodynamics considering multiple turbulence-generating processes have also been developed (e.g., MacIntyre et al., 2021). It has been suggested that models of k_{GHG} should be locally validated whenever possible (e.g., Schilder et al., 2013).

2.2. Upscaling Based on Statistical Prediction

A variety of statistical methods have been used to upscale flux measurements/estimates to the global scale. These methods can be categorized into two groups of statistical upscaling approaches: (a) methods that predict emission rates directly, and (b) methods that first predict A_{IW} , ΔC_{GHG} , and k_{GHG} separately, and combine them using Equations 1 and 2 to estimate the emission flux $F_{\text{GHG,IW}}$.

A simple example for the first group of methods is the use of emission factors (EFs), which has been applied to estimate N_2O emissions from river networks (Beaulieu et al., 2011; Kroeze et al., 2010). Averaged EFs, typically defined as the ratio of N_2O emissions to riverine N loads, were derived from a number of field studies. These EFs were then multiplied by global, spatially explicit estimates of river N loads (e.g., Mayorga et al., 2010) to estimate global riverine N_2O emissions at the same spatial resolution as the riverine N loads. This method assumes that riverine N_2O emissions simply scale linearly to riverine N loads, which is problematic from a reaction kinetics point of view, as discussed in Maavara et al. (2019). As an alternative empirical approach, Hu et al. (2016) used findings from field studies to fit equations predicting riverine N_2O emissions as a nonlinear function of dissolved inorganic N yield and catchment area, thus overcoming some of the limitations of the EF approach.

Another prominent example for the first group of methods is the study by DelSontro et al. (2018), predicting global lake CH_4 emissions empirically. DelSontro et al. (2018) fitted multilinear regression equations to a database of literature studies of 166 water bodies quantifying lake CH_4 emissions, that predict the total (diffusive + ebullitive), annual emission flux from lake size and lake productivity (defined as chlorophyll or phosphorus concentration). The fitted regression equations were then applied to different global data sets/estimates of lake surface area and an assumed statistical distribution of lake productivity across global lakes to estimate the global-scale CH_4 emissions from these water bodies.

Examples for the second group of methods are the studies by Raymond et al. (2013), Lauerwald et al. (2015), and Horgby et al. (2019) that estimated CO_2 emissions from rivers at the global scale or for specific ecoregions (Horgby et al., 2019 focused on alpine streams). These studies all used global data sets including digital elevation models and their derivatives (stream network and channel slope) and gridded estimates of average annual river flow to explicitly estimate stream surface area and k_{GHG} spatially. While Raymond et al. (2013) combined their estimates of A_{IW} and k_{GHG} with regionalized averages of calculated ΔC_{CO_2} , Lauerwald et al. (2015) and Horgby et al. (2019) further used multiple linear regression models to estimate riverine ΔC_{CO_2} from different spatial drivers (like terrestrial Net Primary Productivity—NPP, climate, terrain steepness in Lauerwald et al. (2015), or elevation, soil carbon stocks, and discharge in Horgby et al. (2019)). Note that combining independent estimates of k_{GHG} and ΔC_{GHG} introduces an additional source of uncertainty, as ΔC_{GHG} is in turn controlled by k_{GHG} and its balance with CO_2 resupply rates to the surface water, which is for instance evidenced by low ΔC_{GHG} in turbulent, high alpine streams (Horgby et al., 2019).

2.3. Process-Based Models

Process-based models of varying degrees of complexity have recently been used to assess inland water GHG emission at the global scale (Maavara et al., 2019; Marzadri et al., 2021; Yao et al., 2020). Ideally, such models represent carbon and nutrient transport and transformation processes that drive production, cycling and emission

of GHGs in a water body or along a cascade of water bodies (like a sequence of stream reaches or a cascade of reservoirs along a river network). This representation requires boundary condition data at the global scale and in sufficient quality and quantity. This data requirement is a major limitation for the applicability of process-based models for inland water GHG emissions at the global scale.

A promising strategy to overcome that limitation is the explicit representation of inland waters and associated biogeochemical processes in land surface models (LSMs) that simulate the terrestrial cycling of energy, water, C, nutrients, and GHGs. Using LSMs, the biogeochemical and transport processes that drive the GHG dynamics can be simulated simultaneously for terrestrial and freshwater ecosystems, reducing the need for complex boundary conditions at the land-inland water interface. Developments in that direction have been achieved for the LSMs DLEM (Tian, Yang, et al., 2015; Tian, Ren, et al., 2015; Yao et al., 2020) and ORCHIDEE (Lauerwald et al., 2017, 2020). At global scale, LSM simulations including inland water GHG emissions have yet only been achieved with DLEM (Yao et al., 2020).

When using LSMs, the simulated water fluxes and associated terrestrial C and nutrient inputs to inland waters are already afflicted by considerable uncertainties, including those arising from the overparameterization of these extremely complex models. Thus, an alternative is to use process-based models of only inland waters forced by data driven information. The global river network N₂O modeling studies by Maavara et al. (2019) and Marzadri et al. (2021) follow two different strategies to overcome data limitations to constrain the models. Maavara et al. (2019) followed a metamodeling strategy, for which a box model representing all major processes of N and N₂O cycling in a water body was first set-up. While this process-based model could not be applied at global scale due to data limitations to constrain each biogeochemical process, Maavara and colleagues ran the model across a realistic range of model input parameters using a Monte Carlo approach to derive simple response functions. The resulting response functions relating N₂O emissions to nutrient loads and water residence times were then applied at global scale using loads and residence times derived from available global data sets. Marzadri et al. (2021) applied their process-based model of river N₂O emissions directly at global scale, which required spatially resolved model inputs comprising a detailed set of parameters describing stream hydro-morphology and water quality, which in that form did not yet exist at global scale. To overcome that limitation, machine learning techniques were applied to derive these input data sets from other, available geodata. These input data were then used to feed a process-based model that parametrizes N₂O emissions as a function of river size by means of two Damköhler numbers representing the ratio between a characteristic time of transport and a characteristic time of reaction. The proposed hybrid model (machine learning + process based) allows the consideration of the contribution of surface (e.g., water column) and subsurface (e.g., benthic and hyporheic zones) processes to N₂O emissions (Marzadri et al., 2021).

For aquatic CH₄ emissions, process-based modeling efforts have been mostly dedicated to lake and reservoir systems. For example, an online, open-source predictive model framework “G-res” has recently been developed to provide global, spatially explicit estimates of the form and magnitude of reservoir CH₄ and CO₂ emissions (Harrison et al., 2021; Prairie et al., 2021). G-res uses a series of calibrated empirical models that integrate local (reservoir-specific) and regional (watershed attributes) information to predict GHG emissions (Prairie et al., 2021). The model has been applied to 4,727 reservoirs to estimate global emissions (Harrison et al., 2021). Tan and Zhuang (2015a, 2015b) have developed and applied a process-based model to estimate CH₄ emissions from lakes at pan-arctic scale. That model produces gridded output, resolves seasonal and interannual variability, and permits for projections of long-term trends following global change scenarios.

2.4. Available Data and Previous Estimates of Global Inland Water Surface Area

The first digital global map of inland water surface areas that was used for inland water GHG emission estimates was the Global Lake and Wetland Database (GLWD) by Lehner and Döll (2004). GLWD was derived from a compilation of different global and regional inventories. While GLWD is not globally consistent with regard to detail and reliability of the data sources, it represented the best available data set for more than a decade and was used in numerous studies of inland water GHG emissions (e.g., by Raymond et al. (2013) for lakes and reservoirs, by Aufdenkampe et al. (2011) for all water bodies). Since then, our ability to estimate the global surface area of rivers, lakes, and reservoirs has progressed significantly. This progress has been driven by advances in satellite remote sensing, image processing methods, and geospatial analysis techniques. Several freely available global hydrography data sets have recently become available that can be used to estimate surface area and distribution

of inland water bodies. Here we discuss a selection of high-resolution, freely available data sets that can be useful for global-scale evaluations of greenhouse gas emissions from inland water bodies.

A few global inland water body data sets have been developed using optical remote sensing data. The JRC GSW data sets from Pekel et al. (2016) and the GSWD from Pickens et al. (2020) are two global 30-m-resolution data sets of open surface water extent, created from the Landsat archive. These data sets are multitemporal and highly consistent but they do not distinguish between different water body types (e.g., rivers, lakes, etc.). Classifying water body type is necessary in evaluations of GHG exchange because of differing exchange rates and processes occurring in different aquatic environments. The Global River Widths from Landsat (GRWL) database (Allen & Pavelsky, 2018) contains exclusively river surface areas derived from Landsat imagery.

In addition to these image-based data sets, global topography-based data sets derived from digital elevation models (DEMs) have been used for representing the global extent and distribution of streams and rivers. These include hydrologically conditioned gridded raster data sets like HydroSHEDS (Lehner et al., 2008) and MERIT Hydro (Yamazaki et al., 2019) or vectorized flowline data sets derived from these gridded data sets including HydroRIVERS (Lehner & Grill, 2013) or MERIT Hydro–Vector (Lin et al., 2021). These DEM-based data sets can be used to infer the location and size of narrow rivers and streams too small to be visible from freely available satellite data sets. These data sets can also be used to infer other characteristics of river networks including stream order, slope, upstream area, and topology, which are of potential value for estimating amount and turbulence of river flow, which in turn are important drivers of GHG emissions. Other hydrography data sets innovatively combine DEM-based data sets with other sources of data to produce novel information including machine learning-based estimates of river surface area (Lin et al., 2021) and the extent of non-perennial rivers (Lin et al., 2021; Messenger et al., 2021).

For standing open water bodies like lakes and reservoirs, attempts have also been made to identify water bodies from satellite imagery using automated algorithms. A prominent example is the Global Water Body (GLOWABO) data set (Verpoorter et al., 2014). Due to the unsupervised classification method and the fact that ground truth was evaluated only for Sweden, this data set is however uncertified for other parts of the world and likely contaminated with wrongly assigned riverine, coastal or temporal water bodies (Pi et al., 2022). In addition, inventory-based data sets have further been developed, including the Global Reservoir and Dam database (GRanD) (Lehner et al., 2011) and the HydroLAKES database (Messenger et al., 2016) which gives water surface areas of standing waters distinguishing lakes from reservoirs. Note that HydroLAKES also includes the information from GRanD and GLWD, which makes these products partly redundant. The advantage of inventory-based data sets as GRanD and HydroLAKES is the avoidance of contamination with other water bodies and additional attributes such as names, estimates of water volume and residence time, height and purpose of dam for reservoirs, etc. In particular the distinction between lakes and reservoirs is of major importance for the assessment of inland water GHG emissions. Reservoirs as artificial water bodies deserve special attention, as they represent an anthropogenic source of GHGs and a potential lever for controlling future emissions (Almeida et al., 2019). However, we have to expect an under-classification of reservoirs in inventory data sets such as HydroLAKES, as water bodies for which this information was not available have been categorized as natural lakes by default (Messenger et al., 2016). Smaller hydropower projects which outnumber large hydropower projects by approximately 11:1 (Couto & Olden, 2018) may not always be inventoried and accounted for in regional and global data sets. Recently, new data sets of dams and reservoirs have been created combining remote sensing-based data sets with other sources of information, for example, GOODD (Mulligan et al., 2020) and GeoDAR (Wang et al., 2021), continuously increasing the numbers of reservoirs that are taken up into inventories.

Although considerable progress has been made recently in developing global hydrography data sets, much less work has been done to apply these data sets to estimate global surface area of inland water bodies. For the surface area of rivers, three notable global estimates have been produced by Downing et al. (2012) of between 485,000 and 682,000 km², Raymond et al. (2013) of between 487,000 and 761,000 km², and Allen and Pavelsky (2018) of 773,000 ± 79,000 km². Downing et al. (2012) based their estimate on >400 observations of stream width, data on number and length of streams from HydroSHEDS data set, and statistical scaling relating stream number, width and length to stream order. While this observational data set covers rivers from all over the world, it is clearly dominated by North American rivers and the extent to which the statistical relationships identified in that study is applicable to the global-scale is questionable. Raymond et al. (2013) combined the stream network of HydroSHEDS with gridded runoff data to obtain a distribution of stream lengths and discharge per stream order

of medium to large rivers, to which they then applied empirical, hydraulic equations predicting stream width from discharge. Finally, they used stream-order based scaling laws to estimate stream surface areas for smaller streams. Allen and Pavelsky (2018) used their remote-sensing based GRWL database for surface areas of medium to large rivers, which they complemented with topography- and statistical-based estimates for streams narrower than 90 m to headwater streams as defined by Allen et al. (2018). The GRWL data set is to date the most complete and reliable data set of its kind.

For the surface area of lakes and reservoirs, three notable global estimates have been made by Downing et al. (2006), Verpoorter et al. (2014), and Messenger et al. (2016). Downing et al. (2006) used surface areas from standing water bodies $>10 \text{ km}^2$ from GLWD (Lehner & Döll, 2004) and extrapolated the surface area to smaller water bodies down to 0.001 km^2 assuming power-law relationships (Pareto distributions) between water body size and frequency. Note, however, that a more recent empirical study disproved the hypothesis that number and area of small lakes would follow a power-law distribution (Cael & Seekell, 2016). Verpoorter et al. (2014) used their remote-sensing derived GloWaBo database which includes lakes as small as 0.002 km^2 . Messenger et al. (2016) derived their estimate from their inventory based HydroLAKES database, which contains water bodies $>0.1 \text{ km}^2$. Due to this restriction with regard to minimum lake size, Messenger et al. (2016) obtained the lowest of the three global surface area estimates for standing waters with $2.7 \times 10^6 \text{ km}^2$. The estimate of Downing et al. (2006) is substantially higher with $4.2 \times 10^6 \text{ km}^2$, while for water bodies larger than 0.1 km^2 , their estimate of $2.9 \times 10^6 \text{ km}^2$ is quite comparable to HydroLAKES. The estimate by Verpoorter et al. (2014) is even higher with $5 \times 10^6 \text{ km}^2$, likely due in part to overestimation of lake areas through contamination with other water bodies (Pi et al., 2022). A reliable map of smaller bodies of standing water, such as ponds, which are thought to contribute substantially to the total water surface area and disproportionately to GHG emissions (Holgerson & Raymond, 2016; Rosentreter et al., 2021), is still not achievable.

3. Inland Water CO₂ Budget

3.1. Overview of Existing Estimates

Global estimates for the aquatic CO₂ emission range from 0.84 to $7.33 \text{ Pg CO}_2 \text{ yr}^{-1}$ for streams and rivers, from 0.40 to $2.14 \text{ Pg CO}_2 \text{ yr}^{-1}$ for lakes, from 0.08 to $0.14 \text{ Pg CO}_2 \text{ yr}^{-1}$ for reservoirs (excluding the estimate by Cole et al. (2007), which is discussed at the end of this section), and from 0.89 to $2.35 \text{ Pg CO}_2 \text{ yr}^{-1}$ for estimates that lumped lakes and reservoirs together (Table 1, Figure 1). In general, considerable discrepancies exist in particular between early estimates that relied mostly on lumped estimates of average CO₂ concentrations, k_{GHG} and water surface area, and more recent estimates relying on more complete concentration data sets, more sophisticated upscaling approaches and spatially resolved water surface area estimates. For streams and rivers, the earliest estimates (Cole & Caraco, 2001; Cole et al., 2007, p. 207) were crude and most likely underestimate riverine CO₂ emissions because of their reliance on data from large rivers, which tend to show lower areal CO₂ emission rates than smaller and more upstream systems, as large rivers tend to be less heterotrophic, receive less important inputs of CO₂ enriched groundwater, and show less turbulent stream flow which leads to lower gas exchange velocities (Raymond et al., 2013). Relying on an extensive database for $p\text{CO}_2$, new scaling laws for k_{GHG} and stream hydraulic geometry that allowed for spatially resolved estimates for stream surface areas at the global scale, Raymond et al. (2013) presented the first spatially explicit estimate for the aquatic CO₂ flux and reports a river CO₂ evasion rate that is 3–8 times higher than the earlier lumped estimates (Aufdenkampe et al., 2011; Cole et al., 2007; Tranvik et al., 2009). Moreover, they demonstrated the importance of small headwaters which contribute disproportionately to the total emission flux. Problematic in that approach was that average $p\text{CO}_2$ and k_{GHG} values, from which the fluxes were calculated, were estimated independently from each other. $p\text{CO}_2$ values were taken as average over large regions independent of stream order, while k_{GHG} was estimated per region and stream order with values systematically increasing toward the small stream orders. That bears the risk, in particular for low order streams, that high estimates of k_{GHG} are combined with high estimates of $p\text{CO}_2$, leading to unrealistic high flux rates, whereas empirical studies have shown that under higher k_{GHG} values $p\text{CO}_2$ normally tends to be close to equilibrium with the atmosphere (Rocher-Ros et al., 2019). More recent advancements in stream and river CO₂ evasion estimates involve the development of data-driven statistical models to resolve temporal and finer spatial scale variations of the riverine CO₂ flux (Lauerwald et al., 2015; Liu et al., 2022). For instance, relying on direct C_{CO_2} measurements and seasonally varying estimates for k_{GHG} and river surface area, Liu et al. (2022) demonstrated CO₂ emission from global streams and rivers varied between 411 and $766 \text{ Tg CO}_2 \text{ yr}^{-1}$ per month, that is, by a factor of ~ 2 , with the highest global emissions during northern summer in July.

Table 1
Global Estimates of Inland Water CO₂ Emissions

References	$\Sigma\text{CO}_2\text{em}/\Sigma A_{\text{water}}$ (g CO ₂ m ⁻² yr ⁻¹)	ΣA_{water} (10 ⁶ km ²)	$\Sigma\text{CO}_2\text{em}$ (Pg CO ₂ yr ⁻¹)	Method
Rivers				
DLEM Tian, Ren, et al. (2015)	3,531	0.64	2.24	Model
Liu et al. (2022) ^a	9,900	0.672	7.33 ± 0.73 ^b	Machine learning
Lauerwald et al. (2015) ^a	3,895	0.55–0.67	2.38 (1.77–3.10) ^c	Statistical prediction
Raymond et al. (2013) ^a	10,644	0.62	6.6 (5.7–7.5) ^c	Upscaling from observations + Statistical prediction
Aufdenkampe et al. (2011)	5,009	0.31–0.51	2.05	Lumped estimate
Tranvik et al. (2009)			2.02	Literature review
Cole et al. (2007)	1,492	0.74	0.84	Literature review
Streams and small rivers				
Liu et al. (2022) ^a	23,962	0.202	4.84	Machine learning
Marx et al. (2017)			3.41	Literature review
Lauerwald et al. (2015) ^a	5,842	0.14–0.26	1.16 (0.78–1.61) ^c	Statistical prediction
Mountain streams				
Horgby et al. (2019)	17,490	0.035	0.61	Statistical prediction
Large rivers				
Liu et al. (2022) ^a	4,946	0.47	2.31	Machine learning
Lauerwald et al. (2015) ^a	2,975	0.41	1.22 (0.96–1.54) ^c	Statistical prediction
Lakes and reservoirs				
DelSontro et al. (2018)	414	3.23–5.36	1.99–3.30	Avg. rates
-"	416	4.42	1.84 (1.72–1.98) ^c	Statistical prediction
-"	360	5.36	1.93 (1.80–2.06) ^c	Statistical prediction
-"	276	3.23	0.89 (0.83–0.96) ^c	Statistical prediction
Raymond et al. (2013) ^a	392	3	1.17 (0.22–3.08) ^c	Avg. pCO ₂ + Statistical prediction
Aufdenkampe et al. (2011)	638	2.80–4.54	2.35	Avg. rates
Lakes (including lakes with dams)				
DLEM Tian, Ren, et al. (2015)	312	2.4	0.77	Model
Holgerson and Raymond (2016)	348	5.98	2.14	Avg. rates
Tranvik et al. (2009)			1.94	Literature review
Cole et al. (2007)	257	2	0.4	Literature review
Reservoirs				
DLEM Tian, Ren, et al. (2015)	312	0.27	0.08	Model
Deemer et al. (2016)	451	0.3	0.14 (0.12–0.16) ^c	Avg. rates
Cole et al. (2007)	686	1.5	1.03	Literature review

Note. For each Estimate, the Total Water Surface Area (ΣA_{water}), the Total CO₂ Emission Elux ($\Sigma\text{CO}_2\text{em}$) and the Area Weighted Average Emission Rate ($\Sigma\text{CO}_2\text{em}/\Sigma A_{\text{water}}$) are Reported. Mass units refer to CO₂. For conversion to units of mass of C, divide by 3.67.

^aEstimate accounts for effects of seasonal ice cover. ^bStandard error. ^cLower and upper 90% (Deemer et al., 2016; Lauerwald et al., 2015; Raymond et al., 2013) or 95% (DelSontro et al., 2018) CI.

A process-based model has also been developed (DLEM, Tian, Ren, et al., 2015), which predicts a much lower emission rate than recent data-driven approaches (2.24 vs. 6.60–7.33 Pg CO₂ yr⁻¹) (Table 1). The DLEM estimate is however close to the estimate by Lauerwald et al. (2015) (2.38 Pg CO₂ yr⁻¹) that only accounted for emissions from medium-sized to large rivers (i.e., 3rd order and above). The large discrepancy (i.e., 2.24

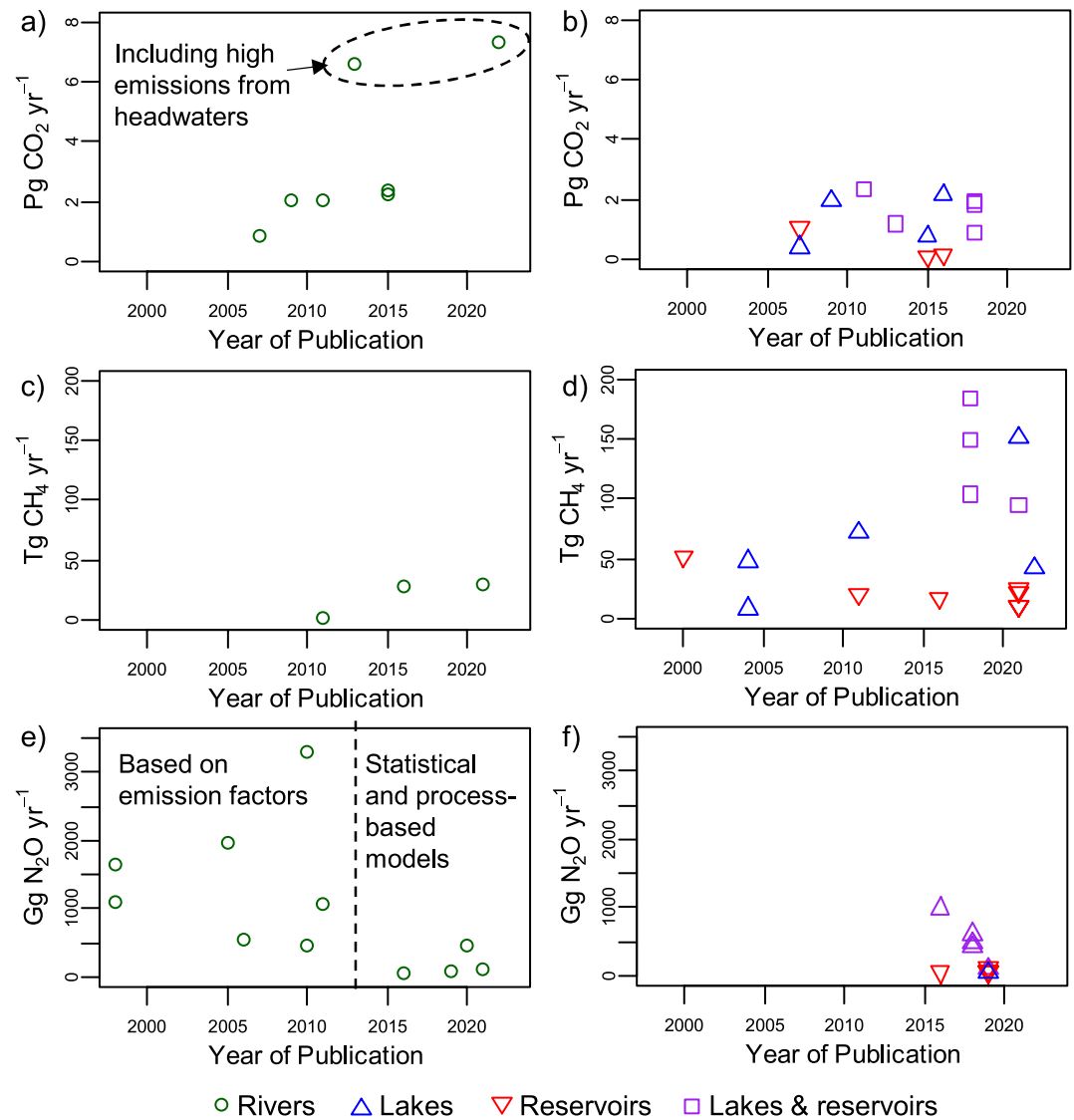


Figure 1. Estimates of inland water GHG emissions by year of publication. For rivers (a, c, e) versus Lakes and reservoirs (b, d, f), and for CO₂ (a, b), CH₄ (c, d), and N₂O (e, f).

vs. 6.60–7.33 Pg CO₂ yr⁻¹) however argues for the importance of the smallest streams in global CO₂ emission from fluvial networks (Marx et al., 2017). In line with this, Liu et al. (2022) estimated emission from the medium-to-large rivers (corresponding roughly to stream order 3 and above as in Lauerwald et al. (2015)) of ~2.31 Pg CO₂ yr⁻¹, while roughly two thirds of the total riverine emissions (~5 Pg CO₂ yr⁻¹) are predicted to be emitted by smaller streams (extrapolated to a minimum stream width of 0.3 m).

For lakes, there is a much larger variation in estimates of water surface area than in average emission rates between different studies (Table 1). In particular, estimates that relied on earlier global lake inventories (Raymond et al., 2013) report lower surface area and total emissions than more recent estimates based on newer lake inventories and extrapolated surface area to account for the smallest water bodies (DelSontro et al., 2018; Hastie et al., 2018; Holgerson & Raymond, 2016). Despite employment of scaling laws (e.g., with lake size and nutrient status) that account for spatial variability due to system size and autotrophic productivity in more recent estimates (DelSontro et al., 2018; Holgerson & Raymond, 2016; Raymond et al., 2013), there seems to be only small difference with regard to global average lake CO₂ emission rates per water surface area between those newer estimates (348–414 g CO₂ m⁻² yr⁻¹) and those of the early crude estimates (257 g CO₂ m⁻² yr⁻¹, Cole et al., 2007). Additionally, though earlier estimates relied more on C_{CO2} calculated from pH and alkalinity

(Cole et al., 2007; Raymond et al., 2013), more recent estimates used more often direct measurements (Delson-tro et al., 2018; Holgerson & Raymond 2016). Differences in lake CO₂ evasion estimates are more driven by variation in estimates of lake area than by areal emission rates. Estimates of global average emission rates per water surface area for lakes and reservoirs (392–638 g CO₂ m⁻² yr⁻¹) are about one order of magnitude lower than those for streams and rivers (1,492–10,644 g CO₂ m⁻² yr⁻¹, Table 1). In comparison to lakes, reported global average emission rates per water surface area for reservoirs are slightly higher (312–686 vs. 257–348 g CO₂ m⁻² yr⁻¹ for reservoirs and lakes, respectively) (Table 1). This may in part be due to the different geographic distribution of both types of standing water bodies; with lakes being particularly abundant in high latitudes where average emission rates tend to be lower (Aufdenkampe et al., 2011). Nonetheless, the current estimates place total CO₂ evasion from reservoirs more than one order of magnitude lower than that from lakes (see Table 1, when excluding the estimate by Cole et al. (2007)), following its low share in the global surface area of standing water bodies. However, the inventory for global reservoirs (which is growing) is far from complete and thus surface area might pose the largest uncertainty for CO₂ evasion from reservoirs. Note that Cole et al. (2007), based on data from St. Louis et al. (2000), estimated a reservoir CO₂ emission of 1.03 Pg CO₂ yr⁻¹, that is, about one order of magnitude higher than the other estimates listed in Table 1. This number is based on a first-order estimate of the total surface area of reservoirs including smallest systems such as farm ponds. This estimated total area is about 5 times larger than that of reservoirs accounted for in recent inventories. While this first order estimate is an eye-opener for the underestimate related to the exclusion of these small systems, it is also highly uncertain and represents an expert opinion rather than a reproducible number. Note further that Cole et al. (2007) estimated a much lower CO₂ evasion rate from lakes, for which they rely on a much more conservative estimate of surface area which excludes smaller systems. In that regard, their emission estimate for standing waters is not consistent. Note finally that other estimates of CO₂ emissions from reservoirs might be underestimated, as in inventories, where the required information is missing, reservoirs might wrongly have been classified as lakes (see discussion in Section 2.4).

3.2. Persisting Shortcomings and Future Challenges

3.2.1. Process Understanding

In the following we discuss our current process understanding of inland water CO₂ emissions in the light of global scale estimates. Figure 2 gives an overview of the main fluxes and processes involved, while Table 2 summarizes the known effects of the major environmental drivers. The most prominent gap in the understanding of the processes that drive inland water CO₂ emissions is the question of where the emitted CO₂ is sourced from. A part of the emitted CO₂ may be produced in situ from the oxidation of allochthonous organic carbon, while another part might stem from inflows of water supersaturated in CO₂ produced during respiration in upland soils and wetlands. Further, this respiration comprises both heterotrophic respiration of plant and soil organic matter as well as autotrophic root respiration. Knowledge about the source of the aquatic CO₂ emissions is of paramount importance for the integration of these fluxes in the overall C budget of continents, as highlighted in the perspective article by Abril and Borges (2019). While earlier studies assumed that the net-CO₂ emissions are entirely the product of heterotrophic respiration and could thus be regarded as a fraction of terrestrial net-primary production (Richey et al., 2002), the contributions of autotrophic root respiration demand consideration of these fluxes as part of total ecosystem respiration that counterbalances gross primary production (Abril & Borges, 2019; Lauerwald et al., 2020).

For streams and rivers, it is assumed that most of the emitted CO₂ is sourced from CO₂ produced by respiration in adjacent wetlands, but also in upland soils, from where the CO₂ is then transported with percolating soil water and groundwater flows (Abril et al., 2014; Liu et al., 2022). The relative importance of groundwater CO₂ inputs are highest in headwaters and decrease downstream (Finlay, 2003; Horgby et al., 2019; Hotchkiss et al., 2015; Liu et al., 2022; Marx et al., 2017), while wetland inputs may be more important in lower reaches as shown for the Congo River (Borges et al., 2019). Moreover, it was shown that due to the very high oversaturation of emerging groundwater, a large part of the emission already takes place over a few hundred meters downstream of the freshwater source (Johnson et al., 2008). It would thus be required to monitor smallest headwaters directly to well capture those hot spots of aquatic CO₂ emission, for which however monitoring data are not available in sufficient quantity (Marx et al., 2017). Assessment of groundwater CO₂ inputs to inland waters would further require knowledge about groundwater C content and residence time (to quantify the outflows) for which data are limited

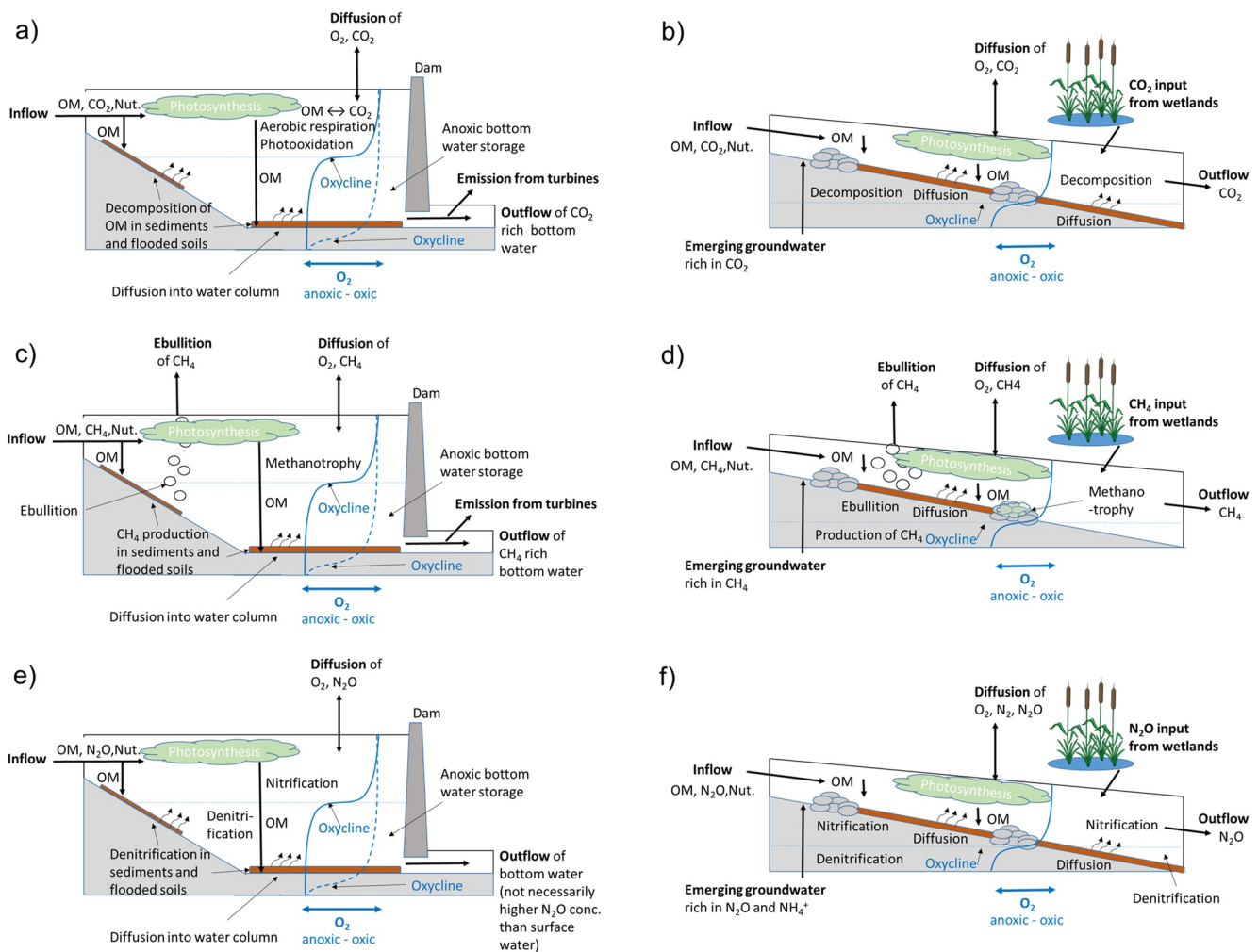


Figure 2. Schematic representation of inland water GHG budgets: for reservoirs as an example of standing waters (a, c, e) and streams/ivers (b, d, f), and for the three GHGs CO₂ (a, b), CH₄ (c, d), and N₂O (e, f). OM = organic matter; Nut. = nutrients.

as well (Downing & Striegl, 2018). While stable C isotopes have been used to estimate source contribution of riverine C loads and CO₂ emissions for single aquatic systems (Telmer & Veizer, 1999), observational data are not yet sufficient for large-scale assessment. Also, these studies do not often include the uppermost parts of the river network where large amounts of external CO₂ inputs are evading to the atmosphere.

In addition, only few existing studies of freshwater CO₂ emissions (e.g., Bogard & DelGiorgio, 2016; Crawford, Lottig, et al., 2014) have attempted to include estimates of aquatic net ecosystem production (NEP), which is the difference between aquatic production and respiration. Most studies currently assess inland waters net-CO₂ emissions rather as a black box that is fed by allochthonous C inputs. The recent study by Battin et al. (2023) has nevertheless demonstrated that the inclusion of NEP estimates can help to disentangle autochthonous CO₂ production from allochthonous CO₂ inputs even at global scale. This study corroborates the assumption that most of the aquatic CO₂ evasion is derived from external CO₂ inputs. However, also availability of aquatic NEP data is limited and does not allow yet for spatially explicit estimates at global scale. More importantly, diurnal variations in NEP may entail a similar variation in pCO₂ and air-water CO₂ exchange. Moreover, predominant sampling during daytime, when CO₂ emissions are lower than at night, may lead to important biases in flux estimations. Gómez-Gener et al. (2021) recently argued that global estimates based on daytime measurements are biased as night time emissions are on average ~30% higher.

To better understand temporal variability and potential “hot-moments” of inland water CO₂ emissions, more process understanding would be required with regard to CO₂ cycling during periods of ice-cover, spring ice-melt,

Table 2
Drivers of GHG Emissions From Different Types of Inland Waters^a

	CO ₂	CH ₄	N ₂ O
Streams and rivers	++ GW ^b inputs	++ GW inputs	++ GW inputs
	++ Inputs from wetlands	++ Inputs from wetlands	++ Inputs from wetlands
	++ Dam outflows	++ Dam outflows	0/- Dam outflows
	+ Allochthonous OM ^c	0/+ Allochthonous OM	+ Allochthonous OM
	+ Temperature	+ Temperature	+ Temperature
	+ Discharge	? Discharge	+ Discharge
	+ O ₂	-- O ₂	? O ₂
	+ Seasonal drying	+ Seasonal drying	0 Seasonal drying
Lakes	+ Allochthonous OM	+ Allochthonous OM	+ Allochthonous OM
	- Productivity	++ Productivity	+ Productivity
	- Depth and Surface Area	-- Depth and Surface Area	++ Nitrogen load
	+ Temperature	++ Temperature	? Depth and Surface Area
	+ O ₂	-- O ₂	+ Temperature
Reservoirs	As lakes, plus ++ OM in flooded soils	As lakes, plus ++ OM in flooded soils	As lakes, plus ++ OM in flooded soils
	+ Turbine degassing	++ Turbine degassing	0 Turbine degassing
	+ Drawdown areas	? Drawdown areas	0 Drawdown areas
			? O ₂

Note. ++, Strong increasing effect; + Increasing effect; 0, No net-effect; -, Decreasing effect; ? Unknown or ambiguous effect.

^aMeaning of effects. ^bGW = groundwater. ^cOM, organic matter.

spring freshet, lake-turnover, and extreme events like floods for which observations are generally rare. Only one of the studies included in our synthesis actually accounts for seasonality (Liu et al., 2022), while the other studies completely ignore seasonality in hydrodynamics, including spring freshet. Further, the estimates of lake and reservoir CO₂ emissions synthesized in our study do not account for contributions during lake-turnover, when emission rates are thought to be highest in boreal to Arctic systems (Sepulveda-Jauregui et al., 2015). Intense emissions have also been reported for periods of ice melt, during which observations are usually rare (Denfeld et al., 2018).

Raymond et al. (2013) account for intermittent drying of streams and rivers by decreasing the annual emission flux relative to the no-flow period. However, existing studies showed that during dry periods, exposed beds might show even higher CO₂ emission rates than from the water surface when inundated (Keller et al., 2020). Marcé et al. (2019) even suggest that taking into account seasonal dry falling and rewetting of bed sediments, estimates of global inland water CO₂ emissions would need to be corrected upward by at least 10%. For reservoirs specifically, existing emission estimates might even be ~50% higher if emissions from dry falling drawdown areas were taken into account (Keller et al., 2021). On the other hand, no estimate of CO₂ emissions from occasionally or seasonally inundated floodplains are available at the global scale. More systematic investigations of flux rates from both temporally dry falling inland water beds and temporally inundated floodplains would help to refine estimates of inland water CO₂ budgets and to better integrate them into continual CO₂ budgets while avoiding gaps and overlaps with terrestrial and wetland ecosystems.

Though some studies (Deemer et al., 2016; DelSontro et al., 2018; Raymond et al., 2013) have linked CO₂ variability in lakes and reservoirs to predictors such as waterbody size, mean annual precipitation, and ecosystem productivity, the controls on within-system CO₂ spatial and temporal variations are not well understood and effective scaling relationships are still in need to better represent CO₂ evasion from lakes and reservoirs. Further, characterizations of spatial variability within water bodies are rather scarce, and the representativeness of the sampling site within an aquatic system is a large source of uncertainty (Colas et al., 2020). Finally, our estimates of reservoir CO₂ emissions do not account for fluxes from hydroelectric turbines and dam outlets, where deep, hypolimnetic water enriched in CO₂ is released (Figure 2a). River reaches directly downstream of dams have been

reported to show increased $p\text{CO}_2$ while this excess CO_2 is being emitted rapidly over a few tenths of river-km (Calamita et al., 2021; Guérin et al., 2006; Teodoru et al., 2015). However, more systematic observations from these parts of the inland water network are needed to quantify this source of CO_2 flux at global scale, and to complete the assessment of reservoir CO_2 emissions.

3.2.2. Spatial and Temporal Resolution

Spatially and temporally resolved estimates of inland water GHG emissions at global scale can help to better understand the role of these fluxes in the overall GHG budget, to include these fluxes in regional budgets, and to evaluate them directly against observations. The realization of spatially and temporally resolved estimates is, however, limited by the availability of observations and using the utilized estimation techniques. For rivers, global empirical, spatially explicit estimates have already been achieved at several different resolutions, specifically, 231 regions (Raymond et al., 2013), gridded at 0.5° (Lauerwald et al., 2015), and for individual river reaches (Horgby et al., 2019; Liu et al., 2022). For lakes and reservoirs, the regionalized estimate based on 231 regions by Raymond et al. (2013) is the only existing spatially explicit estimate of CO_2 emissions at the global scale. Hastie et al. (2018) achieved a spatially explicit, pan-boreal estimate of lake and reservoir CO_2 emissions at 0.5° resolution. As the only process-based model approach at global scale, spatially explicit simulations with the land surface model DLEM have been achieved at 0.5° resolution for rivers and reservoirs (Tian, Ren, et al., 2015). ORCHILEAK—the inland water branch of the land surface model ORCHIDEE has so far only been applied at the continental scale of Europe (Gommet et al., 2022) and in few large-scale basins across the world (Bowring et al., 2020; Hastie et al., 2021; Lauerwald et al., 2020).

With regard to temporal resolution, most of the empirical studies published so far represent climatologies of average annual fluxes, often without precise specification of the time frame covered by the observations (Regnier et al., 2022). The only exception at global scale is the study by Liu et al. (2022) which presents a climatology of average monthly emission fluxes from rivers, thus representing the typical seasonal cycle of riverine emissions. The process-based model DLEM simulates time-series of riverine and reservoir CO_2 emission which reflect both seasonal and interannual variability. In general, a physically based model approach appears to be the most promising strategy to obtain seasonal and interannual variations in response to climate variability, for present day but also for scenario-dependent future projections (Hastie et al., 2021; Lauerwald et al., 2020; Tian, Ren, et al., 2015).

While empirical studies have highlighted the importance of diurnal variation in water-air CO_2 exchange, temporal variations at this time-scale are not yet possible to include in estimates. Process-based models like DLEM (Tian, Ren, et al., 2015) or ORCHILEAK (Lauerwald et al., 2017) represent aquatic CO_2 emissions as net-emissions driven by allochthonous inputs of CO_2 and net-instream respiration. A simulation of the diurnal variations would however require the representation of autochthonous aquatic production, which is not yet possible.

3.2.3. Data Requirements

As for all GHGs, data required to improve inland water CO_2 emission estimates include in the first place direct observation of emission rates. Many earlier estimates relied heavily on partial pressures of CO_2 ($p\text{CO}_2$) calculated from pH and alkalinity (Lauerwald et al., 2015; Raymond et al., 2013), which has been demonstrated to be a significant source of error leading to an overestimation of $p\text{CO}_2$ particularly in freshwaters with low buffer capacity against acidification (Abril et al., 2015; Golub et al., 2017; Hunt et al., 2011; Liu et al., 2020). Liu et al. (2022), relied on direct $p\text{CO}_2$ observations and suggested that average $p\text{CO}_2$ in global streams and rivers obtained by Raymond et al. (2013) is 30% too high.

However, as alkalinity and pH are easier to measure, a vast amount of data is available from a large number of studies and in data sets from environmental agencies, with greater spatial and temporal coverage (Hartmann et al., 2014) than that of direct observation. Nevertheless, as potential biases are hard to correct at large scales, a clear preference should be given to directly observed $p\text{CO}_2$ values. Liu et al. (2022) synthesized 5,910 direct $p\text{CO}_2$ observations from 63 studies, which represents about the latest inventory of available data for stream and river systems at global scale. Delsontro et al. (2018) synthesized literature data for 7,824 lakes and reservoirs. The number of direct $p\text{CO}_2$ observations is still limited, but steadily growing.

Likely for logistical reasons, most observations are from developed countries which contribute most to the research of inland water GHG budgets. For this reason, systems from temperate climate regions are better represented than

tropical or high-latitude systems in remote areas (e.g., compare Deemer et al., 2016; Liu et al., 2022). However, it is these remote areas that play a potentially important role, considering the extensive lake areas in Boreal to Arctic regions, and the large river systems of the humid tropics. There has been notable progress in sampling tropical (Africa (Borges et al., 2015, 2019), Amazon (Abril et al., 2014), and SE Asia (Wit et al., 2015)) and high latitude systems (Siberia (Karlsson et al., 2021; Serikova et al., 2019), Alaska: (Sepulveda-Jauregui et al., 2015)). Despite these advancements, more observations from these poorly monitored areas would help to improve estimates of global inland water CO₂ emissions.

Further, small water bodies require more attention in sampling campaigns. Holgerson and Raymond (2016) have highlighted the potentially important contribution of small lakes and ponds to global inland water CO₂ emissions. However, a regionalized estimate was not yet possible as observations of emission rates are still scarce, and more importantly, as no spatially explicit data set yet exists that would represent such small water bodies. Datasets that present the smallest water bodies (<1 km²) reliably would help to better integrate these important CO₂ sources into regionalized, global estimates.

Finally, increasing the number, variety and representativeness of investigated systems is only one step to reduce uncertainties in large-scale estimates of inland water CO₂ emissions. Temporal and small-scale spatial variations with small stream networks (Natchimuthu, Sundgren, et al., 2017; Natchimuthu, Wallin, et al., 2017) and with lakes (Natchimuthu, Sundgren, et al., 2017; Natchimuthu, Wallin, et al., 2017) and reservoirs (Colas et al., 2020) are substantial, and the choice of one or few sampling locations and a limited measurement period lead to large uncertainties and may introduce biases in the flux estimate for the whole waterbody. Improved investigation of CO₂ budgets of single systems requires measurements at various locations within a stream network or water body. Further, for reservoirs, observations of turbine emissions and of the release of CO₂ rich bottom waters from dams are required to better constrain these important emission pathways that are often ignored in large-scale assessments. In addition, the observations should be taken over a time period long enough to assess seasonal and inter-annual variability, and at a high enough frequency to assess short-term variations, including diurnal variations. In particular, data sets covering longer time periods such as those assembled for the US (Jones et al., 2003), China (Ran et al., 2021) and the boreal biome (Lapierre et al., 2013) are crucially needed to evaluate the extent to which trends simulated by LSMs are realistic (Regnier et al., 2022). The development and deployment of automated data loggers is a promising strategy for achieving this objective (Bastviken et al., 2015).

4. Inland Water CH₄ Budget

4.1. Overview of Existing Estimates

Global estimates of aquatic CH₄ emission range from 1.5 to 30 Tg CH₄ yr⁻¹ for streams and rivers, from 8 to 151 Tg CH₄ yr⁻¹ for lakes, from 9.8 to 52 Tg CH₄ yr⁻¹ for reservoirs, and from 16 to 331 Tg CH₄ yr⁻¹ for estimates that lumped lakes and reservoirs together (Table 3, Figure 1). The range in these emission estimates is generally more dramatic than for either CO₂ or N₂O (see Sections 3 and 5, respectively), with the exception of global CO₂ emission estimates from rivers and streams.

Some of the variation in global CH₄ emission estimates is due to large differences in the waterbody surface areas applied. For example, the earliest estimate of CH₄ emissions from reservoirs used a very rough estimate of surface area, multiplying the surface area of reservoirs in the *World Register of Dams* by a factor of four under the assumption that this would better represent the total surface area including small reservoirs and farm ponds not included in that register (St. Louis et al., 2000). This approach resulted in a surface area that is approximately three times larger than any subsequent estimate. Conversely, the earliest estimate from streams and rivers was conservative in that it applied a surface area for larger rivers only (quantifiable from global maps as the GLWD; Bastviken et al., 2011), resulting in approximately a factor of two reduction compared to subsequent estimates that also account for smaller rivers and streams. While most global estimates have ignored ice cover, Johnson et al. (2021) produced an estimate of reservoir emissions that accounted for this effect and which resulted in a CH₄ emission of 10 Tg CH₄ yr⁻¹ that is half or less of any previous assessment. Harrison et al. (2021) incorporated ice cover correction into their global reservoir emission estimate, resulting in similarly low emissions from reservoir surfaces (9.8 Tg CH₄ yr⁻¹), but still yielded a higher total flux due to the inclusion of reservoir turbine degassing (22 Tg CH₄ yr⁻¹). Rosentreter et al. (2021) also incorporated an ice cover correction into their assessment of global river (30 Tg CH₄ yr⁻¹), lake (151 Tg CH₄ yr⁻¹), and reservoir (24 Tg CH₄ yr⁻¹) emissions

Table 3
Existing Global Estimates of Lake, Reservoir, and River CH₄ Flux

References	$\Sigma\text{CH}_{4\text{em}}/\Sigma A_{\text{water}}$ (g CH ₄ m ⁻² yr ⁻¹)	ΣA_{water} (10 ⁶ km ²)	$\Sigma\text{CH}_{4\text{em}}$ (Tg CH ₄ yr ⁻¹)	Method
Rivers				
Bastviken et al. (2011)	4	0.36	1.5	Avg. rates
Stanley et al. (2016)	41.4 ^a	0.65	27 ^a	Avg. rates
Rosentreter et al. (2021)	66.5	0.77	30	Avg. rates
Lakes and Reservoirs				
DelSontro et al. (2018)	61.7	3.23–5.36	199–331	Avg. rates
-"	33.7	4.42	149 (95–236) ^b	Statistical prediction
-"	34.5	5.36	185 (119–295) ^b	Statistical prediction
-"	32.2	3.23	104 (67–165) ^b	Statistical prediction
Holgerson and Raymond (2016)	2.7 ^a	5.98	16 ^a	Avg. rates
Stavert et al. (2022)	32.4	2.93	95	Statistical prediction
Lakes (including lakes with dams)				
Bastviken et al. (2004)	0.12–122.9	2.8	8–48	Avg. rates
Bastviken et al. (2011)	19.2	3.7	72	Avg. rates
Rosentreter et al. (2021) ^c	54.1	3.71–5.69	151	Avg. rates
Johnson et al. (2022) ^c	15	2.8	42 ± 18 ^d	Model
Reservoirs				
St. Louis et al. (2000)	35	1.5 ^c	52	Avg. rates
Bastviken et al. (2011)	40.1	0.5	20	Avg. rates
Deemer et al. (2016)	58.5	0.31	17 (12–30) ^b	Avg. rates
Rosentreter et al. (2021) ^c	63.8	0.26–0.58	24	Avg. rates
Harrison et al. (2021) ^c	28.3(62.9) ^c	0.35	9.8 (22) ^c	Model
Johnson et al. (2021) ^c	33.3	0.3	10	Model

Note. For each estimate, the total water surface area (ΣA_{water}), the total CH₄ emission flux ($\Sigma\text{CH}_{4\text{em}}$) and the area weighted average emission rate ($\Sigma\text{CH}_{4\text{em}}/\Sigma A_{\text{water}}$) are reported.

^aOnly diffusive emissions. ^bLower and upper 90% (Deemer et al., 2016) or 95% (DelSontro et al., 2018) CI. ^cEstimate accounts for effects of seasonal ice cover. ^dStandard error. ^eIncludes emissions from turbines.

(estimates upscaled from mean emission rates), but in addition also an ice melt overturn correction that reduced the impact of ice cover. Moreover, their corrections did not result in a substantial lowering of the global flux due to increases in the magnitude of areal emission rates applied. The mean areal emission rates applied to upscaling efforts vary by approximately 2, 3, and 10-fold for reservoirs, lakes, and rivers respectively. In general, there is a temporal trend wherein older data sets have lower average areal emission rates than newer data sets. Part of this trend is due to the treatment of ebullition measurements in older emission estimates. Some global estimates summarized diffusive-only estimates of methane emission (Holgerson & Raymond, 2016; Stanley et al., 2016) while others combined diffusive only areal fluxes with ebullitive + diffusive estimates without differentiating one from the other (St. Louis et al., 2000). More recent estimates (Johnson et al., 2021, 2022; Rosentreter et al., 2021) only included studies that estimated both ebullition and diffusion together. Increasing average areal emission estimates may also be due to the increased likelihood of sampling right-skewed data as sample size for water bodies increases (see Wik et al., 2016). For example, a recent data set of lake and reservoir CH₄ emissions contains some of the highest mean areal fluxes, with about 65% of the estimates contained therein published since 2015 (Rosentreter et al., 2021).

Variation in binned areal emissions (e.g., by latitude, size, and chlorophyll-a) are even larger. For example, Rosentreter et al. (2021) reported an average areal CH₄ flux from the smallest lakes (<0.001 km²) that is nearly an order of magnitude higher than from lakes in the 0.1–1 km² size category, making these smallest systems responsible for 38% of the total lake CH₄ emissions (Rosentreter et al., 2021; Table 3). In addition, Bastviken et al. (2011)

reported areal reservoir CH₄ emissions from tropical regions that are an order of magnitude larger than in boreal regions (Bastviken et al., 2011), although follow-up work suggests that this discrepancy may have more to do with a lack of boreal ebullition estimates (Deemer et al., 2016) and the fact that latitude is only a weak predictor for reservoir CH₄ emission (Deemer & Holgerson, 2021; Johnson et al., 2021). Conversely, Rosentreter et al. (2021) report average areal CH₄ emissions from rivers that varied by a factor of about four by latitudinal bin, with the subtropical region (10–25° absolute latitude) producing the highest areal emissions and the temperate region (25–40° absolute latitude) producing the lowest areal emissions (Rosentreter et al., 2021).

While a variety of upscaling methods have been used to estimate inland water CH₄ emission, there does not appear to be any directional bias in the resulting estimates, that is one type of approach does not seem to systematically produce higher or lower emissions than other approaches. Many early estimates and some more recent estimates have applied the simplest empirical upscaling wherein a single areal flux was applied to a global surface area of lakes and/or reservoirs (Deemer et al., 2016; DelSontro et al., 2018; St. Louis et al., 2000), and rivers (Stanley et al., 2016). Other estimates have binned lakes and reservoirs CH₄ fluxes based on latitude (Bastviken et al., 2011), waterbody surface area (Bastviken et al., 2004; Holgerson & Raymond, 2016), primary productivity (e.g., chlorophyll a concentration; DelSontro et al., 2018), or has used some combination of these approaches (Rosentreter et al., 2021). For rivers, binning has so far only been based on latitude (Bastviken et al., 2011; Rosentreter et al., 2021). Finally, the most recent efforts to model lake and reservoir CH₄ flux have used a gridded approach that considers a variety of factors likely to influence the spatial variations in CH₄ emission including temperature, nutrients, and latitudinal variation in emission factors (Harrison et al., 2021; Johnson et al., 2021; Stavert et al., 2022).

4.2. Persisting Shortcomings and Future Challenges

4.2.1. Process Understanding

In the following we discuss our current process understanding of inland water CH₄ emissions in the light of global scale estimates. Figure 2 gives an overview of the main fluxes and processes involved, while Table 2 summarizes the known effects of the major environmental drivers. Significant progress has been made toward describing the drivers of lake and reservoir CH₄ flux, which may help improve our understanding of the spatial and temporal variability in emissions in the future. Specifically, small, shallow, productive, and low latitude lakes and reservoirs have been found to show higher areal methane emissions than larger, deeper, less productive, high latitude systems (Deemer & Holgerson, 2021). In northern systems, methane emissions are often further binned by lake type, with yedoma, peat, and glacial lakes exhibiting different patterns and magnitudes of emission (Kuhn et al., 2021; Matthews et al., 2020; Wik et al., 2016). Ebullition is usually reported to be the major emission path with flux rates nearly one order of magnitude higher than that of diffusion on average (Bastviken et al., 2011), though being less important in deeper parts of standing water bodies (>12 m depth in the study by Grinham et al. (2011)). CH₄ emissions from turbines and from anoxic hypolimnetic water released from dams have been shown to be important emission paths (Delwiche et al., 2022; Harrison et al., 2021; Teodoru et al., 2015), but are still ignored in most global scale assessments. Less is known about the key drivers of river CH₄ flux. Two of the three existing global estimates of river and stream CH₄ flux use latitude to bin emissions, but the latitudinal trend does not appear to describe much of the spatial variability (Rosentreter et al., 2021). The earlier data set compiled by Stanley and others contained many estimates from anthropogenically impacted rivers and streams (Stanley et al., 2016), and could be one explanation for the high global emission estimate despite only considering diffusive fluxes. While for many stream and river systems, CH₄ emissions seem indeed to be dominated by the diffusive path (Rovelli et al., 2022), ebullition has been reported to contribute substantially to stream CH₄ emissions at least locally (Crawford, Lottig, et al., 2014; Rovelli et al., 2022). Further, the effect of nutrient enrichment and productivity on river methane emissions has not been established the way it has been for lake and reservoir CH₄ emissions (Beaulieu et al., 2019). And even for lakes and reservoirs, empirical evidence for this connection between productivity and CH₄ emissions is limited ($R^2 = 0.38$ in Beaulieu et al., 2019).

Temperature is generally considered an important predictor of aquatic CH₄ emission and relationships between temperature and CH₄ flux have been used to scale seasonal emissions from reservoirs (Harrison et al., 2021; Johnson et al., 2021; Prairie et al., 2021). Such temperature-corrections address biases in many flux observations where measurements are focused during the spring-to-fall period whereas lower emissions during the ice-free winter period are typically not recorded. Temperature is considered a main driver of CH₄ production (Yvon-Durocher

et al., 2014), which limits both ebullitive and diffusive emissions. It was however found that ebullition responds much more intensely to temperature, in particular with regard to long-term trends. In a series of mesocosm experiments around the Northern Hemisphere, Aben et al. (2017) found that an increase in temperature by 4°C led to 51% higher annual ebullitive emissions, while diffusive emissions did not seem to be affected. While there is compelling cross-ecosystem evidence of increasing CH₄ emission with increasing temperature (Yvon-Durocher et al., 2014) there are also examples of systems where CH₄ oxidation is able to keep pace or surpass CH₄ production at higher temperatures (Duc et al., 2010; Shelley et al., 2015). A recent synthesis of CH₄ oxidation in lakes and reservoirs showed that CH₄ oxidation efficiency declines with ecosystem productivity (e.g., trophic status, D'Ambrosio & Harrison, 2021). In contrast, other studies have shown an increase in CH₄ oxidation with productivity (Grasset et al., 2020). Similarly, Sawakuchi et al. (2021) showed experimental evidence that CH₄ oxidation may be phosphorus-limited in northern lakes, also providing further evidence of more complex interactions between lake CH₄ dynamics and nutrient levels. The synergistic effects of productivity and temperature as driving factors of CH₄ emissions, and in particular ebullition as dominant emission path, have been shown in experimental studies (Davidson et al., 2015, 2018). Future work could improve our process understanding of methane emission dynamics by disentangling the role of temperature and productivity in driving both total emission and the balance between methane production and consumption.

Within a single waterbody, CH₄ emissions generally vary substantially in space and time (Wik et al., 2016), and this variation is likely more substantial than for either CO₂ or N₂O. This spatial and temporal variability has been shown to cause bias in upscaling, where too few measurements in either space or time can lead to underestimation of fluxes (Wik et al., 2016). While the regionalization exercise carried out in our companion paper (Lauerwald et al., 2023) begins to address seasonality by applying an ice cover and ice melt correction, future work should aim to better constrain temporal variability in methane fluxes within single water bodies. Temporal variability can arise from seasonal dynamics such as ice melt (Denfeld et al., 2018), fall turnover (Mayr et al., 2020), seasonal water-level changes (Varadharajan et al., 2010), or in response to phytoplankton blooms (Waldo et al., 2021). Diel variation can also be important. Daytime sampling might overestimate CH₄ flux in lakes (Sieczko et al., 2020), but may underestimate it in wetlands (Anthony & MacIntyre, 2016; Godwin et al., 2013; Poindexter et al., 2016). Episodic events can also be the source of large temporal variation such as water level drops in reservoirs (Harrison et al., 2017), storm-driven drops in hydrostatic pressure (Mattson & Likens, 1990) or increases in wind shear stress (Joyce & Jewell, 2003). In connection to seasonal or occasional water level drops, it was found that dry falling bed sediments tend to emit CH₄ at lower rates than when inundated, but are still a stronger source than upland soils (Paranaíba et al., 2022). For rivers, elevated discharge can lead to higher methane fluxes, especially in small high-gradient streams where methane is sourced predominantly from groundwater (Natchimuthu, Sundgren, et al., 2017; Natchimuthu, Wallin, et al., 2017). Spatial variability in aquatic CH₄ fluxes can arise for both biological and physical reasons. In lakes and reservoirs, the main drivers are the spatial variability of sedimentation of allochthonous and autochthonous organic matter (Maeck et al., 2013) and the reactivity of the sediment organic matter (Sobek et al., 2012; Wilkinson et al., 2015). Accordingly, higher fluxes are observed in inlets (DelSontro et al., 2011), near the shores (Natchimuthu et al., 2016; Peixoto et al., 2015) and behind run-of-river dams (Maeck et al., 2013). Further, spatial variability in CH₄ fluxes may arise from the heterogeneity of the sediment matrix and associated seeps (Walter Anthony & Anthony, 2013). In rivers, physical features such as waterfalls can be particularly important sites for CH₄ emissions (Natchimuthu, Sundgren, et al., 2017; Natchimuthu, Wallin, et al., 2017). At larger scales, high gradient headwater streams comprising <1% of catchment stream surface area can contribute 30% of catchment emissions, emphasizing the need to sample throughout a catchment rather than attempting to capture network-wide flux via single measurements at river mouths (Natchimuthu, Wallin, et al., 2017).

4.2.2. Spatial and Temporal Resolution

At global scale, gridded estimates of inland water CH₄ emissions exist for reservoirs (Johnson et al., 2021), lakes (Johnson et al., 2022) and lakes and reservoirs (Stavert et al., 2022). For rivers, disaggregating global fluxes over broad latitudinal zones (Bastviken et al., 2011; Rosentreter et al., 2021) still seems to be the best possible practice. Most existing global estimates for lakes and rivers represent climatologies of annual fluxes that do not resolve the seasonal and interannual variability, and longer-term trends. Using a relatively simple, process-based model, Johnson et al. (2021, 2022) were able to represent the seasonality in lake and reservoir CH₄ emission forced by temperature and ice-cover as drivers. Long-term trends in lake CH₄ emissions due to climate change have been predicted for the hol-arctic/boreal region using a more complex process-based model (Tan &

Zhuang, 2015a, 2015b). Other complex, process-based models of lake CH₄ cycling have been developed (e.g., Lake 2.0, Stepanenko et al., 2016), but have not been applied at large-scales. A recent study has hinted at the potential of exploiting global lake physical models to estimate changes in lake CH₄ cycling (Jansen et al., 2022). Combining simulated lake temperature profiles with an empirical equation linking sediment CH₄ production rates and temperature (Yvon-Durocher et al., 2014), Jansen et al. (2022) predicted relative increases in CH₄ production rates around the globe. In contrast to CO₂ and N₂O, no efforts to model river CH₄ emissions at regional to global scales have been published yet, which may partly be due to the relative small role of rivers in inland water CH₄ emissions as well as to the complexity of processes involved and the scarcity of data for model calibration and evaluation.

4.2.3. Data Requirements

One critical uncertainty for the inland water CH₄ budget is the inability to resolve the location and total surface area of the smallest lakes and impoundments. Waterbodies <0.001 km² have been recently estimated to comprise 37% of the lentic methane flux (Rosentreter et al., 2021). Along the same lines, Grinham et al. (2018) have estimated for Australia that CH₄ emissions from impoundments <0.1 km² equal about 10% of the national land use, land use change and forestry sector emissions. Given high variability in areal emissions from these smallest water bodies it is also important to increase effort in sampling these systems to reduce uncertainty. In addition to very small lakes and impoundments, sampling effort should be increased for large lakes (>1 km²) (Deemer & Holgerson, 2021). Given the additional importance of depth and productivity in regulating lentic CH₄ flux, spatially resolved information about the depth, chlorophyll *a*, and oxygen concentrations as well as quality and quantity of deposited organic matter in lakes and reservoirs will also help improve regional and global budgets (and overall upscaling efforts). More generally, systematic, long-term monitoring programs are needed which account for the high spatio-temporal variability in areal emission rates, in particular for ebullition, to better constrain the emissions even for individual, monitored systems. Long time-series of observations may finally help to better constrain the evolution of CH₄ production and emission in response to environmental change and climate extremes like droughts and heatwaves. This need for more and better observational data can hardly be satisfied with conventional methods, but would require the deployment of automatized observation systems and the use of remote sensing data, for which more research and development is needed. For reservoirs, finally, more observations of CH₄ emissions from turbines and of downstream release of CH₄ rich waters from dams are required to better constrain these important emission pathways, and to better assess the full impact of river damming on the CH₄ budget.

For rivers, many estimates of CH₄ emission rely on pairing concentration data with estimates of gas transfer (k_{GHG}) especially in low order streams (see Section 2.1 for further discussion). These low order systems have been observed to contribute disproportionately to CH₄ emissions at the catchment network scale despite very low CH₄ concentrations (Natchimuthu, Sundgren, et al., 2017; Natchimuthu, Wallin, et al., 2017), highlighting the need to constrain local values of k_{GHG} and/or perfect a universal physical model. Further, while most observations of stream CH₄ emissions concentrate on the diffusive pathway, ebullitive emissions can be important locally in small streams (Crawford, Stanley, et al., 2014). The development and application of empirical methods to directly measure the total GHG flux from low order streams would help constrain emissions from these systems.

5. Inland Water N₂O Budget

5.1. Overview of Existing Estimates

N₂O emissions from inland waters are poorly constrained at the global scale, which is visible in the largely divergent global estimates listed in Table 4 and shown in Figure 1: 0.05–3.3 Tg N₂O yr⁻¹ for streams and rivers and 0.1–0.6 Tg N₂O yr⁻¹ for lakes and reservoirs. Most existing global estimates of riverine N₂O emissions are based on modeled N loads from watersheds and emission factors (EFs), in stark contrast to CO₂ and CH₄ global estimates, which are calculated mainly by empirically upscaling local observations. N₂O is produced as an intermediate product in denitrification, that is, the reduction of nitrate to N₂, but also as a by-product in the process of nitrification, that is, the oxidation of ammonium to nitrate (Canfield et al., 2010). The amount of N₂O produced and emitted due to these processes depends on environmental and hydrological factors including water temperature, N availability and speciation, water body depth, oxygen availability, pH, and labile carbon concentrations (Clough et al., 2007; Hu et al., 2019; Outram & Hiscock, 2012; Rosamond et al., 2012; Venkiteswaran

Table 4
Global Scale Estimates of Inland Water N₂O Emissions

References	$\Sigma N_2O_{em} / \Sigma A_{water}$ (mg N ₂ O m ⁻² yr ⁻¹)	ΣA_{water} (10 ⁶ km ²)	ΣN_2O_{em} (Gg N ₂ O yr ⁻¹)	Method
Rivers				
Seitzinger and Kroeze (1998) and Seitzinger et al. (2000)			1,650 (300–2,940) ^c	Emission factors
Kroeze et al. (2005)			1,975	Emission factors
Mosier et al. (1998)			1,100	Emission factors
De Klein et al. (2006)			550	Emission factors
Kroeze et al. (2010)			470–3,300	Emission factors
Beaulieu et al. (2011)			1,070	Emission factors
Hu et al. (2016)			51 (19–105) ^a	Statistical prediction
Maavara et al. (2019)			72–78 ^c	Model
Yao et al. (2020)			458 ± 92 ^b	Model
", stream orders 1–3			387 ± 93 ^b	Model
", stream orders ≥ 4			71 ± 23 ^b	Model
Marzadri et al. (2021)			114	Machine learning + Model
", stream orders 1–3			76	Machine learning + Model
", stream orders ≥ 4			38	Machine learning + Model
Lakes and reservoirs				
DelSontro et al. (2018)	78	3.23–5.36	252–424	Avg. rates
DelSontro et al. (2018)	106	4.42	470 (300–710) ^a	Statistical prediction
DelSontro et al. (2018)	112	5.36	600 (380–860) ^a	Statistical prediction
DelSontro et al. (2018)	127	3.23	410 (250–600) ^a	Statistical prediction
Soued et al. (2016)	235	4.20	985 ± 465 ^b	Avg. rates
Lauerwald et al. (2019)	34	2.93	98 ± 64 ^c	Model
Lakes (including lakes with dams)				
Lauerwald et al. (2019)	17	2.68	46 ± 29 ^c	Model
Reservoirs				
Deemer et al. (2016)	152	0.31	47 (31–110) ^a	Avg. rates
Maavara et al. (2019)	148–250 ^c	0.45	67–112 ^c	Model
Lauerwald et al. (2019)	185	0.25	52 ± 33 ^c	Model

Note. For each estimate, the total water surface area (ΣA_{water}), the total N₂O emission flux (ΣN_2O_{em}) and the area weighted average emission rate ($\Sigma N_2O_{em} / \Sigma A_{water}$) are reported.

^aLower and upper 90% (Deemer et al., 2016; Lauerwald et al., 2015; Raymond et al., 2013) or 95% (DelSontro et al., 2018) CI. ^bStandard error. ^cMin and max estimate.

et al., 2014). EFs can be defined as average ratios of N₂O emission to denitrification and nitrification fluxes. However, it is difficult to quantify nitrification and denitrification fluxes for entire river systems, and even more so at the global scale. Therefore, EFs have traditionally been established by linking N₂O emissions directly to riverine N loads, implicitly assuming a certain fraction of riverine N loads to be nitrified and denitrified. Mosier et al. (1998) assumed that N leached to the river network was denitrified once and nitrified twice along the river network. Further assuming that 0.25% of both nitrified and denitrified N is emitted as N₂O, they concluded that 0.75% of the total N leached to the river is emitted as N₂O. Applying that percentage as EF directly to riverine N loads, they estimated a global riverine N₂O emission of 1.1 Tg N₂O yr⁻¹. The methodology and EFs established by Mosier et al. (1998) also served to assess the river N₂O emissions in the 5th Assessment Report of the IPCC (2013).

In a similar approach, Seitzinger and Kroeze (1998) and Seitzinger et al. (2000) estimated N₂O emissions from only the dissolved inorganic fraction (nitrate, nitrite and ammonium) of N (DIN) leached to rivers. Applying

EFs of 0.3% and 3% relative to riverine DIN load they estimated a global riverine N_2O emission of 1.7 (range 0.3–2.9) Tg $\text{N}_2\text{O}/\text{yr}$. Over the following decade, these two EF approaches, that is, the one of Seitzinger and Kroeze (1998) and the IPCC approach derived from Mosier et al. (1998), were updated, yielding consistently large emission fluxes. Kroeze et al. (2005) estimated 2 Tg $\text{N}_2\text{O}/\text{yr}$, and later Kroeze et al. (2010) revised their estimate to 0.5–3.3 Tg $\text{N}_2\text{O}/\text{yr}$, both using modeled DIN loads and EFs of 0.3% and 3%. De Klein et al. (2006) predicted a global riverine N_2O emission of 0.6 Tg $\text{N}_2\text{O}/\text{yr}$, while Beaulieu et al. (2011) calculated an emission flux of 1.1 Tg $\text{N}_2\text{O}/\text{yr}$, both using the IPCC approach.

Studies conducted over the last 5–7 years (Hu et al., 2016; Maavara et al., 2019; Marzadri et al., 2021; Yao et al., 2020) consistently calculate N_2O emissions for rivers that are substantially lower than those of the decades before (Figure 1). Hu et al. (2016)'s empirical approach estimated global riverine N_2O emissions of 51 (19–105) Gg $\text{N}_2\text{O}/\text{yr}$. Further, the authors report EFs relative to riverine DIN loads of 0.16%–0.19% to be realistic, suggesting the EFs used by Seitzinger and Kroeze (1998) to be unrealistically high. Maavara et al. (2019)'s spatially resolved stochastic-mechanistic river-continuum model is the first to explicitly represent N transformation processes, and results agreed well with Hu et al.'s predictions, with a global flux of 72–78 Gg $\text{N}_2\text{O}/\text{yr}$. Moreover, Maavara et al. (2019) estimated that only 7% and 9% of the total N loads are respectively denitrified and nitrified in the global river network. Thus, the assumption behind the IPCC AR5 approach that all N leached to rivers is once denitrified and twice nitrified also appears to be unrealistic and responsible for gross overestimations.

The studies by Yao et al. (2020) and Marzadri et al. (2021) are complementary as they provide estimates that also account for small streams that contribute disproportionately to the overall riverine N_2O emissions, but which were ignored in earlier estimates. Marzadri et al. (2021), using a machine learning based approach, reach an estimate of about 114 Gg $\text{N}_2\text{O} \text{ yr}^{-1}$, of which about half is contributed by headwater streams. Note that this is only a near-global estimate which excludes high latitudes $>60^\circ \text{ N}$, which can however be assumed to be small contributors to the global emission due to low N loads of the corresponding river systems (Maavara et al., 2019). Yao et al. (2020), using the land surface model DLEM, estimate riverine N_2O emissions at even higher values of $458 \pm 92 \text{ Gg } \text{N}_2\text{O} \text{ yr}^{-1}$, of which 80% stems from small stream emissions up to stream order 3. In their simulations, emissions from these small streams are largely fed by N_2O inputs from groundwater and saturated soils, which are not accounted for in the other studies. Marzadri et al. (2021), although not representing groundwater N_2O inputs, still estimate that about $\frac{2}{3}$ of total riverine N_2O emissions is contributed by small streams of orders 3 and lower (see Table 4). In these small streams, nitrification-denitrification processes occur mainly within hyporheic and benthic zones, whereas in larger rivers, the contribution of water column exceeds that of subsurface environments in contributing to N_2O production (Marzadri et al., 2021). The estimates for larger rivers only by Yao et al. (2020) and Marzadri et al. (2021) agree better with those by Hu et al. (2016) and Maavara et al. (2019).

For lakes and reservoirs, the first global estimates were only published recently. Soued et al. (2016) and DelSontro et al. (2018) gave estimates for the entirety of lakes and reservoirs, without distinguishing between both types of systems while Deemer et al. (2016) estimated N_2O emission from reservoirs only. Maavara et al. (2019), in their stochastic-mechanistic model of N_2O emissions from river networks, included explicit emission estimates for reservoirs. Lauerwald et al. (2019) then adapted that model to estimate N_2O emission from both reservoirs and lakes. Soued et al. (2016) performed a simple upscaling based on averaged observed N_2O emissions rates for three latitudinal zones, which yielded with $985 \pm 465 \text{ Gg } \text{N}_2\text{O} \text{ yr}^{-1}$ the highest of the emission fluxes from lakes and reservoirs listed in Table 4. A major limitation of this study was the poor global coverage of observations. While they used data from 298 systems worldwide, they had observations from only six systems for their low latitude estimate, all belonging to the reservoir-class from the study of Guérin et al. (2006). This fact is critical as in their upscaling, lakes and reservoirs from that zone contributed about 80% of their global estimate of N_2O emissions. Moreover, some of these reservoirs showed extremely high emission rates due to the fact that soils and biomass had not been removed before dam closure, which contributed massively to GHG production and emission (Guérin et al., 2006). It is thus highly probable that these reservoirs are not representative for low latitude lakes and reservoirs, as later discussed in detail in Lauerwald et al. (2019).

Similar to their estimates of lake and reservoir CO_2 and CH_4 emissions (see Sections 3 and 4), DelSontro et al. (2018) followed two distinct methodological approaches to obtain their global estimates: a direct upscaling approach based on a global average of observed N_2O emission rates ($252\text{--}346 \text{ Gg } \text{N}_2\text{O} \text{ yr}^{-1}$), and a statistical approach using lake/reservoir size classes and classes of chlorophyll-a concentrations as predictors

(409–597 Gg N₂O yr⁻¹). Note that the second approach did not lead to a spatially explicit estimate, as only global, statistical distributions of size classes and chlorophyll-*a* concentrations were used for upscaling. Further, the statistical upscaling equation had a very low predictive power with an *R*² below 0.1.

Deemer et al. (2016) performed a simple upscaling to estimate N₂O emissions from reservoirs only, obtaining a global flux of about 47 Gg N₂O yr⁻¹. Despite the very different approach, Maavara et al. (2019) and Lauerwald et al. (2019) estimated global N₂O emissions from reservoirs that are comparable to those by Deemer et al. (2016) (see Table 4). For the entirety of lakes and reservoirs, Lauerwald et al. (2019) by far the lowest global estimate of 99 ± 64 Gg N₂O yr⁻¹, and which is only about one tenth of what was estimated by Soued et al. (2016). Comparing their spatially explicit estimate to regional estimates based on direct upscaling, Lauerwald et al. (2019) found that their model results are reasonable. Moreover, they estimated that lakes, although contributing more than 90% of the global surface area of standing water bodies, contribute only about half of the emission flux as a result of their much lower average emission rates. This indicates that it is problematic to lump together lakes and reservoirs in global upscaling exercises.

5.2. Persisting Shortcomings and Future Challenges

5.2.1. Process Understanding

In the following we discuss our current process understanding of inland water N₂O emissions in the light of global scale estimates. Figure 2 gives an overview of the main fluxes and processes involved, while Table 2 summarizes the known effects of the major environmental drivers. While a basic understanding of processes involved in aquatic N₂O cycling exists from a certain number of field studies, the quantification of these processes in large-scale estimates is still difficult due to their complexity and the unavailability of sufficient data sets to support their assessment. For this reason, empirical EFs have long been used to estimate riverine N₂O emissions directly from N loads, assuming a constant fraction of N loads to be nitrified and denitrified within the rivers, independent of the size of the river network and its ecoclimatological setting. While newer, model-based studies proved the worth of calculating more precise estimates of nitrification and denitrification fluxes taking into account physical constraints such as water residence time and temperature (Maavara et al., 2019; Marzadri et al., 2021; Yao et al., 2020), the actual production and/or emission of N₂O related to these processes is still based on simple, empirical factors. As N₂O is formed only as a by-product of nitrification and as an intermediate product in the denitrification process, the actual fraction of N₂O produced from these processes is highly variable and not yet possible to reproduce based on mechanistic formulations.

A better assessment of inland water N₂O cycling would require the representation of the vertical profile of oxygen concentrations through the water column and benthic sediments. Of particular importance is the position of the oxycline, that is, the rather narrow zone of steep decrease in oxygen concentrations, separating an oxic upper layer, where nitrification is the dominant process, from an anoxic lower layer, where denitrification dominates. The oxycline itself is a hot-spot of N₂O production because here, anoxic water rich in ammonium mixes with oxygen-rich waters, promoting nitrification, while in turn nitrate produced from nitrification diffuses down and fuels denitrification in the anoxic zone (Beaulieu et al., 2015). Depending on water depth, water column mixing and sediment oxygen consumption, the oxycline lies either in the water column or sediment column. Following the conceptual model by Marzadri et al. (2017, 2021), the position of the oxycline, and thus the relative importance of water column versus sediment processes in N₂O production, changes along the river network. In headwaters, the oxycline is situated in the bed sediments and nitrification of emergent, ammonium-rich groundwater in streambed sediments is the dominant source of N₂O. The importance of nitrification decreases downstream while the oxycline moves up from the sediment into the water column, until finally denitrification in the lower water column is the dominant source of N₂O. Further, it was shown that dissolved N₂O inputs from groundwater and waterlogged soils feed an overproportional contribution of headwaters to riverine N₂O emissions (Billen et al., 2020; Yao et al., 2020). Note that the global assessments of river N₂O emissions by Marzadri et al. (2021) and Maavara et al. (2019) do not account for groundwater N₂O inputs which represent an important part of inland water N₂O emissions.

For lakes and reservoirs, the importance of processes in the benthic zone has been implicitly taken into account by the use of “hydraulic load” to scale denitrification rates (Harrison et al., 2009). Hydraulic load has been defined as the ratio of water inflow to water surface area, which is identical to the ratio of average lake or reservoir

depth over water residence time (Harrison et al., 2009). The process of denitrification is assigned an “apparent settling velocity” which expresses rates of nitrification or denitrification in the benthic zone relative to water column depth. The deeper the average lake or reservoir, the longer it takes until the whole volume is nitrified or denitrified. However, this approach does not take into account the actual shape of the lake/reservoir bed and the proportions of shallow, littoral zones, where the oxycline and thus main source of emitted N_2O lies in the bed sediments (Liikanen et al., 2003; Zhu et al., 2015), versus the deeper zones, where processes in the water column are the dominant source of N_2O (Mengis et al., 1997). While streams and rivers are usually well mixed, deeper lakes and reservoirs may be temporally stratified, with a pronounced oxycline within the water column—and important consequences for N_2O cycling, which have so far not been taken into account in large-scale assessments. During stratification, only the top layer (epilimnion) is exchangeable with the atmosphere and thus well oxygenated. Then, nitrification in the epilimnion is the main source of N_2O emissions (Beaulieu et al., 2015; Mengis et al., 1997). In anoxic parts below the oxycline (hypolimnion), denitrification prevails, which can be a source or sink of N_2O , depending on the availability of nitrate for reduction (Beaulieu et al., 2015; Mengis et al., 1997). As this anoxic water may also be rich in ammonium from the in situ decomposition of organic matter, mixing with more oxygenated, epilimnetic waters during lake turn-over may represent a “hot moment” for nitrification and N_2O emissions (Beaulieu et al., 2015; Roland et al., 2017). However, a quantitative assessment of this hot-moment at large scales is not yet possible due to the lack of observational data. Moreover, while a certain number of studies report measurements of N_2O concentrations in the shallow, easy to reach epilimnion, studies investigating the deeper profile of N_2O concentrations through the hypolimnion are scarce (Beaulieu et al., 2015; Mengis et al., 1997).

Further, also resolving the horizontal zonation would help to better assess the overall N_2O budget of a lake. Within larger lakes, shallow littoral zones have been shown to contribute disproportionately to lake N_2O emissions relative to their areal extent (Zhu et al., 2015). Here, benthic sediments contribute most to N_2O production, while in the deeper, pelagic zone, N_2O is produced in the water column, and more specifically, under stratified conditions with a pronounced oxycline, in the epilimnion (Liikanen et al., 2003; Mengis et al., 1997). Yet, most observations are constrained to pelagic zones, which dominate lakes and reservoirs with regard to surface area, but not necessarily emissions. Further, strong horizontal gradients in N_2O emissions rates may be formed towards the points of riverine inflows of reactive N (Miao et al., 2020). However, few studies conduct systematic sampling which could reveal and account for these internal spatial variations. In contrast to for instance CH_4 or CO_2 , drawdown areas of reservoirs do not appear to emit more N_2O than upland soils (Hao et al., 2019). Also, seasonal streambed drying was not found to increase riverine N_2O emissions (Tonina et al., 2021).

In general, observational studies are skewed towards temperate, eutrophic systems in developed countries, which are easily accessible for sampling and which represent potentially important N_2O sources related to water quality issues caused by agricultural non-point sources and sewage water injections. In boreal regions where N loads are usually lower, it was demonstrated that a substantial proportion of aquatic systems is undersaturated with N_2O and thus rather act as sinks for this GHG (Kortelainen et al., 2020; Soued et al., 2016). Further, as for CO_2 and CH_4 , observations of smaller water bodies are generally underrepresented. Interestingly, though small, agricultural ponds could be hypothesized to be strong GHG emitters, a study of 101 such systems across Canada (Webb et al., 2019) has shown that about two thirds of these systems are on the contrary N_2O sinks. In conclusion, global inland water N_2O emissions may have been overestimated due to the bias in observed systems outlined above. Moreover, most estimation approaches, particularly the use of EFs, do not permit for representing inland waters as N_2O sinks. Finally, samples from temperate and high latitude systems are skewed towards summer months, while the full seasonal cycle is only rarely covered in observational studies. Kortelainen et al. (2020) have demonstrated in their study on Finnish lakes that there is a strong seasonality in N_2O concentrations and emission rates, with much higher values in winter when low autotrophic production allows for higher nitrate concentrations. A flux estimate based on summer-time observations only would thus have led to an underestimation by a factor of four. Assuming that similar seasonal patterns are to be observed in other temperate to high latitude systems, and that in particular lake turnover as hot moment of N_2O emissions is not well captured in observations, we can hypothesize that the non-representativeness of sampling times might have introduced a negative bias in upscaling exercises. Note finally that the non-representativeness of observations does not only affect estimates based on direct upscaling of average emission rates. With the lack of representative observational data for calibration and validation, also model-based studies will remain of limited validity.

5.2.2. Spatial and Temporal Resolution

For riverine systems, global, spatially explicit estimates of N_2O emissions have been achieved by numerous studies. The spatial resolution ranges from large river basins (Hu et al., 2016), over gridded estimates (Yao et al., 2020) to estimates per stream segment (Marzadri et al., 2021). For lakes, Soued et al. (2016) have resolved N_2O emissions for three broad latitudinal bands and the only published spatially explicit, gridded estimate is that of Lauerwald et al. (2019). Most of these studies represent a climatology of average annual fluxes. Only the process-based model by Yao et al. (2020) allows for spatio-temporally resolved simulation results which cover seasonality, interannual variability and long-term temporal trends. So far, this model includes rivers and reservoirs. For lakes, a global scale, process-based model that permits for temporally varying N_2O emissions is still missing. In line with what was discussed in the preceding subsection, such a model would need to couple lake physics and biogeochemistry.

5.2.3. Data Requirements

To achieve better global estimates of inland water N_2O emissions, more observational data is needed, in particular from high latitude and tropical areas. Also, small lentic water bodies, including natural and farm ponds, and ponds used for aquaculture, are so far undersampled. Finally, the bulk of available observation data is biased by a tendency to study eutrophic systems that promise high emission rates, while oligotrophic systems that may even be sinks are underrepresented (Soued et al., 2016). In general, more systematic observational programs permitting the quantification of seasonality and the impact of seasonal ice cover, lake turn-over and algae blooms to annual emissions are needed to avoid biased upscaling of annual flux estimates. Finally, long-time series are needed to assess the long-term evolution of inland water N_2O emissions and to evaluate process-based models.

To support the application of more advanced upscaling approaches in the estimation of inland water N_2O budgets, including process-based models, better data on environmental drivers and boundary conditions are required. That includes the representation of reactive N species to inland waters. While global estimates of total N and DIN inputs to the river network exist (Mayorga et al., 2010), it would be even better to have information on the more specific inputs of nitrate, ammonium, and dissolved N_2O to set the boundary conditions for processes involved in N_2O production, reduction and emissions. Further, similar to what was pointed out for CH_4 (Section 4.2.3), the model representation of N and N_2O cycling in inland water would profit from data sets on bed morphology of the water body and properties of bed sediments. Marzadri et al. (2021) have used a machine learning approach to estimate all these boundary conditions for the application of their model of stream N_2O production. While this seems a promising strategy, this approach could be steadily improved with new findings from field observations and improved data sets of predictor variables. Also for lakes and reservoirs, only estimates of volume and average lake depth are available (Messenger et al., 2016), which could be steadily improved using a similar strategy. Finally, for the better assessment of lake and reservoir N_2O budgets, physical processes such as stratification, mixing and ice cover would need to be represented dynamically. For example, process-based models of lake physical processes have been developed and even implemented into land surface models for global scale application (Subin et al., 2012) and the outputs of such models may be included in future lake and reservoir N_2O models.

6. Conclusions and Outlook

The number of global scale estimates of inland water GHG emissions is constantly increasing, at an accelerated step. For CO_2 and CH_4 , we see a tendency for increasing numbers in estimates of global scale fluxes following the inclusion of water bodies which contribute significantly to the overall water surface area, and disproportionately to overall emissions. For water bodies above a certain size (e.g., stream orders, lake size), estimates of average emission rates seem to converge in latest estimates. Major discrepancies persist however with regard to the assumed water surface area and the statistical distribution of water body size classes, in particular for small lakes and impoundments (<10 ha), and ponds. For riverine N_2O emissions on the contrary, we find newer estimates to be lower than older estimates, following a change in methodologies moving away from the application of emission factors toward more process oriented modeling. For lake N_2O emissions, discrepancies in assumed water surface area and distribution of lake size classes still play a role as well, but not as strongly as for CO_2 and CH_4 because small water bodies do not appear to contribute disproportionately to the total emission flux.

While uncertainties in global scale assessment of inland water GHG emissions persist, developments in monitoring and upscaling techniques are envisageable to overcome these uncertainties, as discussed in this review paper and summarized in Table A1. There is ongoing work to improve spatially explicit data sets of inland water surface

areas that will help improve global scale estimates. For streams and rivers, global scale estimates have recently been largely improved combining high resolution remote sensing of water surface areas and statistical prediction for headwater streams which are too narrow to be detected (GRWL—Allen & Pavelsky, 2018). A similar strategy may also be the solution for lakes and reservoirs, combining data from inventories and remote sensing. Inventories are more reliable but less comprehensive as they exclude smallest water bodies and are susceptible to geographical biases due to differences between national data sources. Remote sensing is able to detect smaller water bodies but is prone to contaminations with wrongly attributed water surface areas if unsupervised algorithms are applied and checks for ground truth in sufficient quantity and quality are not possible.

More importantly, improving inland water GHG emissions estimates requires more fieldwork to improve quantity and quality of observational data. In particular, we need more data from systems in remote areas of the high latitudes and the tropics, and systematic measurements with time-series of sufficient length and frequency of observations to better capture seasonal and inter-annual variability in fluxes as well as long-term trends in response to environmental change. In addition, more attention has to be paid to hot-spots and hot-moments of inland water GHG emissions, which likely contribute a substantial fraction of overall emissions.

For upscaling and predictions to achieve better global scale estimates, recent developments of machine learning-based approaches and process-oriented models seem promising. These approaches help to better constrain the spatial-temporal variability in global scale estimates, which allows to better include inland water GHG emissions in regionalized budget efforts such as RECCAP-2, but also in top-down approaches based on atmospheric inversions, further reducing uncertainties in global estimates.

Appendix A

Table A1 summarizes the main take home messages from our review.

Table A1

General Uncertainties and Knowledge Gaps That Persist With Regard to Regional to Global Scale Estimation of Inland Water GHG Emissions, and Recommendations for Monitoring and Upscaling

Uncertainty/knowledge gap	Recommendations for monitoring	Recommendations for upscaling
Source attribution of emitted GHGs difficult: produced in situ or imported from upstream, surrounding soils or groundwater?	Watershed scale monitoring with systematic observations of GHG sources and sinks in interconnected upland, wetland and inland water systems, and of lateral GHG transfers along terrestrial-aquatic continuum.	Process based models that represent sources and sinks of GHGs within the inland waters and their catchment, including the reactive transport along the terrestrial-aquatic continuum.
Smallest water bodies contribute overproportionally to CO ₂ and CH ₄ emissions, but are not well constrained yet at global scale.	More systematic observations of GHG emissions from small lakes, ponds and streams are needed.	Need for reliable spatial data sets of small inland waters from remote sensing with extensive ground truthing. Categories of climate and land use may be useful for upscaling.
Internal variability needs to be better constrained, especially in large waterbodies with diffuse inputs, and in reservoirs that flood heterogeneous landscapes.	Systematic observations along internal spatial gradients (along depth gradients, distance to shore, position to inflows) are required.	More detailed information on geometry of waterbodies, on inflows of water and sediment, and on reservoir management required to predict internal heterogeneity of GHG fluxes.
Temporal variability (diurnal, event-based, seasonal, interannual) and contributions from hot moments (e.g. spring freshet, flood events, ice-out, lake turn-over, algae blooms, reservoir drawdowns) are poorly constrained at global scale.	Time series of observations needed, which: - are long enough to capture inter-annual variability and the effects of climate extremes such as droughts and heat waves; - are frequent enough to capture hot moments (automatic sensors) -also cover night-time fluxes.	Statistic or process based models that use the main meteorological (temperature, radiation, wind speed, air pressure, precipitation) and hydrodynamic (fluctuations of water flows and water table, thermal stratification vs. mixing over lakes) drivers of the physical and biological processes behind inland water GHG dynamics.
Uneven geographic distribution of observations may lead to biases in upscaling. Higher uncertainties persist for high latitude, high elevation, arid, and tropical systems.	More long-term monitoring networks in remote tropical, arid, high elevation, and high latitude areas are required.	Upscaling techniques are required that bin data per climate zone, or use relevant climatic drivers as predictors or as input for process based models.
Contributions from ebullition versus diffusion, and other paths like emission from turbines not well constrained.	More systematic observations of inland water GHG budgets are needed that cover all important emission paths.	For upscaling, the different emissions paths need to be explicitly represented and their specific drivers taken into account.

Data Availability Statement

The data that support the findings of this study are available at figshare (<https://doi.org/10.6084/m9.figshare.22336465.v1>).

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