Lawns and meadows in urban green space — A comparison from perspectives of greenhouse gas, drought resilience and biodiversity perspectives

plant functional types

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Abstract. Nowadays, city planners design urban futures by considering climate change and biodiversity loss. Here, we studied the greenhouse gas fluxes of urban lawns and meadows and linked the observations with plant functional diversity types and soil properties. In eight lawns and eight meadows in the Helsinki Metropolitan Area, Finland, carbon dioxide (CO₂), methane (CH₄) and nitrous oxide (N₂O) fluxes, as well as plant diversity, were measured using manual chambers and plant functional types were recorded in eight lawn–meadow pairs in the Helsinki Metropolitan Area, Finland. Among the four of these sites, i.e., an irrigated lawn, an old mesic meadow, a non-irrigated lawn, and a young dry meadow and an old mesic meadow, were more intensively studied especially using manual chamber measurements in 2021–2022. The process-based ecosystem model JSBACH was utilized together with the momentary observations collected approximately every second week on CO₂ exchange to quantify the annual carbon (C) balance of these intensive study sites. On another hand, the remaining sites, we studied the initial dynamics of conversion from lawns to meadows were studied with the six other pairs, whereby transforming part of lawns to meadows in late 2020 and conducting measurements were conducted from 2020 to 2022 and the transformation of half of the sites had been performed in late 2020. We found that without considering the impact of management and unstabilized soil, lawns are clear sink of carbon, whereas sinks of carbon (NEE = −336 ± 187 and −157 ± 139 g CO₂ m⁻² yr⁻¹) compared with meadows (NEE = −151 ± 198 and −63 ± 134 g CO₂ m⁻² yr⁻¹), and the conversion from a lawn to a meadow did not affect the fluxes of CH₄ and N₂O. Moreover, the mesic meadow was more resistant to drought events than the non-irrigated lawn. Moreover, according to our. Lastly, the proportion of flowering plants (forbs, including legumes) was found to be higher in meadows than in lawns. Even though social and economic aspects also steer urban development, these results, the conversion from lawn to meadow did not affect the fluxes of CH₄ and N₂O. Last, the relation between C and N cycle and plant diversity was unclear and would need further investigations, can guide planning when considering biodiversity and carbon-smartness.

Keywords, carbon balance, photosynthesis, respiration, CH₄, N₂O, JSBACH, plant diversity, functional types, urban nature
1 Introduction

The impacts of climate change and biodiversity crises are increasingly evident. The crises have feedbacks with each other as climate change threatens biodiversity, and the loss of biodiversity and possible degradation of ecosystems affects the climate system. Therefore, it is crucial to understand the role that various ecosystems in the land-use sector play in regulating climate and supporting biodiversity, and to find land management practices that can effectively mitigate the climate and biodiversity crises simultaneously. Well-known management practices for the land use sector include, for example, conservation and restoration, and integrated land-use planning taking into account the relationships between different land uses (Niemelä et al., 2010; Pörtner et al., 2021).

Species richness can affect carbon sequestration potential in several ways. Generally, ecosystems with higher species richness are shown to have higher photosynthetic productivity than less diverse systems, because of the more efficient use of resources by the greater variety of species contributing to carbon uptake (Fornara and Tilman, 2008; Lange et al., 2015; Yang et al., 2019; Chen et al., 2020). In addition, increased species richness can lead to a more resistant functioning of the ecosystem in extreme weather events, as the responses by different species can respond differently to changes in environmental conditions (De Keersmaecker et al., 2016; Hossain et al., 2022). This means that, for example, during a drought some species may be more resistant, making the functioning of a diverse ecosystem less vulnerable compared to a habitat with lower species richness. However, the relationship between species richness, productivity and resistance is not straightforward, but influenced by many factors, such as ecosystem management, species cortege and environmental conditions (Vogel et al., 2012; De Keersmaecker et al., 2016; Jung et al., 2020).

In the urban context, Urban green spaces not only sequester atmospheric carbon (C) but provide vital ecosystem services, such as cooling, recreation, purification of air and water, and risk reduction for flooding (Niemelä et al., 2010; Belmeziti et al., 2018; Lampinen et al., 2021; Shen et al., 2023). Cities actively seek optimal green area planning and management practices to mitigate and adapt to climate change and there are also numerous initiatives aimed at promoting urban green spaces and infrastructure (Haaland and van den Bosch, 2015). One of those is Article 6 in the Nature Restoration proposal launched by the European Commission (European Commission, 2022), which sets targets for increasing green urban spaces in cities, towns and suburbs.

Within urban landscapes, lawns constitute one of the most common features of the green space (Hedblom et al., 2017; Ignatieva et al., 2020), and are usually subjected to frequent and intense management regimes to fulfill social, aesthetic and recreational purposes (Ignatieva et al., 2015, 2017; Zobec et al., 2020; Paudel and States, 2023). Although many types of lawns exist, most of the urban green spaces worldwide are dominated by turfgrass lawns. These lawns typically contain certain selected species of grasses, such as in Southern Finland Agrostis capillaris, Alopecurus pratensis, Dactylis glomerata, Festuca ovina, F. pratensis, F. rubra and Poa annua (Tonteri and Haila, 1990), yet they can also harbor other species of forbs and grasses that are spontaneously established, thus giving lawns the ability to behave like semi-natural grasslands (Thompson et al., 2004; Fischer et al., 2013). Reducing the conventional turfgrass lawn management has been proved to enhance the abundance, species richness and diversity of vegetation and arthropods (Venn and Kotze, 2014; Watson et al., 2020), thus leading these passively created urban grasslands to have a positive effect on biodiversity. An alternative approach to creating more environmentally friendly and biodiverse urban grasslands consists of substituting or modifying part of the lawnscape from the short, monocultural, homogeneous setting of grass species into meadows, a flowering setting with extensive management and possible active incorporation of forbs (Southon et al., 2017; Lane et al., 2019; Norton et al., 2019; Brezel et al., 2020; Marshall et al., 2023). This practice is becoming increasingly common as more cities and other communities look for ways to create sustainable and low-maintenance green space (Smith et al., 2015; Unterweger et al., 2017; Norton et al., 2019; Ignatieva et al., 2020). However, the relative impact of urban lawns and meadows, and the transformation...
between them, on soil and vegetation dynamics, carbon balance and greenhouse gas (GHG) exchange, remains poorly understood.

Within the context of climate mitigation and adaptation, the capacity of urban lawns to assimilate carbon varies widely due to climatic, edaphic and biological factors. Furthermore, management factors (e.g., mowing and irrigation) can make lawns behave either as carbon sinks or sources (Selhorst and Lal, 2013; Kong et al., 2014; Wang et al., 2022). For instance, soil organic carbon content in this type of urban green space tends to be larger than in natural grasslands (Kaye et al., 2005; Pouyat et al., 2006). Yet, carbon emissions from urban lawn soils have also been shown to be higher than those from natural and extensively managed urban grasslands (Kaye et al., 2005; Upadhyay et al., 2021). Unlike lawns and other types of grasslands, meadows do not only comprise grasses but also additional plant functional types such as legumes and other forbs, and the relative proportions of different plant functional types may influence the overall carbon dynamics and GHG exchange. However, the relative impact of urban lawns and meadows on soil and vegetation dynamics, carbon balance and greenhouse gas (GHG) exchange, remains poorly understood.

It is known that natural ecosystems with higher vegetation species richness generally show higher photosynthetic productivity than less diverse ones, due to a more efficient use of resources by a more diversified vegetation contributing to carbon assimilation (Fornara and Tilman, 2008; Lange et al., 2015; Yang et al., 2019; Chen et al., 2020). Additionally, they could lead to a more resistant functioning of the ecosystem in extreme weather events, as different grasses and forbs can respond differently to changes in environmental conditions (De Keersmaecker et al., 2016; Hossain et al., 2022). This means that, for example, during a drought event some plant species may be more resistant, making the functioning of a diverse green space less vulnerable than a less diverse setting. However, the relationship between species richness, productivity and resistance is not straightforward but influenced by many factors, such as management, vegetation community and environmental conditions (Vogel et al., 2012; De Keersmaecker et al., 2016; Jung et al., 2020).

The aim of this study is to determine the climatic impact of transforming urban lawns into urban meadows in northern Europe. In practice, we had four specific research questions:

1. Does the carbon balance differ between urban lawns and urban meadows?
2. Are urban meadows more tolerant to extreme weather events than urban lawns?
3. Does the transformation from an urban lawn to an urban meadow increase GHG emissions?
4. Does plant diversity functional types affect GHG fluxes in urban grasslands?

In addition, we were interested to see if we could detect any connections between different plant functional types and carbon and nitrogen (N) pools. To answer these questions, we measured GHG fluxes in urban lawns, in recently created urban meadows and in an older urban meadow in a more advanced phase around the Helsinki Metropolitan Area in Finland. Observations of the sites were used to set up a land-surface model, which was used to simulate annual carbon balances forced by reanalysis data. The empirical data set included intensive sites with high temporal coverage of CO₂ exchange to estimate the carbon balance and satellite sites with high spatial coverage to study the transformation process.

2 Materials and method

2.1 Site description

2.1.1 Study region

The measurements were collected within the Helsinki Metropolitan Area (60°10'N, 24°57'E, Fig. 1), which is situated on the northern coast of the Gulf of Finland. In Helsinki, the mean annual precipitation and temperature were 653 mm and 6.5
2023). According to the Köppen climate classification, the climate is humid continental (Dfb) (Kottek et al., 2006). The data collection took place on intensive and satellite sites which all are considered urban.

2.1.2 Intensive sites

Two in order to study the CO₂ exchange with high temporal coverage and to compare the carbon balances of urban grasslands, two pairs of urban lawns and urban meadows, situated 4 kilometers apart, were selected for the intensive measurement sites (Fig. 1). In Kumpula (KMP) neighborhood, an old and mesic and mesotrophic urban meadow, which could be associated to the “Mesic perennial anthropogenic herbaceous vegetation (V39)” in the EUNIS classification (European Environment Agency, 2023) and hereinafter referred to as “mesic meadow”, was paired with a highly managed urban lawn situated 150 meters away inside the Kumpula botanical garden, hereinafter referred to as “irrigated lawn”. The meadow is an old agricultural field on which farming practices were abandoned about 40 years ago. Nowadays, Currently, Aegopodium podagraria, Lupinus polyphyllus, (non-native invasive), Dactylis glomerata, Anthriscus sylvestris, Elymus repens, Lamium album and Urtica dioica are the dominant species. The meadow is cut once a year in autumn and most of the vegetation clippings are taken away. It is neither irrigated nor fertilized. The lawn, mainly covered by Poa pratensis, was installed about 15 years ago. It is currently mowed automatically by a mowing robot that operates daily between 6 PM and 11 AM, and the grass clippings are pounded and left on the site. It is irrigated during the summer when needed and was last fertilized in spring 2021. The mesic meadow and the irrigated lawn were situated 150 meters apart from each other.

In Viikki (VKI) neighborhood, the urban lawn and urban meadow – hereinafter referred to as “non-irrigated lawn” and “dry meadow” – are situated inside a public park 60 meters apart from each other. The urban lawn of Festuca sp. was sown in 2005 and is managed as a “utility lawn” (Viherympäristöliitto, 2023). It is mown regularly so that the grass height is 4–12 cm, and the clippings are mostly left on the site. There is no irrigation, and the lawn has not been fertilized for several years. In 2020, a section of the park’s lawns was transformed into a dry and nutrient-poor urban meadow by replacing the topsoil with a layer of recycled sand and sowing seeds (Riikonen and Karilas, 2021). This young dry urban meadow tends toward a “Dry perennial anthropogenic herbaceous vegetation (V38)” in the EUNIS classification (European Environment Agency, 2023). In 2022, the dominant species were Trifolium repens, Tripleurospermum inodorum, Lecanthes vulgare, Centaurea jacea, Phleum pratense and Plantago major, and some rare species such as Dianthus deltoides, Campanula rotundifolia and Galium verum were also observed. It was fertilized in spring 2021 and mown for the first time in autumn 2022. Most After mowing, most of the clippings were removed from the site.

2.1.3 Satellite sites

Six to study the transformation process from an urban lawn into an urban meadow, six other locations belonging to local student housing associations were selected for high spatial coverage GHG measurements in the urban green space. Six urban lawns over 20 years old, each located near the developing urban meadows and all subject to regular lawn management, were selected and studied as control cases. All of the six lawns were predominantly covered by grasses (mostly Poa pratensis, Festuca rubra and Lolium perene). In each of the locations, 50 m² of the already established lawn was transformed into an urban meadow site by volunteering and community participation.

Site transformations were conducted during late autumn 2020 by manually turning over the existent soil up to a depth of ca. 35 cm, followed by the sowing of 14 pollinator-friendly forbs at a density of 0.706 g m⁻². The commercial seed mixture (Suomen Nittyisimen Oy, Koski TI, Finland) also included four grass species at a density up to 0.67 g m⁻². Consequently, the hemiparasitic forb Rhinanthus minor, which is known to have negative effects on grass cover (Chaudron et al., 2021), was sown at a 0.65 g m⁻² density.
Every year since the lawn transformation, each of the developing urban meadows was managed by manually removing on-site tree seedlings (mainly Acer platanoides) before a one-time mowing event in autumn by hand scythe. Lastly, all mowing clippings were removed from the sites. Six urban lawns over 20 years old, each located nearby the developing urban meadows and all subject to regular lawn management, were selected and studied as control cases. All of the six lawns were predominantly covered by Poaceae sp.

Figure 1. Locations of the intensive (yellow) and satellite (green) measurement sites around the Helsinki metropolitan area. Background maps built with topographic database by the National Land Survey of Finland (2023) and global administrative borders by GADM (2023).

Table 1. Site characteristics at the intensive and satellite sites. Sown. The sown mixture indicates seeds of pollinator-friendly forbs, grass species and Rhinanthus minor which were sown in late 2020. veg. refers to vegetation. (± standard deviation) calculated for at least four replicates, at each VKI site there was only one replicate for each variable. Due to Finnish classification, approximations were made for sand (0.06–2.0 mm), silt (0.002–0.06 mm) and clay (<0.002 mm).

<table>
<thead>
<tr>
<th>Site ID</th>
<th>Experiment</th>
<th>Management</th>
<th>Soil texture</th>
<th>Type of vegetation</th>
<th>Vegetation’s characteristic</th>
<th>Irrigation</th>
<th>Sand (%)</th>
<th>Silt (%)</th>
<th>Clay (%)</th>
<th>pH</th>
<th>SOC (%)</th>
<th>SON (%)</th>
<th>CN ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>VKI lawn</td>
<td>Intensive</td>
<td>MP</td>
<td>Sandy loam</td>
<td>Poa grasses</td>
<td>69.5 ± 2.7</td>
<td>72.2 ± 2.6</td>
<td>22.1 ± 2.3</td>
<td>5.6 ± 0.3</td>
<td>23.8 ± 1.2</td>
<td>0.19 ± 0.09</td>
<td>14.5 ± 2.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VKI meadow</td>
<td>intensive</td>
<td>No</td>
<td>Sandy loam</td>
<td>Xerophilic mesic veg</td>
<td>59.1 ± 3.2</td>
<td>72.4 ± 2.6</td>
<td>31.8 ± 1.0</td>
<td>6.5 ± 0.4</td>
<td>2.5 ± 1.8</td>
<td>0.42 ± 0.16</td>
<td>14.0 ± 2.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>KMP lawn</td>
<td>intensive</td>
<td>Yes</td>
<td>Sandy loam</td>
<td>Poa grasses (Fescue)</td>
<td>72.6 ± 2.0</td>
<td>72.6 ± 2.0</td>
<td>42.7 ± 2.7</td>
<td>5.6 ± 0.3</td>
<td>32.8 ± 0.01</td>
<td>0.25 ± 0.14</td>
<td>14.6 ± 1.7</td>
<td></td>
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</tr>
<tr>
<td>KMP meadow</td>
<td>intensive</td>
<td>No</td>
<td>Sandy loam</td>
<td>Xerophilic mesic</td>
<td>72.4 ± 1.0</td>
<td>72.4 ± 1.0</td>
<td>42.7 ± 2.7</td>
<td>6.5 ± 0.3</td>
<td>2.5 ± 0.01</td>
<td>0.46 ± 0.14</td>
<td>14.2 ± 1.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JMT3C satellite</td>
<td>lawn</td>
<td>No</td>
<td>Silt loam</td>
<td>Poa grasses</td>
<td>5.2 ± 0.2</td>
<td>68.9 ± 2.6</td>
<td>26.0 ± 4.3</td>
<td>6.4 ± 0.4</td>
<td>5.0 ± 0.4</td>
<td>0.46 ± 0.04</td>
<td>10.8 ± 0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JMT3T satellite</td>
<td>Meadow</td>
<td>No</td>
<td>Loamy sand</td>
<td>Sown mixture</td>
<td>73.7 ± 2.3</td>
<td>73.7 ± 2.3</td>
<td>28.3 ± 4.3</td>
<td>6.5 ± 0.4</td>
<td>4.6 ± 0.7</td>
<td>0.18 ± 0.03</td>
<td>25.7 ± 0.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JMT7C satellite</td>
<td>lawn</td>
<td>No</td>
<td>Loamy sand</td>
<td>Poa grasses</td>
<td>82.7 ± 2.8</td>
<td>14.9 ± 2.4</td>
<td>2.6 ± 4.3</td>
<td>6.5 ± 0.4</td>
<td>4.0 ± 0.5</td>
<td>0.26 ± 0.04</td>
<td>15.0 ± 0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>JMT7T satellite</td>
<td>Meadow</td>
<td>No</td>
<td>Loamy sand</td>
<td>Sown mixture</td>
<td>81.4 ± 2.8</td>
<td>16.0 ± 2.6</td>
<td>2.6 ± 4.3</td>
<td>6.5 ± 0.4</td>
<td>3.2 ± 0.3</td>
<td>0.21 ± 0.02</td>
<td>15.3 ± 0.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>KO4C satellite</td>
<td>lawn</td>
<td>No</td>
<td>Sandy loam</td>
<td>Poa grasses</td>
<td>66.2 ± 2.6</td>
<td>29.4 ± 4.4</td>
<td>4.4 ± 4.3</td>
<td>6.4 ± 0.3</td>
<td>4.4 ± 1.2</td>
<td>0.29 ± 0.08</td>
<td>15.4 ± 0.6</td>
<td></td>
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</tr>
</tbody>
</table>
In this study, the lawn and the meadow within a pair were paired considering the short distance between each other and the two sites were mostly visited on the same day. GHG flux measurements were conducted at each site along a transect. All the data was collected at four fixed quadrats of approximately 1 m² distributed along a single transect. At the intensive sites, most of the quadrats were about 10 meters apart, but to fit the study setup to the landscape design, there were some variations mainly due to passing walking paths: at the non-irrigated lawn, the quadrats were separated by 10 m, 5 m and 10 m; at the dry meadow, the quadrats were separated by 17 m, 10 m and 10 m; and last, the quadrats on the irrigated lawn were located in two separate transects 15 meters away from each other, with the two quadrats on each transect being 3 meters apart. At the satellite sites, from the centroid of the site, a 10m long transect was established along the length of the site, with four quadrats of 1-m² area each uniformly distributed.

The soil temperature, soil moisture, soil samples and plant inventories were collected close to the GHG measurement quadrats except for the irrigated lawn where we used the soil samples collected from the same plot by Ahongshangbam et al. (2023).

2.2 Flux measurements

2.2.1 Measurement protocol

The GHG flux measurements were conducted with manual chambers using different setups at the intensive and satellite sites. At the intensive sites, the carbon dioxide (CO₂) exchange measurements were conducted more frequently than at the satellite sites, and they also included the light response (LR) of net ecosystem exchange (NEE). At the satellite sites, the measurements also included methane (CH₄) and nitrous oxide (N₂O) exchanges, but measurements were conducted with opaque chambers, thus disabling photosynthesis. The chamber measurements were always performed within the above-mentioned quadrats.

At the intensive sites, the light response of NEE was measured fortnightly between June and September 2021–2022. Additional measurements of NEE, but mainly of total ecosystem respiration (TER), were also conducted in May and in October–November. Over the two years of measurements, KMP lawn was visited 23 times, including 15 days of LR, KMP meadow 23 times, including 16 days of LR, VKI lawn and meadow 20 times, including 15 days of LR, and at each site, most of the time the four collars were measured at a site (Table S2). The measurement setup consisted of a transparent chamber attached to a CO₂ and H₂O analyzer (Li-840A, LI-COR, Inc., Nebraska, USA), an air temperature and humidity sensor (BME280, Bosch Sensortec GmbH, Reutlingen, Germany), and a photosynthetically active radiation (PAR) sensor (PQS1, Kipp & Zonen, Delft, Netherlands). The air was circulated through the analyzer at a flow rate of 1 L min⁻¹, and the chamber was equipped with a small fan to ensure air mixing within the chamber headspace. We used two differently sized chambers, either 0.072 m³ or 0.288 m³, and three collars of different heights (18.3 cm, 20.8 cm and 50 cm) to individually match the height of the chamber+collar combination to the vegetation we were measuring during each measurement day. Chamber measurements
were performed on four fixed plots established on the lawn and the meadow at each of the two locations. At VKI lawn and meadow as well as at KMP meadow, the plots were located about 10 meters apart, along a single diagonal transect. The plots on KMP lawn were located in two separate transects 15 meters away from each other, with the two plots on each transect being 3 meters apart. The transparent chamber was placed on the ground while avoiding damaging the plants, and a tight seal was created by using sand-filled cloth bags as insulation. In meadows, when a collar was used, it was placed on the ground and sealed with the cloth bags, and the chamber was placed on top of the collar. The duration of each chamber closure was at least 2 minutes. To determine the light response of NEE, we repeated the measurement five times at each plot/collar under different PAR intensities — 100%, 50%, 25%, 12.5% and 0% of the prevailing sunlight level — created by shading the chamber with netted or opaque fabrics. We first measured without covering the chamber and then covered it with 1–3 layers of mesh cloth.

Eventually, we measured the last recording with an opaque cover. The chamber was kept well ventilated — a pause of at least one minute in between the single light intensities to ventilate the chamber and to allow the plants some time to acclimatize to the changing light level. While conducting the measurements, we on the field, we allowed for ±10% variation in the PAR level during a single closure. TER was obtained from the measurement conducted with the opaque cover.

At the satellite sites, GHG measurements were conducted with an opaque chamber monthly, between May–September 2021–2022. Additional background measurements were conducted in July, August and October 2020, before the lawn transformations took place. In total, the sites were visited thirteen times: three times before transformation, including two times with CH₄ and N₂O measurements and one time with only CO₂, and ten times after transformation, including six times with CH₄ and N₂O measurements and four times with only CO₂ (Table S2). A single measurement was performed at each of the four selected 1-m² quadrats at each site (Table S2). To allow a good representativity of CO₂ measurements, but also measurements of N₂O and CH₄, two different types of devices were used over the different months (Table S2). The measurement setup in August and October 2020, as well as in May, July and September 2021 and 2022 consisted of a CO₂, H₂O, N₂O and CH₄ analyzer (DX4015, Gasmet Technologies Oy, Vantaa, Finland) connected to an opaque chamber equipped with a small fan. This Fourier Transform Infrared-based analyzer was zero calibrated at the beginning of each measurement day. The chamber volume was either 0.00456 m³ or 0.268 m³ depending on the height of the vegetation measured. An additional collar of 20–cm height was needed to allow enough room for tall vegetation in July 2021 and 2022. The measurements in July 2020 and in June and August in 2021 and 2022 were conducted with a setup consisting of an opaque chamber (volume 0.007434 m³) equipped with a CO₂ probe (GMP343, Vaisala Oyj, Vantaa, Finland), a relative humidity and air temperature sensor (HMP75, Vaisala), and a small fan. This chamber is introduced in detail by Ryhti et al. (2021). Four study plots were established both on the lawn and the meadow at each of the six locations. From the centroid of the site, a 10 m-long transect was established along the length of the site, with four quadrats of 1 m² area each uniformly distributed, and the chamber measurements were always performed within these quadrats. When measuring only CO₂ (i.e., July 2020, June and August 2021 and 2022), the duration of each chamber closure was 4–5 minutes. We extended the closure time to 15 minutes when measuring also N₂O and CH₄ (i.e., August and October 2020, May, July and September 2021 and 2022).

Soil temperature and soil moisture were always measured together with the chamber measurements at both the intensive and the satellite sites. One replicate of soil temperature was measured next to each flux measurement plot at a 10 cm depth with a handheld soil thermometer (HH376, Omega Engineering Inc., Connecticut, USA), and 1–5 replicates of soil moisture at 5 cm depth were measured with a handheld setup (ML3 Theta Probe and HH2 Moisture Meter, Delta-T Devices Ltd., Cambridge, UK). All plots were also photographed from above on each measurement day to calculate the green cover percentage of the basal area within the chamber using Canopeo (Patrignani and Ochsner, 2015).
2.2.2 Flux calculation

The measurement represents measurements representing the total ecosystem fluxes: total CH\textsubscript{4} fluxes (gCH\textsubscript{4} m\textsuperscript{2} s\textsuperscript{-1}), total N\textsubscript{2}O fluxes (gN\textsubscript{2}O m\textsuperscript{2} s\textsuperscript{-1}) and NEE (gCO\textsubscript{2} m\textsuperscript{2} s\textsuperscript{-1}). NEE includes possible photosynthetic input (GPP, gCO\textsubscript{2} m\textsuperscript{2} s\textsuperscript{-1}) and the outflow by respiration (TER, gCO\textsubscript{2} m\textsuperscript{2} s\textsuperscript{-1}), which includes both autotrophic and heterotrophic respiration. This can be mathematically written as:

\[
\text{NEE}(t) = \text{TER}(t) + \text{GPP}(\text{PAR}(t)).
\]

where PAR (µmol m\textsuperscript{-2} s\textsuperscript{-1}) is photosynthetically active radiation and \(t\) is time. Here we use meteorological notation i.e., we consider the inflow of carbon to be negative (GPP ≤ 0) and therefore NEE can be either negative (i.e., sink of CO\textsubscript{2}), when the absolute value of GPP is higher than TER, or positive (i.e., source), when TER is higher than the absolute value of GPP.

NEE or the total flux of other GHG fluxes during each chamber closure was calculated with the following Equation (2):

\[
\text{flux} = \left(\frac{dC(t)}{dt}\right)_{t=0} \frac{M_{\text{GHG}} P V}{R T A},
\]

where \(\left(\frac{dC(t)}{dt}\right)_{t=0}\) is the time derivative (ppm s\textsuperscript{-1} = 10\textsuperscript{-6} s\textsuperscript{-1}) of the linear regression, \(M_{\text{GHG}}\) is the molecular mass of the GHG (44.01 g mol\textsuperscript{-1} for CO\textsubscript{2}, 16.05 g mol\textsuperscript{-1} for CH\textsubscript{4}, and 44.02 g mol\textsuperscript{-1} for N\textsubscript{2}O), \(P\) is the air pressure (Pa), \(V\) is the system (chamber + possible collar) volume (m\textsuperscript{3}), \(R\) is the universal gas constant (8.31446 J mol\textsuperscript{-1} K\textsuperscript{-1}), \(T\) is the mean temperature inside the chamber headspace during the closure (K) and \(A\) is the soil surface i.e. the basal area of the chamber (m\textsuperscript{2}).

TER, CH\textsubscript{4} and N\textsubscript{2}O fluxes were obtained from the measurement conducted with the opaque cover (intensive sites) or with the opaque chamber (satellite sites).

To ensure a proper air mixing inside the chamber, at least ten seconds were removed from the beginning of each closure before fitting a linear regression into the GHG concentration values. Otherwise, the quality control and the length of the measurements included in the flux calculations differed between the instruments and GHGs. For the Vaisala equipment, CO\textsubscript{2} fluxes were calculated with the first minute of the recordings with R (version 4.2.2) after visual validation of linearity. Python (version 3.9.7) was used to calculate the fluxes measured by the two other devices. With the Gasmet analyzer, CO\textsubscript{2} fluxes were calculated from the first five minutes of the recordings whereas CH\textsubscript{4} and N\textsubscript{2}O fluxes, which were significantly lower fluxes and closer to the detection limits of the equipment, were calculated with the first seven minutes of the recordings. For the intensive sites, where the LI-COR analyzer was used, all fluxes were calculated from at least the first minute of the recording. There, possible poor-quality measurements were eliminated from the end of the measurement period if there was at least 1 minute of good quality data. The quality control measures were stability of light conditions inside the chamber during the closure and normalized root mean squared error (NRMSE) of the fit. Varying light conditions during the closure were inspected by calculating the standard deviation of PAR. If the standard deviation was larger than 150 µmol m\textsuperscript{-2} s\textsuperscript{-1} during the closure, the measurement was discarded. Yet, it is worth noting that in cases where the flux is nearly zero, NRMSE can be very large even though the measurement is not erroneous. Therefore, we did not discard measurements based on NRMSE if \(\left(\frac{dC(t)}{dt}\right)_{t=0}\) was below 0.1 ppm s\textsuperscript{-1}. Otherwise, measurements were discarded if NRMSE was larger than 0.05. Accordingly, 41 NEE measurements out of 1225 recordings were discarded from the intensive site dataset, but none of them referred to the opaque chamber measurements, i.e., TER. Thus, TER was represented by 23 and 20 days of data at KMP and VKI sites, respectively. Regarding the satellite sites, no fluxes were discarded. Regarding the satellite sites, the filtering process described above was also applied to the Gasmet dataset resulting in discarding 23 measurements of TER out of 384 recordings over the three years of measurements, but no CH\textsubscript{4}, nor N\textsubscript{2}O measurements (384 recordings of each in total). All TER measurements (240 TER recordings in total) measured with the Vaisala device were considered as valid (Table S2).
2.2.3 Light response (LR) fitting and daily GPP

In order to fit LR curves to the CO₂ flux data, we first subtracted TER from the other NEE rates to obtain an estimate for GPP (Eq. 1). GPP(PAR=0) was set to zero. The LR curve was expected to be a rectangular hyperbola (Ruimy et al., 1995), as follows:

\[ \text{GPP} = \frac{\text{PAR} \alpha \text{GP}_{\text{max}}}{\text{PAR} \alpha + \text{GP}_{\text{max}}} \]  

(3)

where PAR (µmol m⁻² s⁻¹) is photosynthetically active radiation, α (mg µmol⁻¹) is the initial slope between GPP and PAR, and GPmax (mg CO₂ mg CO₂⁻¹ m⁻² s⁻¹) is the light-saturated photosynthesis rate. The parameters α and GPmax were estimated using non-linear least squares minimization. α was allowed to vary between −0.1 and −1*10⁻⁸ mg µmol⁻¹ and GPmax between −5 and −1*10⁻⁷ mg CO₂ mg CO₂⁻¹ m⁻² s⁻¹.

In order to fit the LR curve for one collar, a minimum of three measurements done at different light intensities was required. After processing the light response calculation, LR curves where α had standard error over 0.0008 mg µmol⁻¹ or GPmax standard error was higher than 10 mg CO₂ mg CO₂⁻¹ m⁻² s⁻¹ were discarded from the dataset. In addition, LR was discarded if the highest PAR intensity in the response was under 500 µmol m⁻² s⁻¹. That resulted in 154 LR curves out of 212, representing 14 days of data at KMP meadow, VKI lawn and meadow and 10 days at KMP lawn, which were used in further analyses.

Daily GPP was calculated for each plot collar and for each measurement day using Eq. 3, fitted parameters for the plot collar and day, and continuous PAR measurements from SMEAR III station (Institute of Atmospheric Research, 2023). (Institute of Atmospheric Research, 2023). First, GPP was estimated for each half hour with the fitted parameters and the 30-min average PAR. Then, all half-hourly GPPs were summed up to obtain the daily GPP of the plot collar on the particular day.

2.3 Soil sampling analyses

2.3.1 Soil temperature and soil moisture

Soil temperature and soil moisture were always measured together with the chamber measurements within the 1-m² quadrats at both the intensive and the satellite sites. One replicate of soil temperature was measured within the quadrats at a 10-cm depth with a handheld soil thermometer (HH376, Omega Engineering Inc., Connecticut, USA), and 4–5 replicates of soil moisture at 5 cm depth were measured with a handheld setup (ML3 ThetaProbe and HH2 Moisture Meter, Delta-T Devices Ltd., Cambridge, UK).

2.3.2 Soil characterization

At the intensive sites, overall soil characteristics at the intensive sites were analyzed at a commercial lab (Eurofins Viljavuuspalvelu Oy, Mikkeli, Finland). We, soil density and C and N contents were analyzed.

To determine the overall soil characteristics, we collected altogether 1 L of soil up to 25 cm depth at each of the sites. At KMP irrigated lawn, the pooled sample comprised of 16–18 individual samples systematically collected in 2020 around the site with a thin auger (d = 2.3 cm). At the rest of the remaining intensive sites, we used the samples were collected in 2021 with a slightly larger auger (d = 5.0 cm) to collect 4 individual samples that were then pooled together. The fresh samples were stored in a fridge and sent to the lab within 1–3 days after collection. From these samples, particle size distribution was analyzed according to Elonen (1971). (1971) and the overall soil characteristics were analyzed at a commercial laboratory (Eurofins Viljavuuspalvelu Oy, Mikkeli, Finland).

Soil density samples were collected by digging a small pit and carefully inserting a steel cylinder (V = 0.151 dm³) horizontally into an undisturbed pit wall at 10 cm depth. When the cylinder was fully inserted, it was gently detached by removing the surrounding soil while ensuring that all soil within the cylinder remained in place in order to achieve volumetric accuracy.
Five replicates were collected at KMP lawn and three replicates at the other intensive sites. After collection, the samples were dried at 105 °C for 48 h and the dry weights were weighed.

Other soil samples were collected to determine C and N content. At KMP sites, samples for C and N were collected with soil auger (d = 2.3 cm) down to a 30 cm depth. Altogether, six individual samples were collected systematically at the lawn and the meadow. At VKI, we used soil auger (d = 1.7 cm) to collect eight samples and then pooled them together. The fresh irrigated lawn samples were first sieved (with a 2-mm mesh size = 2 mm) and the fresh weights of the smaller and larger soil fractions were weighed. The soil was then dried at 105 °C for 24 h. At the other intensive sites, the samples were first dried at 105 °C for 48 h, and then sieved with a 2-mm mesh sieve. Total soil C and N contents were determined from the dried and milled samples of soil with grain size smaller than 2 mm with an elemental CN analyzer (LECO, Michigan, USA). Total soil C and N contents were assumed to be only organic components without carbonates i.e., soil organic carbon (SOC) and soil organic nitrogen (SON).

At the satellite sites, overall soil characteristics, soil C and N contents and soil particle distribution were also determined. All soil samples from the satellite sites were collected during early July 2022, and all samples were extracted within a 1-m² radius from the plots in which GHG fluxes were measured. 1-m² quadrats. A total of four soil cores, dug by means of a soil auger of 5 cm diameter down to ca. 15 cm depth, were pooled together to create a composite soil sample. In each of the sites, a total of four composite soil samples were collected in ziplock bags and kept on ice during soil collection. All soil samples were then stored at −20 °C, until shipped and analyzed at a commercial lab (Eurofins Viljavuuspalvelu Oy, Mikkeli, Finland). Total soil C and N contents were assumed to be only organic components without carbonates, and SOC and SON were determined from freeze-dried and milled samples with an elemental CN analyzer (LECO, Michigan, USA). For the case of soil particle size distribution, a fraction of each of the soil samples was repeatedly washed with a 30% H₂O₂ solution to remove its organic matter content. The resulting soil samples were then dried out at 105 °C for 48 h, before sieving (mesh size = 0.6 mm) the final mineral soils. The sieved soil samples were then mixed with a sodium pyrophosphate solution (0.05 M), and their sand, silt and clay proportions were calculated by means of laser diffraction particle size analyzer (Bekman Coulter Inc., California, USA).

### 2.4 Biodiversity

Vegetation inventories were conducted in all intensive and satellite sites. Four quadrats of 1 m² each were distributed along on June 27th and July 21st 2022, respectively, according to the transects, in the same location as the chamber measurements-quadrat sampling method. In each of the quadrats, (see Section 2.1.4), the cover of each plant species was estimated and classified in one of the following plant functional type categories: forb, grass, horsetail, legume, moss, sedge or tree. Species richness (S) was estimated at site level, considering all grass species to be a combined, singular entity. Site species richness was used for the estimation of the site’s Shannon diversity index value (H′), as Equation (4):

\[ H' = -\sum_{i=1}^{S} p_i \ln(p_i) \]  

where \( p_i \) is the proportion of \( i \)th species, and \( S \) is the number of species in the site.

Lastly, species evenness (E) at the site proportion was calculated as, Equation (5) from the four quadrats to estimate the overall proportion of each plant functional type at a site.

\[ E = \frac{\mu^2}{\ln(S)} \]

All collars were also photographed from above on each measurement day to calculate the green cover percentage of the basal area within the chamber using Canopeo (Patrignani and Ochsner, 2015).
2.5 Ecosystem modelling

To derive annual estimates of NEE, TER and GPP, the intensive sites were simulated using the JSBACH (Reick et al., 2013), which is the land model in the Earth system models of the Max Planck Institute for Meteorology (Reick et al., 2013), the land component of the Max Planck Institute Earth system model (MPI-ESM) (Giorgetta et al., 2013). JSBACH is a process-based model and calculates the dynamic carbon cycle and key driving factors including seasonal dynamics in leaf area, momentary CO₂ fluxes, evapotranspiration, soil moisture, litter production and soil carbon dynamics.

The model was driven with hourly data (2005-2022) of air temperature, precipitation, shortwave and longwave radiation, relative humidity and wind speed. The driver data were derived from observations from the Kumpula weather station (60°12'1.1" N, 24°57'4.7" E–60°12'14.0" N, 24°57'38.9" E) operated by the Finnish Meteorological Institute (Finnish Meteorological Institute, 2023a). The data was gapfilled by observations from the nearby urban measurement station SMEAR III (Järvi et al., 2009). Hourly ERA5-Land data (Muñoz-Sabater et al., 2021) were used to fill a small number of remaining gaps. Leaf area index (LAI) data for the simulated sites were obtained from Sentinel-2 satellite data (Nevalainen et al., 2022; Nevalainen, 2022).

The vegetation in JSBACH is represented by plant functional types (PFT). Here, the model was set up for simulating only one PFT, C3 grass, for each site. The Logistic Growth Phenology (LoGro-P) model (Böttcher et al., 2016) is used to describe the phenology in JSBACH, where the temporal development of the leaf area index (LAI) of grass depends on both temperature and soil moisture. The maximum LAI for each site (Table S23) was set based on Sentinel-2 data (Nevalainen et al., 2022; Nevalainen, 2022), whereas the model simulated the seasonal LAI dynamics driven by temperature and precipitation. In addition, in the simulation of the mesic Kumpula meadow the shedding of the grass was activated 65 days after the growth started, in order to better follow-simulate the observed LAI.

According to the soil particle size distribution the soil texture at all sites was sandy loam. The parameters describing the soil properties follow the recommendations by Hagemann and Stacke (2015). However, the volumetric field capacity and wilting point for each site were adjusted based on the soil moisture measurements (Table S32). The root depths of the lawns were assumed to be shallower than for the meadows (Table S32).

The photosynthesis of C3-plants in JSBACH is described by the model by Farquhar et al. (1980). The photosynthesis is calculated once to get the unstressed canopy conductance, which is then scaled based on the soil moisture in the root zone to get the canopy conductance and photosynthesis under water stress. The available water in the root zone depends on the field capacity, root depth and a scaling factor \( f=\text{Wcrit}/(\text{Wcrit-Wwilt}) \). The scaling factor is applied for relative soil moisture values between \( \text{Wcrit}=\text{fcrit}^*\text{Wmax} \) and \( \text{Wwilt}=\text{fwilt}^*\text{Wmax} \), where \( \text{Wmax} \) is the maximum moisture content in the root zone. No water is available to the vegetation when the soil moisture reaches \( \text{fwilt}^*\text{Wmax} \). The factors \( \text{fcrit} \) and \( \text{fwilt} \) are given in Table S23. Moreover, the daily GPP values were used to adjust the photosynthetic parameters to meet the observations at each intensive site.

The adjusted model, forced with the observed weather, was used to derive the annual average GPP, TER and NEE of the intensive sites (KMP, VKI) for the period 2005–2021.

The model was used to derive the annual average GPP, TER and NEE for the period 2005–2022. The simulations were set up to represent habitats where the soil organic matter accumulates over time from the litter of standing vegetation. This was achieved by running a long spin-up period (thousands of years). As it is often the case in urban areas, the soils at the intensive sites had not been accumulated from the litter of the current vegetation, and therefore the soil carbon pools in the model are not equal to the ones present at the sites. Due to this, the simulations may not reproduce the observed TER, but instead represent a more general situation for these habitats.
were adjusted to meet the observed TER values in 2021 and 2022. The agreement between the simulated and observed carbon fluxes was evaluated by $R^2$ and RMSE.

2.6 Study of drought events

2.6.1 Drought definition

Since drought can be defined by the deficit of precipitation and the increase in evaporation (Wang et al., 2021), we estimated the drought period at our intensive sites with the Standardized Precipitation-Evapotranspiration Index (SPEI). We calculated the drought period at a 14-day scale with the SPEI R package (Vicente-Serrano et al., 2010; Beguería et al., 2014). The Penman-Monteith equation (Monteith, 1965) was used to calculate the Potential Evapotranspiration (PET) with the FAO-56 method of the FAO (Allen et al., 1998). The altitude was fixed at 10 m and the latitude at 60°11’ N.

The minimum temperature, maximum temperature, wind speed, air cloud amount, dew point temperature, relative humidity and air pressure, all collected at an hourly scale at the Kumpula meteorological station and averaged at a daily level, were downloaded from the Finnish Meteorological Database at the Kumpula meteorological station (Finnish Meteorological Institute, 2023). (Finnish Meteorological Institute, 2023b).

The drought periods were weekly defined with SPEI under −1.5, corresponding to the highest value describing a severe drought (Wang et al., 2021): a week with at least one daily SPEI value under −1.5 was considered as a drought week.

2.6.7 Statistical analyses

2.7.1 Statistical analyses regarding drought stress

With the collected data, the resilience of the different grassland vegetation types habitats to drought events was estimated. As resilience is the result of both components—resistance and recovery—we studied in this paper only the resistance, which was defined as “the magnitude of disturbance that a system can absorb before shifting from one state to another” (Capdevila et al., 2021).

Thus, to estimate the direct impact of drought on lawns and meadows, we chose to primarily compare the old mesic meadow (KMP meadow) and the non-irrigated lawn (VKI lawn), since both of these habitat vegetation types were older than 15 years and non-irrigated. In addition, we also included the irrigated lawn (KMP lawn) in the comparison to see the effect of irrigation.

The yearly summer means of TER, daily GPP, green cover, soil temperature and soil moisture were calculated for each site, by considering that the summer season starts on week number 24 and ends on week number 34 (data collected from May 31st–August 24th 2021 and June 3rd–August 25th 2022). Lastly, we calculated the differences between the measured values of the variable and the summer average of the same variable at the site level. These calculated differences were then normalized at a site and year level by subtracting the annual means of the differences and dividing the whole by the annual standard deviation of the differences.

With the summer data, we defined two categories: data collected during the drought events and data collected outside of the drought events. Finally, with the Mann-Whitney U tests, we compared, pair by pair, VKI lawn, KMP lawn and KMP meadow in the different summer conditions.

2.7 Statistical analyses

2.7.2 Dynamics of GHG fluxes at the satellite sites

2.7.1 Dynamics of GHG fluxes

All statistical analyses were run using R (version 4.2.2). Considering the transformation from lawns into meadows at the satellite sites, Shapiro-Wilk tests were used to evaluate the normal distribution of each variable (i.e., CO$_2$ fluxes, CH$_4$ fluxes, NO$_2$ fluxes, soil moisture and soil temperature) by year and type (meadow vs. lawn). Then, differences in given
variables between lawns and meadows were assessed either by t-tests or Mann-Whitney U tests depending on the normality of the data.

2.7.23 Statistical analyses of plant variable contribution

The effects of plant diversity and vegetation functional categories on SOC, SON and GHG fluxes were studied in all intensive and satellite sites with the plant and fluxes data collected in 2022. The fluxes were averaged for the growing season, from May to September. The Mann-Whitney U tests were used to compare the proportions of grasses, legumes and forbs between lawns and meadows.

The effects of vegetation were assessed by applying linear mixed-effects models (LMMs), using the R lme4 package (Bates et al., 2015), where the factor location describing the locations of the lawn/meadow pairs (Fig. 1) was included as a random effect. 18 mixed-effect linear regressions were built using one by one. This factor was introduced to consider the spatial variability between the sites, as the environmental conditions of the lawn and meadow within each of the biodiversity indices (i.e., species richness, Shannon index and species evenness) as site were considered to be similar. If predictive variables and each of the GHG fluxes as well as SOC and SON values as response variables. To test the contribution of each plant e., plant functional category on the GHG fluxes, as well as on SOC and SON values types correlated over 0.70, one of the correlated variables was discarded from the LMM according to the lowest cover proportion and the lowest occurrence in the dataset. Next, the normal distribution of each regression’s residuals was visually checked with Quantile-Quantile (Q-Q) plots. In total, 6 mixed-effect multiple regressions including the proportions of all the selected plant functional categories (i.e., legumes, grasses, forbs, trees (saplings), sedges, horsetails and mosses) types as predictive variables were run by applying a descending backward selection. At each iteration, predictive variables with p-values higher than 0.1 were removed and variables with p-values lower than 0.05 were considered statistically significant. Before testing any regression, the distributions of each of the LMMs were visually checked with Quantile-Quantile (Q-Q) plots. P-values were adjusted by a Kenward-Roger approach (Kenward and Roger, 1997) from R lmerTest package (Kuznetsova et al., 2017) and a conditional R²-squared was used to assess the quality of the model.

3 Results

3.1 Drought occurrences

There was 7% and 12% more precipitation in 2021 and 2022 compared to the 653-mm value of the reference period 1991–2020 (Jokinen et al., 2021), but in both years July was drier than the long term average, with 17% and 15% less precipitation in 2021 and 2022 respectively (47 mm in 2021 and 49 mm in 2022 compared to 57 mm in 1991–2020). At the same time, 2021 and 2022 were 0.1 and 0.4 °C warmer than the reference which was 6.5 °C. The summers were on average hotter than the reference period with 2.5 °C and 1.8 °C higher temperatures in 2021 and 2022, respectively, compared to 16.6 °C. These particularly hot and dry weeks were observed from June 14th to August 1st 2021 and from June 6th to July 17th 2022 (Fig. 2a).

At the intensive sites, the measured soil moisture (Fig. 2b) also showed a drastic decrease at all sites during these two periods, except at the irrigated lawn (KMP lawn), where the soil moisture remained relatively high during the whole summer. The highest soil temperature was also reached during the same period (Fig. 2c). In 2021 and 2022, those hot and dry weeks also corresponded to the decrease phase of LAI, except at the dry meadow (VKI meadow), where LAI increased slowly from spring until August (Fig. 2d). The measured green cover was notably following the same patterns as the satellite LAI data (Fig. 2e).

Using the SPEI, two drought periods were defined: the first one from June 14th to August 8th in 2021, and the second one from June 27th to July 17th in 2022 (Fig. 2). Thus, the defined drought event in 2021 happened early in the summer season and
lasted eight weeks, whereas the drought event in 2022 arrived later in the summer and lasted only three weeks. These two drought events co-occurred with the high temperature and low soil moisture periods described above.
Figure 2. (a) Weekly precipitation (Prec.) and mean air temperature (Tav) recorded by FMI meteorology station. (a), (b) manually measured soil moisture at a 5 cm depth, (c) manually measured soil temperature at a 10 cm depth, (d) leaf area index from satellite Sentinel-2, and (e) manually measured green cover at the intensive study sites with standard deviations during the campaign years 2021–2022 calculated for each site at a daily scale. The irrigated lawn is KMP lawn, the non-irrigated lawn is VKI lawn, the mesic meadow is KMP meadow, and the dry meadow is VKI meadow. Red rectangles indicate the drought periods according to the SPEI and light grey rectangles represent the summer season. In panel (a), Tmax and Tmin represent the weekly instantaneous maximum and minimum mean temperatures.

3.2 CO₂ fluxes

3.2.1 Observed seasonal variation

At all intensive sites, the measured TER increased at the beginning of the measurement campaign and declined during the drought events at the non-irrigated sites. When comparing the differences between the four sites, the mean momentary TER was the highest at the mesic meadow (KMP meadow: 0.44 and 0.31 mgCO₂ mg CO₂ m⁻² s⁻¹ in 2021 and 2022, respectively) and the lowest at the dry meadow (0.29 and 0.26 mgCO₂ mg CO₂ m⁻² s⁻¹ in 2021 and 2022,
The highest recorded mean values \(0.86, 0.84, 0.59, 0.59 \text{mg CO}_2 \text{mg CO}_2 \text{m}^{-2} \text{s}^{-1}\) were measured at VKI\text{non-irrigated} lawn, KMP\text{mesic} meadow, KMP\text{dry} meadow, KMP\text{irrigated} lawn, on June 7th, July 2nd, July 19th, August 11th, respectively. In 2021, the highest values \(0.71, 0.49, 0.65, 0.55 \text{mg CO}_2 \text{mg CO}_2 \text{m}^{-2} \text{s}^{-1}\) were recorded at VKI\text{meadow}, KMP\text{lawn}, KMP\text{dry meadow}, VKI\text{irrigated lawn, mesic meadow, non-irrigated} lawn on June 23rd, June 30th, July 1st, August 17th, respectively. In general, daily TER were higher at KMP sites (irrigated lawn and mesic meadow) compared with VKI sites (non-irrigated lawn and dry meadow), although the highest values were not reached in a specific order.

There were site-specific variations in the seasonal pattern of GPP (Figure S1). In the first campaign year, after the first measurements, the daily GPP sink decreased at all sites except VKI\text{dry} meadow, for which it was the first growing season after establishment. In 2022, the daily GPP sinks first increased and then started to decrease around the end of June (Figure 3). The highest recorded daily uptakes \(-0.48, -0.51, -0.40, -0.46 \text{mg CO}_2 \text{mg CO}_2 \text{m}^{-2} \text{s}^{-1}\) were measured at VKI\text{non-irrigated} lawn, KMP\text{mesic} meadow, VKI\text{dry} meadow, KMP\text{irrigated} lawn, on June 7th, June 18th, August 5th, August 11th, respectively, in 2021. In 2022, the order had also slightly changed as the \text{daily highest uptake} \(-0.37, -0.58, -0.42 and -0.47 \text{mg CO}_2 \text{mg CO}_2 \text{m}^{-2} \text{s}^{-1}\) were recorded at VKI\text{non-irrigated} lawn, KMP\text{mesic} meadow, KMP\text{irrigated} lawn, VKI\text{dry} meadow on June 10th, June 13th, June 13th and June 23rd. Thus, the measured GPP values were higher at KMP sites (irrigated lawn and mesic meadow) than at VKI sites, (non-irrigated lawn and dry meadow).

### 3.2.2 JSBACH performance

In order to represent the average fluxes of the different habitats, the model was primarily initialized to reach a steady state with each vegetation type and therefore, the simulated soil carbon pool represents the result of long-term carbon input of the site itself. The simulated soil moisture (Fig. S2) varied according to irrigation, precipitation and evaporation and showed lower values in summer, particularly in June and July, and also in August in 2022. The modelled seasonal dynamics in soil moisture followed the observations in general; however, some of the observations were considered to represent moisture content above the volumetric field capacity. The maximum soil moisture in the model is limited by the field capacity (Fig. S2). At the same time, the modelled soil temperature between 10 and 15 cm depth agreed with the measured soil temperature at 10 cm depth (Fig. S3). Modelled LAI varied in accordance with the observations especially at KMP\text{mesic} meadow and was the second year at VKI\text{dry} meadow (Fig. S4). For VKI\text{dry} meadow, was established very recently and was found to be sparse and inhomogeneous with respect to the vegetation, and therefore the drought response was not captured challenging to capture in the model simulation particularly in early summer 2021 during the first growing season of the meadow. The drought response for VKI\text{non-irrigated} lawn agreed with the observations in 2021, while in 2022 the recovery was too strong in the simulation. At KMP\text{irrigated} lawn, the simulated LAI decreased later in 2022 than the observed LAI (Fig. S4).

Simulated TER fits the observations at KMP lawn in 2021–2022 and at KMP meadow in 2022 (Fig. 3, Fig. S5). The measured instantaneous TER was sometimes higher than modelled TER in summer, while it overlapped with the modelled TER in spring and in autumn (Fig. 3 and S5). Simulated daily mean GPP followed the observations closely (Fig. 3), except at VKI meadow where the model overestimated the observations in early season after the sowing (Fig. S5h), which is probably due to the discrepancy in LAI between the simulation and observations.

Simulated TER followed the dynamics in the observed seasonal cycle and, for example, showed a similar response to the 2022 drought but generally underestimated the overall level of the non-irrigated lawn and the dry meadow. The drought response was stronger in the mesic and dry meadow (Fig. 3e-h, Fig. S5e-h). For the irrigated lawn the benefits of irrigation may have been overestimated, seen as a weaker drought response in the model than in the observations. The irrigation used in the model was estimated as an average over the whole area, while there probably was less irrigation where the measurement equipment was installed. The highest \(R^2\) values for TER were observed at the irrigated lawn and mesic meadow and the lowest at the dry
meadow (Table 2). Adjusting the carbon pools to fit closer to observed TER values in 2021 and 2022 (Fig. S6-7) mainly decreased RMSE but did not notably improve R² values at the different vegetation types in 2021 and 2022 (Table 2). The model was able to simulate the overall level and the seasonal cycle in GPP at the irrigated lawn, non-irrigated lawn, and mesic meadow (Fig. 3a-c, Fig. S5a-c, Table S5). At the dry meadow, the model overestimated the increase in GPP in the early season during its first growing season in 2021 (Fig. S5d). The vegetation cover in the newly established meadow was very low in the early growth season in 2021, which was not accounted for in the model. The agreement was better in the following year (Fig. 3d, Table S5). The benefit of irrigation was also somewhat overestimated for GPP during the dry periods. Simulated daily mean GPP followed the observations closely for non-irrigated lawn and mesic meadow (R² between 0.41 and 0.85, Table 2), while the dry meadow was difficult to simulate especially in 2021 (Table 2, Fig. S5).

3.2.3 Annual balances

During 2005–2021, simulated annual GPP and TER were on average the highest at KMP the irrigated lawn and mesic meadow (Table 23). GPP and TER were the lowest at VK in the dry meadow (Table 23) being approximately 40% of those of KMP. The unirrigated lawn’s GPP and TER were approximately 70% of those of the irrigated lawn. The lawns resulted had more negative NEE i.e., a greater sink of carbon, than meadows, and looking at the ratio between standard deviation and NEE, meadows showed higher year-to-year variations than the lawns (Table 23). The sink of the irrigated lawn (KMP lawn) was approximately 53% higher than that of the unirrigated lawn (VKI lawn). Additionally, it was found that 13–16% of TER occurred between November and March.

According to the simulations that were adjusted to meet the TER observations, annual NEE were 328, 2663, 1476 and 2140 g CO₂ m⁻² yr⁻¹ in 2021 and –195, 1496, 571 and 1205 g CO₂ m⁻² yr⁻¹ in 2022, at the irrigated lawn, the non-irrigated lawn, the mesic meadow and the dry meadow respectively (Table S4).
Figure 3. Seasonal dynamics of mean total ecosystem respiration (TER, abcd) and daily photosynthesis (GPP, efgh) in the four intensive sites in 2022. Grey continuous lines represent JSBACH simulations and diamonds and dots represent the mean of manual measurements with standard deviation bars. (a, e) Irrigated lawn = KMP lawn; (b, f) non-irrigated lawn = VKI lawn; (c, g) mesic meadow = KMP meadow; (d, h) dry meadow = VKI meadow. Red rectangles indicate the drought periods according to the SPEI and light grey rectangles represent the summer season. The model simulated the whole year but for clarity, Jan-March and Dec are not visible as there were no measurements during those months.

<table>
<thead>
<tr>
<th>Table 2.</th>
<th>R² and root mean square error (RMSE, mg CO₂ m⁻² s⁻¹) calculated for total ecosystem respiration (TER) and gross primary production (GPP) in the standard simulation where the soil carbon was stabilized based on the standing vegetation, and for the adjusted simulations (adj soil c) where the soil carbon pool was set so that simulated TER met the observations in 2021 and 2022.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Year</td>
<td>Site</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>2021</td>
<td>irrigated lawn</td>
</tr>
<tr>
<td></td>
<td>non-irrigated lawn</td>
</tr>
<tr>
<td></td>
<td>mesic meadow</td>
</tr>
<tr>
<td></td>
<td>dry meadow</td>
</tr>
<tr>
<td>2022</td>
<td>irrigated lawn</td>
</tr>
<tr>
<td></td>
<td>non-irrigated lawn</td>
</tr>
<tr>
<td></td>
<td>mesic meadow</td>
</tr>
<tr>
<td></td>
<td>dry meadow</td>
</tr>
</tbody>
</table>

Table 3. Mean annual (± standard deviations calculated for 2005–2022) of ecosystem respiration (TER), photosynthetic uptake (GPP) and net ecosystem exchange (NEE) modeled with JSBACH at the four intensive sites during the years 2005–2022.

<table>
<thead>
<tr>
<th>Sites</th>
<th>Characteristics</th>
<th>TER (g CO₂ m⁻² yr⁻¹)</th>
<th>GPP (g CO₂ m⁻² yr⁻¹)</th>
<th>NEE (g CO₂ m⁻² yr⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KMP lawn</td>
<td>Irrigated lawn</td>
<td>4331 (± 177)</td>
<td>-4667 (± 218)</td>
<td>-336 (± 187)</td>
</tr>
<tr>
<td>VKI lawn</td>
<td>Non-irrigated lawn</td>
<td>3039 (± 251)</td>
<td>-3196 (± 231)</td>
<td>-157 (± 139)</td>
</tr>
<tr>
<td>KMP meadow</td>
<td>Mesic meadow</td>
<td>4127 (± 291)</td>
<td>-4278 (± 233)</td>
<td>-151 (± 198)</td>
</tr>
<tr>
<td>VKI meadow</td>
<td>Dry meadow</td>
<td>1765 (± 136)</td>
<td>-1828 (± 161)</td>
<td>-63 (± 134)</td>
</tr>
</tbody>
</table>
3.3 Extreme weather resistance

Green cover, TER and daily GPP were significantly decreased at the unirrigated lawn (VKI lawn) compared to the records outside the drought and to the mature mesic meadow (KMP meadow) and the irrigated lawn (KMP lawn) during the prolonged drought in 2021 (Fig. 4ace) and during the shorter drought in 2022 (Fig. 4bdf). In 2022, TER was significantly higher at the irrigated lawn and at the mesic meadow during the drought than outside of the drought period (Fig. 4d).
Figure 4. The resistance indices of an irrigated lawn (KMP lawn), a non-irrigated lawn (VKI lawn) and an old mesic urban meadow (KMP meadow) to drought events in 2021 (first column) and 2022 (second column), calculated for the green cover (a, b), the TER (c, d) and the daily GPP converted to positive values (e, f). The values are the differences between the measured values and the summer average, normalized at a yearly level. Horizontal lines represent the normalized summer means of each site. Summer includes Jun–Aug and droughts were defined in Fig. 2 (June 14th–August 8th, 2021 and June 27th–July 17th, 2022). (*) for p-value ≤ 0.10; * for p-value ≤ 0.05; ** for p-value ≤ 0.01; and *** for p-value ≤ 0.001.
Figure 5. Box and whisker plots of measured (a) total ecosystem respiration (TER), (b) CH$_4$ and (c) N$_2$O fluxes of lawns and transformed meadows at the satellite sites before (grey background) and after transformation (white background), which happened in the end of 2020 only at the transformed meadows. The environmental conditions were considered to be similar between the two treatments. Statistical differences between both types of green spaces were tested at the year level (¤ for p-value ≤ 0.10; and * for p-value ≤ 0.05).

The measurements at the control lawn and transformed meadow of each site were conducted consecutively during the same day, and therefore the environmental conditions were considered to be as similar as possible. Indeed, soil moisture and temperature did not differ between the treatments during the measurements (Fig. S9). At the satellite sites, before the transformation in 2020, there were no differences between lawn and meadow sites in terms of TER, CH$_4$ and N$_2$O fluxes measured in darkened conditions (Fig. 5). There was some general year-to-year variation in the median and range of observed CO$_2$ and CH$_4$ fluxes, but the fluxes did not differ between lawns and transformed meadows in 2021 or 2022 (p > 0.1, Fig. 5ab). The mean N$_2$O fluxes were slightly lower in meadows than in lawns in 2021 and 2022 (p-value < 0.1, Fig. 5c). Soil moisture and temperature did not differ between the treatments during the measurements (Fig. S7).

3.5 Biotic Plant functional type predictor variables for C and N cycles

According to the linear regressions, TER, CH$_4$ fluxes, N$_2$O fluxes, SOC, SON and C/N ratio were not related with species richness, Shannon index, nor Species evenness according to the data collected in 2022 (Table 3).

With the six multiple regressions, we found that TER was. On average, the total vegetated area of the studied meadows (Table 4) did not significantly differ from those on lawns (U = 40.000, p-value = 0.442), but the proportions of the main plant functional types shared by both vegetation types differed. Meadows had a significantly lower proportion of grasses (i.e., 25%) than lawns (65%, U = 10.000, p-value = 0.002). Most of the meadow area was covered by forbs other than legumes (i.e., 65%).
a proportion significantly larger ($U = 16,000$, $p$-value $= 0.002$) than that found in lawns (i.e., 19%). Nevertheless, the proportion of legumes was found to be statistically similar ($U = 39,000$, $p$-value $= 0.505$) in meadows (11%) and lawns (6%).

Next, we explored possible correlations between plant functional types and variables related to C and N cycles. First, we discarded sedges from the analysis since it was correlated with horsetails ($> 0.7$, Table S7) and only present at one site in a low proportion (Table 3). None of the functional plant types are significantly associated to TER, nor SON. CH$_4$ flux appeared to be positively related with almost all vegetation categories, except mosses and just with minor significance positively with sedges and trees, and moss proportion (Table 5), and N$_2$O flux was significantly negatively with horsetails (Table 4). CH$_4$ appeared to be significantly and negatively correlated with sedges associated to forb proportion and positively with horsetails (Table 4), and N$_2$O was mainly negatively associated with forbs and positively with horsetails (Table 4) to horsetail proportion (Table 5). SOC was slightly positively explained by forbs proportion whereas SON appeared to be significantly positively connected with grass proportions and almost with forbs (Table 4). The C/N ratio was significantly and negatively connected associated with legume, grass and forb proportions (Table 45).

Table 3. Effect of plant diversity (i.e., species richness, Shannon index and species evenness) on Table 4. Cover proportions (%) of the different plant functional types: grasses (Poaceae), legumes (Fabaceae), forbs (other families of flowering vascular plants, which do not belong to one of the listed categories), trees, sedges (Carex), horsetails (Equisetum), and mosses (Bryophyta), inventoried on June 27th and 21st July 2022. The “total” column is the sum of all the plant functional type cover, a fully covered quadrat with only one layer of vegetation should have a 100% cover, >100% indicates layered vegetation (short and tall grassland plants) and <100% indicates the presence of bare soil.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Location</th>
<th>Management</th>
<th>Grasses</th>
<th>Legumes</th>
<th>Forbs</th>
<th>Trees</th>
<th>Sedges</th>
<th>Horsetails</th>
<th>Mosses</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intensive</td>
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<td>lawn</td>
<td>70.5</td>
<td>6.0</td>
<td>21.5</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>98.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>meadow</td>
<td>30.0</td>
<td>16.5</td>
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<tr>
<td></td>
<td>VKI</td>
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<td>92.3</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>100.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>meadow</td>
<td>6.3</td>
<td>38.3</td>
<td>23.3</td>
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<td>0.0</td>
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<td>0.0</td>
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<tr>
<td>Satellite</td>
<td>JMT-3</td>
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<td>0.0</td>
<td>0.0</td>
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<tr>
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<tr>
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<td>37.5</td>
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<td>0.0</td>
<td>79.4</td>
</tr>
<tr>
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<td></td>
<td>meadow</td>
<td>31.9</td>
<td>13.0</td>
<td>68.3</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
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</tr>
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<td>KO-4</td>
<td>lawn</td>
<td>78.8</td>
<td>3.9</td>
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<td>0.0</td>
<td>0.3</td>
<td>0.0</td>
<td>98.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td>meadow</td>
<td>10.0</td>
<td>0.4</td>
<td>53.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
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</tr>
<tr>
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<td>0.0</td>
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<td>0.0</td>
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<tr>
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<td>SK-1</td>
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<td>65.0</td>
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<td>24.3</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>99.0</td>
</tr>
<tr>
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<td></td>
<td>meadow</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>136.0</td>
</tr>
<tr>
<td></td>
<td>SMT-12</td>
<td>lawn</td>
<td>66.3</td>
<td>2.3</td>
<td>27.8</td>
<td>1.3</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>97.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>meadow</td>
<td>15.8</td>
<td>1.8</td>
<td>106.0</td>
<td>0.3</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>123.8</td>
</tr>
</tbody>
</table>

Table 5. The connection between the proportions of different plant functional types and total ecosystem respiration of CO$_2$ (TER), CH$_4$ fluxes, N$_2$O fluxes, Soil C content (SOC), Soil N content (SON) and C/N ratio. Estimate ($p$-value for $p$-value $< 0.05$; * for $p$-value $< 0.01$; ** for $p$-value $< 0.001$) indicates values that were only measured at satellite sites.
| C/N ratio | 0.23 (0.14) | 0.06 | 2.51 (0.14) | 0.14 | 8.46 (0.12) | 0.18 |

Table 4. The connection between the proportions of different plant functional categories and total ecosystem respiration of CO$_2$ (TER), CH$_4$ fluxes, N$_2$O fluxes, Soil C content (SOC), Soil N content (SON) and C/N ratio. The plant functional categories are grasses (Poaceae), legumes (Fabaceae), forbs (other families of flowering vascular plants, which do not belong to one of the listed categories), trees, sedges (Carex), horsetails (Equisetum), and mosses (Bryophyta). (NS for p-value > 0.10; * for p-value ≤ 0.10; * for p-value ≤ 0.05; ** for p-value ≤ 0.01; and *** for p-value ≤ 0.001). sat. indicates values that were only measured at satellite sites.

<table>
<thead>
<tr>
<th>Predictive variables</th>
<th>Legumes</th>
<th>Grasses</th>
<th>Forbs</th>
<th>Trees</th>
<th>Sedges</th>
<th>Horsetails</th>
<th>Mosses</th>
<th>R$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TER (mg CO$_2$ m$^{-2}$ s$^{-1}$)</td>
<td>0.003 *</td>
<td>0.002 **</td>
<td>0.001 *</td>
<td>0.063 *</td>
<td>17.658 H</td>
<td>-0.297 H</td>
<td>NS</td>
<td>0.02</td>
</tr>
<tr>
<td>CH$_4$ fluxes (ng CH$_4$ m$^{-2}$ s$^{-1}$) sat.</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>-2.2039.32 *</td>
<td>366.476 *</td>
<td>2.429 H</td>
<td>0.22</td>
</tr>
<tr>
<td>N$_2$O fluxes (ng N$_2$O m$^{-2}$ s$^{-1}$) sat.</td>
<td>0.588 H</td>
<td>NS</td>
<td>-0.279 **</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>0.98</td>
<td></td>
</tr>
<tr>
<td>SOC (% ≤ 2 mm)</td>
<td>NS</td>
<td>NS</td>
<td>0.011 H</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>0.56</td>
</tr>
<tr>
<td>SON (% ≤ 2 mm)</td>
<td>NS</td>
<td>0.002 *</td>
<td>0.002 H</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>0.28</td>
</tr>
<tr>
<td>C/N ratio</td>
<td>-0.281 **</td>
<td>-0.159 ***</td>
<td>-0.079 **</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>0.69</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Predictive variables</th>
<th>Grasses</th>
<th>Legumes</th>
<th>Forbs</th>
<th>Trees</th>
<th>Horsetails</th>
<th>Mosses</th>
<th>R$^2$</th>
</tr>
</thead>
</table>
| TER (mg CO$_2$ m$^{-2}$ s$^{-1}$) | NS | NS | NS | NS | NS | NS | /
| CH$_4$ fluxes (ng CH$_4$ m$^{-2}$ s$^{-1}$) sat. | NS | NS | NS | NS | NS | 2.698 H | 0.63 |
| N$_2$O fluxes (ng N$_2$O m$^{-2}$ s$^{-1}$) sat. | NS | NS | -0.279 ** | NS | 1.092 ** | NS | 0.98 |
| SOC (% ≤ 2 mm) | NS | NS | 0.011 H | NS | NS | NS | NS | 0.56 |
| SON (% ≤ 2 mm) | NS | NS | NS | NS | NS | NS | /
| C/N ratio | -0.159 *** | -0.281 ** | -0.079 * | NS | NS | NS | /

4 Discussion

It is well known that converting lawns into meadows can increase biodiversity in cities (Venn and Kotze, 2014; Wastian et al., 2016; Chollet et al., 2018). In this study, we wanted to understand better how transforming lawns into meadows in Nordic urban areas could affect their GHG fluxes and how their vegetation diversityplant functional types could influence the C and N cycles. The studied sites, an irrigated lawn, a non-irrigated lawn, an old mesic spontaneous meadow and a young dry meadow, were mainly C sinks. Moreover, with a focus on the transformation dynamic from lawns into meadows, no significant differences in measured GHG fluxes (CO$_2$, CH$_4$ and N$_2$O) were found between the six control lawnsites and six transformed meadowsites in the two following years after the conversion. However, the studied mesic meadow appeared to be more resistant to drought stress than an unirrigated lawn. This resistance could notably be explained by plant diversity, as discussed later. Furthermore, as expected, the forbs proportion was found to be higher in meadows than in lawns and seems to be negatively associated with N$_2$O fluxes and C/N ratio, and positively to SOC, even though some further studies would be necessary to understand better the interaction between the plants and the C and N cycles, we found that the mixture of plants functional categories showed some links with the C and N cycle.
4.1 Fluxes in urban grasslands

Carbon neutrality, or even a net positive scenario, is nowadays one of the major goals for cities and states in mitigating climate change (IPCC, 2022). Our aim was to understand how CO₂ sequestration differs between lawns and meadows by studying contrasting vegetation types: habitats, a mesic mesotrophic very fertile meadow and a dry nutrient very poor meadow together with irrigated and non-irrigated lawns, to determine the full range of their GHG exchange. We found that the photosynthetic production (GPP) of an irrigated lawn and a mesic meadow were roughly equaled on an annual scale whereas the GPP of a non-irrigated lawn was notably lower. At an annual level, the irrigation increased the GPP of a lawn by over 40% and the sink (NEE) by more than 100%. It has already been found that water input improves carbon uptakes (Thienelt and Anderson, 2021) and demonstrated by Zirkle et al. (2011) that irrigated lawns store up to 10 g C m⁻² yr⁻¹ more carbon in soil than non-irrigated ones. Also, whereas our analysis on annual NEE study indicated that an irrigated lawn is a stronger sink as much as approximately 50 g C m⁻² yr⁻¹ higher than a non-sink on an irrigated lawn.

Here, the model estimates for NEE had some uncertainties as the model was unable to simulate some of the observed momentary total ecosystem respiration (TER) values during summertime (Fig. 3) which caused the full carbon balance estimate to be more uncertain than the estimated GPP. In this experiment, we were measuring the collars plots at various times of day due to practical reasons, even though Pavelka et al. (2018) suggest measuring ecosystem respiration in the morning around 9–10 to get the average level of daily respiration. Moreover, with manual chamber measurements, we were able to measure only once or twice a month and most of the time under a sunny weather, whereas with automatic chambers or eddy covariance (Hiller et al., 2011; Thienelt and Anderson, 2021), it would have been possible to estimate the daily and monthly variations at a finer scale and to reduce the bias caused by hot and dry days. Therefore, especially during the summertime, the measured values most probably are overestimations of the daily average whereas in the early spring and in autumn, when the diurnal cycle in soil temperature and plant activity are smaller, the observations fit the model estimate closely. Decina et al. (2016) found that during the growing season, TER was about 0.198 ± 0.006 mg CO₂ m⁻² s⁻¹ at urban lawns in Boston, USA, which is lower than the values measured at our sites. Last, it has been shown that during cold days, under snow and/or frozen ground, grasslands are a source of CO₂ (Hiller et al., 2011; Jasek-Kamińska et al., 2020). The model was also run for winter months with the heterotrophic soil respiration decreasing at the same time as the soil temperature until reaching a minimum in late winter. Although the highest ecosystem activities and emissions take place in the warm summer months, to fully validate the annual balance, it would be useful to have some additional wintertime observations as well. On the other hand, nevertheless, we can be quite confident in the simulated photosynthetic production, since GPP principally occurs during the snow-free seasons.

Furthermore, soil respiration is highly related to soil carbon quality and quantity (Davidson and Janssens, 2006). Here, it seems evident that the studied soil was not stabilized but, rather, that growing media with unknown properties was either brought to the site or some earlier vegetation type built the soil carbon storage. This is most evident in the young sites (less than 15 years old) and most managed sites i.e., the lawns and the dry meadow. It highlights unpredictable features of urban soils which are characterized by high anthropogenic disturbances, unknown origins and changes in land use. Thus, the quality and quantity of organic matter cannot be connected to a linear history even at a city scale (Pouyat et al., 2006; Setälä et al., 2016; Ivashchenko et al., 2019; Sushko et al., 2019; Cambou et al., 2021). Here, we chose to estimate the carbon balance over an extended time in stabilized conditions and all our sites were estimated to be small sinks of carbon varying between 63–336 g CO₂ m⁻² yr⁻¹. However, adjusting the model parameters to reproduce values closer to the observed TER values turned most of the sites to sources of atmospheric CO₂ during the measurement years 2021 and 2022. Even if those values represent the current situation at the study sites, those represent also an unstabilized state. Whereas the extended runs stand for long-term carbon balances of these vegetation types whereas the results of earlier studies are conflicting.
Many have reported urban grasslands to be C sources at least in certain conditions (Allaire et al., 2008; Hiller et al., 2011; Bezyk et al., 2018; Hundermark et al., 2024), but others have also reported sinks of about 20–180 g CO₂ m⁻² yr⁻¹ (Thienelt and Anderson, 2021). However, most of the studies focus on the respiration rate, omitting GPP (Kaye et al., 2005; Decina et al., 2016; Lerman and Contosta, 2019; Sushko et al., 2019; Upadhyay et al., 2021). Jasek et al. (2020) found that the annual mean of TER for a mix of different types of urban grassland is 424 ± 43 g C m⁻² yr⁻¹ in Krakow, Poland, and Kaye et al. (2005) measured that irrigated lawns in Northern Colorado, USA, emits around 2777 ± 273 g C m⁻² yr⁻¹, where our annual TER values vary between 1181 ± 48 g C m⁻² yr⁻¹ at the irrigated lawn and 481 ± 37 g C m⁻² yr⁻¹ at the dry meadow. In the end, we found that lawns were stronger sinks than the meadows which is in line with Peoplau et al. (2016) who found that soil C was higher in lawns than in meadows, which was mainly attributed to the clippings left at the site and fertilization.

It is noteworthy that lawns in Finland have a certain requirement are well maintained to support soil fertility (Viherympäristöliitto, 2023); whereas meadows grow on various types of soil ranging from poor and dry to mesic and fertile. We found that the mesic meadow, with high (max. 1.3 m) and dense vegetation in summer, had higher photosynthetic production and annual sink than the dry meadow sown in late 2020. However, during the campaign that took place 0–2 years after the establishment, the dry meadow was still in an initial phase i.e., its carbon sequestration potential may not have been fully realized, with about six-times lower CO₂ uptake than the mesic meadow. In addition, in the same meadow, there were dissimilar features among the four sampled plots: one of them was scarce in plants, another one had a tall and dense vegetation, and the two others had a short dense vegetation, which have made it difficult to compare if the ecosystem model fitting simulations and the analysis even more challenging. For these reasons, it would be useful to compare in the future the fertile site to naturally dry and homogenous meadows with similar varying fertility – from mesic mesotrophic to dry nutrient-poor, but comparable age.

With the satellite sites Nevertheless here, we particularly focused on the transformation process with the satellite sites and found no negative climate impacts in terms of greenhouse gases during the campaign which covered two growing seasons after the transformation. However, it is possible that we missed some peaks of emissions as the measuring frequency was just once a month. Nonetheless, it is quite safe to conclude that even if such unmonitored peaks occurred, their significance on an annual level would be minor as there was no indication of any differences during the measurements. On the other hand, it is evident that destroying vegetation during the transformation process decreases photosynthetic input at least during the following autumn and spring when lawns continue photosynthetic activities but when the new meadow has not developed yet.

Therefore, it would have been interesting to measure GPP as well and conduct the experiment over a longer period of time, as it has been shown that even grassland restoration requires of many more years (Müller et al., 1998; Waldén and Lindborg, 2016; Kose et al., 2021). In addition, the CO₂, H₂O, N₂O and CH₄ analyzer used at the satellite sites has a low sensitivity for small fluxes, at least in certain conditions (Kohl et al., 2019), in comparison with more advanced analyzers. However, we were only interested in the possible difference between the two treatments and not the actual rates nor annual balances and therefore, we find the choice to be acceptable. In any case, it would be important to study the CH₄ exchange. N cycle and particularly N₂O fluxes in urban grasslands in more detail. In addition, the applied transformation process was the same at all the satellite sites and it would be interesting to also study the impact of different transformation processes on GHG fluxes.

Finally, even though the spectra of our four intensive sites was wide, the four sites were unique in their environmental conditions, vegetation and management. In order to provide a stronger categorization, it would have been beneficial to study more replicates intensively. Moreover, mowing frequency was not studied, nor its carbon footprint, although studies have found notable impacts on source and sink due to managements (Allaire et al., 2008; Poepplau et al., 2016; Lerman and Contosta, 2019; Thienelt and Anderson, 2021) leaving room for improvements for further studies.
4.2 CO₂ fluxes impacted by drought events

Extreme climate events may occur more often in the coming years and during the two years of measurements in this study, dry summers impacted the ecosystems. Here, we focus only on the resistance component, i.e. the capacity of the system to absorb the drought disturbance during the event (Vogel et al., 2012; Capdevila et al., 2021). According to the comparison between drought and the outside-of-drought periods, the unirrigated lawn reached a more desiccated stage during the dry period of the summer, whereas the meadows seemed to endure the deficit of water and the hot weather as well as the irrigated lawn. In unirrigated lawns, such a suppression is a physiological reaction to drought stress reducing photosynthesis (GPP) and respiration (TER), as also found in other studies (Allaire et al., 2008; Hiller et al., 2011; Vogel et al., 2012). Yet, it has been also demonstrated in several studies that species-rich grasslands better endure extreme weather events (Vogel et al., 2012; De Keersmaecker et al., 2016).

The greater resistance of meadows compared to non-irrigated lawns could be explained by several factors, such as plant complementarity, microclimate and management (Vogel et al., 2012; De Keersmaecker et al., 2016; Bernath-Plaisted et al., 2023). (1) Lawns are usually sown, weeded and managed in order to maintain the desired low species composition and aesthetic appeal; on the other hand, plant diversity is usually much richer in meadows. Yet, the sampling effect theory (Tilman et al., 1997; Loreau and Hector, 2001) argues that in a more diverse grassland, there is more chance to find at least one species, which is well adapted to drier habitats. For this reason, there is a higher chance for at least one plant species to survive during drought events in meadows than in lawns. (2) The second theory is the niche complementarity, which results from niche differentiations and the benefits of interspecific interactions among them between species (Tilman et al., 1997), and induces a better individual species performance (Tilman et al., 1997; Loreau and Hector, 2001). Such a high diversity of plants and niches, as we can find in meadows, leads to a better nutrient, light and water-use efficiency (De Boeck et al., 2006; Walde et al., 2021) leads to a better water use efficiency as described by De Boeck et al. (2006), and helps to face water shortage. In relation to (1) and (2), meadows usually contain vegetation with deeper root systems, which allow plant individuals to supply themselves with water and reduce competition for the available water in the upper soil layers during drought periods. (3) Moreover, this complementarity in meadows is reinforced by microclimate anomalies (Bernath-Plaisted et al., 2023). The tall and dense vegetation of meadows creates a buffer to face climate extreme events by offering shade to the lower layers and keeping the humidity beneath the meadow canopy. (4) Finally, management, and notably mowing frequency, is also described as one of the factors impacting drought resistance. Regrowth of vegetation is more sensitive to extreme weather stress than vegetation at a later stage of its growth dynamics (Vogel et al., 2012). Hence, lawns are more affected, since they are typically mown every two to three weeks during the growing season, whereas meadows are cut one to two times a year. Thus, vegetated urban meadows are able to function almost as well as irrigated lawns and as well as during regular summertime weather since, e.g., during and outside drought periods, the green cover and the photosynthetic uptake are comparable.

In this study, we did not have a reference year without a drought, as according to daily SPEI calculation, both years of measurements were affected by extreme drought stress. Further, it would be important to understand how these ecosystems function during a normal climate year, especially to compare with a year similar as 2021 when the drought period covered 8 weeks out of 13 weeks of summer, even though the pre-drought reference and post-drought stable stages used in this study would better estimate the resistance itself (Capdevila et al., 2021), even though the pre-drought reference and post-drought stable stages used in this study would better estimate the resistance itself (Capdevila et al., 2021), especially in 2021 when the drought period covered 8 weeks out of 13 weeks of summer.

4.3 Biodiversity effects Effect of plant functional types on C and N stock cycles

It is known that biodiversity, including plant diversityits component associated with vegetation, increases drastically after conversion from lawn to meadow (Venn and Kotze, 2014; Chollet et al., 2018; Norton et al., 2019). Yet, we wanted to
understand how plant diversity, the composition of vegetation, could affect the SOC and the GHG fluxes at our sites. However, contrary to what was revealed in other studies (Fornara and Tilman, 2008; Lange et al., 2015), we did not find any link between the plant diversity and the C and N cycles, i.e. TER, CH$_4$, N$_2$O fluxes and C and N soil contents. Indeed, it has been found that vegetation richer in plant species, a large variety of plants, could explain a higher rate of TER (Dias et al., 2010) and could further enhance accelerate more N and C storage in the soil (Fornara and Tilman, 2008; Oelmann et al., 2011; Mueller et al., 2013; Cong et al., 2014; Lange et al., 2015). Nevertheless, Wei et al. (2017) demonstrated that under scenarios in which vegetation was more diverse in its plant functional groups, with high plant functional group diversity, instead of species diversity, the nitrate pool and the N mineralization rate were lowered. Mueller et al. (2013) also found that high diversity enhances N transformation and NH$_4^+$ pool, but a too high plant diversity could also have the opposite effect. One of the issues in our study could be that we studied the influence of diversity with only a one-year dataset in order to include the satellite as well as the intensive sites. The second issue may be that we used the SOC and SON instead of their accumulation over the years, because a three-year period is not enough to appreciate a C or N accumulation in the soil, where the changes happen slowly, i.e. a changing rate lower than 2% per year (Fornara and Tilman, 2008), whereas the standing pools of C and N are reflecting also historical land use.

To go further into the diversity, thus, in our study, we explored tested the possible connections the link that the different plant functional type categories, i.e. grasses, legumes, forbs, trees, horsetails and mosses, could have on the C and N cycles. As expected, according to our analysis, TER increased with most of the plant categories and CH$_4$ flux was connected to hygrophytes, such as horsetails and mosses, even though the relation between soil moisture and CH$_4$ fluxes in urban grassland seem to be hardly predictable (Groffman and Pouyat, 2009; Costa and Groffman, 2013; Bezyk et al., 2023). Although, even though some of the fluxes were accounting to the analysis linked to trees, sedges, horsetails and mosses, it would be beneficial to validate the results with a broader dataset since some plant functional types were scarcely represented: mosses and sedges were present in only one site, horsetails in two sites and tree saplings were surveyed in only four sites, whereas there were four of the sixteen sites included in the study (Table 4). Nevertheless, we showed that SOC increases with forbs, of which the proportion is higher in meadows. Hence, the meadows could store more carbon than lawns because also: (1) because of the transformation, some SOC could have been incorporated into the soil due to the site transformation; (2) meadows could have produced more litter that enriched the soil in carbon; and (3) with a deeper and strong root system, forbs could have allocated more carbon into the soil. Indeed, it has been shown that sites with woody plants, which also have deeper root systems, have a higher stock of carbon than lawns (Setälä et al., 2016; Lindén et al., 2020). However, such an increase in soil carbon was not revealed by the plant diversity analysis or supported by the flux analysis. One of the shortfalls in our study regarding C and N cycles and plant functional types could be that it was studied with only a one-year dataset in order to include both the satellite and the intensive sites. We utilized SOC and SON pools in our analyses, even though their accumulation rate would have been a more interesting metric, as the standing pools also reflect the historical land use of the sites. However, a three-year period would not have been long enough to observe changes in the pools due to their slow changing rate of less than 2% per year (Fornara and Tilman, 2008). Furthermore, the SON model was not significant according to the R-squared, but SON was slightly related to grasses and forbs, and not to legumes; however, in the transformed meadows, the proportion of legumes was rather low.

Some other studies have also tried to estimate the correlation between plant functional type and soil characteristics, by studying C3 and C4 grasses and legumes (Fornara and Tilman, 2008; Yang et al., 2019). Fornara and Tilman (2008) estimated that C4 grasses may increase soil carbon accumulation by 193%, and legumes by 522%. Moreover, even though legumes did not appear to be one of the significant groups in our SOC model, and negatively correlated to C/N ratio, legumes are known to improve the soil N availability and the litter quality by reducing the C/N ratio (Fornara and Tilman, 2008; Yang et al., 2019). Thus, plant functional types undoubtedly impact the C and N cycles; however, the low representativity of some plant functional types and the low number of replicates limit any further interpretation of the robustness of our analyses, but
To draw a stronger conclusion on the impact of the functional types on carbon and nitrogen dynamics, it would be necessary to study a larger number of sites and also compare chosen mixtures of plants on soil with the same soil properties.

5 Conclusions

In order to slow down the biodiversity loss, cities around the world are considering converting their lawns into meadows. Another priority is to mitigate climate change and achieve carbon neutrality. In this study, we focused on urban grasslands from a climate and biodiversity vegetation perspective by studying maintained lawns in comparison with mature and newly transformed meadows in northern Europe at the border between the boreal and temperate zones. The transformations from lawns to meadows did not demonstrate any additional negative climate effects in terms of ecosystem respiration, CH$_4$ and N$_2$O fluxes. Meadows do increase biodiversity, but, however, while meadows are known to have higher species richness and diversity of plant functional types, we found that meadows do not increase carbon sequestration compared with lawns on an annual scale, and even that NEE is usually more negative in lawns than in meadows. Nevertheless, in a climate warming perspective, with more frequent extreme events, meadows resist drought events better than lawns and are better able, at least in the short term, to adapt cities to extreme drought events induced by climate change. Nonetheless, it would be necessary to check this result with a larger number of sites and investigate the overall resilience with longer time series dataset, to help stakeholders and city planners to make the right decisions to optimize land-use. Thus, considering biodiversity, climate warming and carbon neutrality, it is necessary to find the right trade-off between lawns and meadows in cities. Regarding the plant diversity, the species richness does not seem to affect significantly the C and N cycles, even though some vegetation categories seem, forbs proportion was found to be higher in meadows and some plant functional type proportions, including grasses, legumes, forbs, horsetails and mosses, are correlated to specific C and N status. Nevertheless, as this analysis was done with only one-year dataset, it would be necessary to investigate with more sites, urban grasslands, on the link between plants and C and N cycle. Moreover, it would be beneficial to further study the carbon storage process and its longevity, as well as the biodiversity dynamics during transformation from lawns to meadows or drought resilience processes, and to consider the social aspects of the conversion of lawns to meadows.

Data availability

The data used in the manuscript can be downloaded from Finnish Meteorological Institute’s data storage: http://hdl.handle.net/11304/9a597a75-6f3b-48ee-b3e6-d23598817a5b, https://doi.org/10.23728/fmi-b2share.6cfb9e97b72648d6ad57fc3a08cfc6734 (Trémeau et al., 2023).

Code availability

The scripts used to calculate LI-COR and Gasmet fluxes can be found at: https://github.com/hvekuri/Chamber-codes/ (Vekuri, 2023).
Author contributions

LK, BO, EK and JT designed the research; BO, EK and JT participated in the data collection; HV, LK, EK and JT participated in the flux calculations; JT and BO conducted the statistical data analyses; LB performed the JSBACH simulations; all of the authors participated in writing the paper.

Competing interests

The authors declare that they have no conflict of interest.

Acknowledgements

This research takes part in the CO-CARBON project supported by the Strategic Research Council working under Academy of Finland (grant no. 335204), in the CarboCity project supported by the Academy of Finland (grant no. 325549), in the ACCC Flagship program of the Academy of Finland (grant no. 337552), in the Lawns into Meadows project supported by Koneen säätö (grant no. 201902880), and was also supported by the Maj and Tor Nessling foundation (grant no. 202000391).

We warmly thank Olivia Kuuri-Riutta, Pinja Rauhamäki and Elisa Vainio for their help with field data, Suvi Orttenvuori for her help with the preliminary JSBACH simulations, Elina Nieminen for supporting the research in its early phase, Olli Nevalainen for the Sentinel-2 data, Joséphine Couet, Helena Rautakoski and Thomas Merrien for support in the statistical analyses, Mika Korkkakoski, Stéphanie Gerin and Helena Rautakoski for their help with the chamber measurement analysis, Jack Chapman for the proofreading and valuable comments and finally, we also thank the reviewers, for their supports. We also thank the Stara company, represented by Inkeri Salo, the City of Helsinki, represented by Tuuli Ylikotila, and the Finnish Museum of Natural History, represented by Mikael Lindholm, for providing crucial information on the landscape management. Finally, we express gratitude to the City of Helsinki, the Finnish Museum of Natural History, AYY and HOAS for allowing us to transform and/or conduct our measurements on their properties.

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