

1 Cost Effective Off-Grid Automatic Precipitation Samplers for 2 Pollutant and Biogeochemical Atmospheric Deposition

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19 20 21 22 **Abstract**

23 An important transport process for particles and gases from the atmosphere to aquatic and
24 terrestrial environments is through dry and wet deposition. An open-source, modular, off-grid, and
25 affordable instrument that can automatically collect wet deposition samples allows for more
26 extensive deployment of deposition samplers in fieldwork and would enable more comprehensive
27 monitoring of remote locations. Precipitation events selectively sampled using a conductivity
28 sensor powered by a battery-based supply are central to off-grid capabilities. The prevalence of
29 conductive precipitation - that which initially contains high solute levels and progresses through
30 trace level concentrations to ultrapure water in full atmospheric washout, depends on the sampling
31 location but is ubiquitous. This property is exploited here to trigger an electric motor via limit
32 switches to open and close a lid resting over a funnel opening. The motors are operated via a
33 custom-built and modular digital logic control board, which have low energy demands. All
34 components, their design and rationale, and assembly are provided for community use. The
35 modularity of the control board allows operation of up to six independent wet deposition units,

36 such that replicate measurements (e.g., canopy throughfall) or different collection materials for
37 various targeted pollutants can be implemented as necessary.

38 We demonstrate that these platforms are capable of continuous operation off-grid for integrated
39 monthly and bimonthly collections performed across the Newfoundland and Labrador Boreal
40 Ecosystem Latitudinal Transect (47° to 53° N) during the growing seasons of 2015 and 2016.
41 System performance was assessed through measured power consumption from 115 volts of
42 alternating current (VAC; grid power) or 12 volts of direct current from battery supplies during
43 operation under both standby (40 or 230 mA, respectively) and in-use (78 or 300 mA, respectively)
44 conditions. In the field, one set of triplicate samplers was deployed in the open to collect incident
45 precipitation (open fall) while another set was deployed under the experimental forest canopy
46 (throughfall). The proof-of-concept systems were validated with basic measurements of rainwater
47 chemistry including: i) pH ranging from 4.14 to 5.71 in incident open fall rainwater; ii)
48 conductivity ranging from 21 to 166 $\mu\text{S}/\text{cm}$; and iii) dissolved organic carbon concentrations in
49 open fall and canopy throughfall of 16 ± 10 mg/L and 22 ± 12 mg/L, respectively; with incident
50 fluxes spanning 600 to 4200 mg C $\text{m}^{-2} \text{a}^{-1}$ across the transect. Ultimately, this demonstrates that
51 the customized precipitation sampling design of this new platform enables more universal
52 accessibility of deposition samples to the atmospheric observation community – for example, those
53 who have made community calls for targeting biogeochemical budgets and/or contaminants of
54 emerging concern in sensitive and remote regions.

55

56 **1.0 Introduction**

57 Atmospheric deposition is the central loss process for particles and gases to terrestrial and
58 aquatic surfaces (Pacyna, 2008). Particles and gases can be deposited by both dry and wet
59 deposition processes. Dry deposition is facilitated by the direct interaction of gases and particles
60 with boundary layer surfaces such as water, vegetation, and/or soil, while wet deposition involves
61 in-cloud scavenging and below-cloud interception of gases and aerosols by, e.g., rain droplets and
62 snow crystals (Fowler, 1980; Lovett and Kinsman, 1990). Dry and wet deposition are global
63 processes coupled to regional synoptic scale conditions, but their relative importance depends on
64 local sources and global transport of atmospheric analytes of interest. Dry deposition consists of a
65 variety of mechanisms for particles and gases, with fine mode particles and their chemical
66 constituents (compared to ultrafine and coarse mode particles) being more likely to undergo

67 atmospheric long-range transport prior to being deposited (Farmer et al., 2021). Wet deposition
68 occurs when such long-lived atmospheric particles and gases are included and/or scavenged into
69 cloud water and transported to the surface of the Earth in precipitation (e.g., snow and rain). With
70 the size and number of droplets in the atmosphere largely controlling the rate, wet deposition
71 depends on a variety of meteorological factors affecting precipitation, such as the size distribution
72 and concentration of ice and droplet nucleating particles, as well as the solubility, concentration,
73 and reactivity of gases (Lovett, 1994). Ultimately, deposition plays an important role in pollutant
74 distribution and biogeochemical cycling of long-studied major nutrients (e.g., nitrogen and sulfur
75 in acid rain) and those with increasing recognition of importance such as dissolved organic carbon
76 (DOC) (Meteorological Service of Canada, 2005; Vet et al., 2014; Safieddine and Heald, 2017;
77 United States Environmental Protection Agency, 2020).

78 Recognizing the significance of atmospheric trace chemical deposition has led to the
79 establishment of monitoring networks. For example, long-term wet deposition monitoring
80 networks, like the Canadian Air and Precipitation Monitoring Network (CAPMoN) and the
81 National Atmospheric Deposition Program (NADP), aim to provide critical data on the spatial and
82 temporal patterns of wet and dry deposition. As a result, this has allowed for the estimation of
83 regional and continental deposition rates of species regulated by national or international policies
84 (Lovett, 1994). Data from these networks have been critical to understanding the efficacy of policy
85 to reduce environmental issues like acid rain (Likens and Butler, 2020). In particular, the Oslo and
86 Geneva protocols have achieved an 80% decrease in both North American and European SO₂
87 emissions since 1980 (Grennfelt et al., 2020). Despite these successes, reduction in acid deposition
88 has had unexpectedly slow recovery in ecosystems leaving them sensitized – necessitating
89 continued deposition monitoring (Stoddard et al., 1999; Kuylenstierna et al., 2001).

90 Over the past 60 years, the precipitation chemistry community has made advancements in
91 deposition collectors to better understand atmospheric processes (Siksna, 1959). While bulk
92 deposition collection (i.e., a collection bucket or jug fitted with a funnel open at all times; Hall,
93 1985) is both a simple and economically feasible sampling method utilized by monitoring
94 networks, it is subject to bias through collection of inputs other than atmospheric deposition (e.g.,
95 bird droppings, insects, plant debris). As a result, bulk collectors can overestimate total deposition
96 and underestimate wet deposition in a variety of locations (Lindberg et al., 1986; Richter and
97 Lindberg, 1988; Stedman et al., 1990). Sequential precipitation collection methods include

98 manually segmenting samplers (requiring only a shelter, clean surface, and an operator), linked
99 collection vessels (sample containers that are filled in sequence via gravitational flow), amongst
100 others and have been developed to analyze rainwater composition and measure parameters such
101 as pH and conductivity (Gatz et al., 1971; Reddy et al., 1985; Vermette and Drake, 1987; Laquer,
102 1990). Sequential sampler designs have also been adapted to collect precipitation in remote field
103 sites (Germer et al., 2007; Sanei et al., 2010). Although it is a more costly and time intensive
104 method when compared to bulk deposition collection, the major appeal of measuring isolated wet
105 deposition is the ability to isolate this individual atmospheric process. Further innovation can
106 reduce bias and improve the preservation of samples, such as the use of sensors to automate
107 isolation of collected precipitation or the addition of polymeric mesh barriers to reduce debris input
108 in windy environments (Lovett, 1994) - yet commercial solutions often come at a substantial
109 expense.

110 When targeting biogeochemically relevant species in deposition collectors, additional
111 standard practices have been developed to improve the representativeness of sample composition.
112 First, an appropriate monitoring site must be selected. Three categories of siting criteria,
113 established by organizations such as CAPMoN and the NADP, are of particular importance: (i)
114 site representativeness and physical characteristics, (ii) distance from potential pollution sources,
115 and (iii) operational requirements (Canadian Air and Precipitation Monitoring Network, 1985a;
116 National Atmospheric Deposition Program, 2009). This means that each site must be a location
117 that receives precipitation representative of the hydrologic area; is ideally not within 500 m of
118 local pollution sources, such as wood-burning stoves, garbage dumps, and vehicle parking lots;
119 and is accessible for daily collections, maintenance, and can be serviced by reliable 115 volts of
120 alternating current (VAC) electrical power (Canadian Air and Precipitation Monitoring Network,
121 1985a; National Atmospheric Deposition Program, 2009). Despite these guidelines, there are many
122 reasonable scenarios in which these siting conditions cannot be met. As an example, remote sample
123 collections are often required for global assessments on persistent contaminants or nutrients of
124 biogeochemical importance. Remote locations, however, can result in sampling sites with no
125 power provision, infrequent sample collection, and/or the infrastructure-bearing location itself
126 being a source of the targeted pollutants. As a result, innovation in collection strategies such as
127 time-integrated off-grid sampling, with modularity in the deployment of replicates, as well as

128 materials for quantitative collection of environmental targets, is still needed to expand and/or
129 modify networks to meet current and future monitoring and policy needs.

130 In biogeochemical cycles, for example, improvement of constraints in atmospheric carbon
131 linkages to terrestrial and aquatic processes is necessary. This would play a critical role in correctly
132 assessing climate feedbacks and reducing uncertainty in Earth system models. The measurement
133 of atmospheric DOC transport to surfaces has been limited and impedes landscape scale carbon
134 balance from being obtained (Casas-Ruiz et al., 2023). The pool of compounds from which DOC
135 is derived in the atmosphere has also been limited and is only recently seeing an increase in
136 research intensity. Reactive organic carbon (ROC) is defined as the sum of nonmethane organic
137 gases and primary and secondary organic aerosols (Safieddine and Heald, 2017). The major
138 removal mechanism of water-soluble organic compounds produced through oxidation from the
139 atmosphere is by dry deposition of particle-bound pollutants and scavenging by rainfall (Jurado et
140 al., 2004, 2005). When ROC is scavenged into rainfall, it becomes DOC and enters terrestrial and
141 aquatic systems. Deposition measurements of ROC compounds are needed since they play a
142 crucial role in the formation of secondary species such as ozone, particulate matter, and carbon
143 dioxide (CO₂) (Safieddine and Heald, 2017; Heald and Kroll, 2020).

144 There are several evolving drivers around studying atmospheric ROC; for example, light-
145 absorbing organic carbon that can affect global radiative balance and undergo photochemical
146 transformations in the condensed phase (Saleh, 2020; Wang et al., 2021; Washenfelder et al., 2022;
147 George, 2023). Reactive organic carbon can also influence cloud formation and contribute to
148 precipitation acidity (Avery et al., 2006; Ramanathan and Carmichael, 2008). Measurements of
149 speciated ROC are difficult due to the chemical complexity of emitted compounds and oxidation
150 products (Heald and Kroll, 2020). To circumvent this, monitoring and quantifying DOC can be
151 used as a proxy to estimate the total ROC in precipitation. However, quantitative measurements of
152 DOC in precipitation samples are sparse due to its relatively low concentration of 0.1 to 10 mg C
153 L⁻¹ (Iavorivska et al., 2016; Safieddine and Heald, 2017). Recently, calls for carbon closure on
154 atmospheric processing of ROC make this measurement of increasing importance (Kroll et al.,
155 2011; Heald et al., 2020; Barber and Kroll, 2021). Similarly, to obtain net landscape or watershed
156 carbon exchange, studies require effective methods for capturing and preserving atmospheric DOC
157 deposition to constrain biogeochemical linkages at global interfaces as outlined above.

158 In this work, we present the design of a custom-built automated array of precipitation
159 samplers that can be operated both on- and off-grid for wet deposition collection. The purpose of
160 these samplers is to enable cost-effective collection of integrated water-soluble conductive
161 atmospheric constituents deposited in remote environments without grid power or routine access.
162 A sensor interfaces with a custom-built motor control board capable of operating up to six
163 independent wet deposition units such that canopy throughfall (TF) and incident precipitation
164 (open fall, OF) measurements are possible to collect in replicate. The materials used can be easily
165 changed in order to optimize collection and preservation of a wide array of target analytes, such
166 as DOC, when using high density polyethylene and mercuric chloride (HgCl₂). We demonstrate
167 that these platforms are capable of continuous operation off-grid for monthly wet deposition
168 collection of precipitation across the Newfoundland and Labrador Boreal Ecosystem Latitudinal
169 Transect (NL-BELT) during snow-free periods in 2015 and 2016. Extremes in system performance
170 were evaluated by testing the power consumption of a sampling array from spring through fall
171 when paired with a solar top-up system, and during snow-free winter conditions using only a
172 battery. The two years of field samples were collected using an array of six collection units, with
173 triplicate collection of both incident precipitation and throughfall from rain passing through a
174 forest canopy. Samples were analyzed in terms of deposition volumes relative to total bulk
175 volumes, reproducibility of replicate samples, and to determine the fraction of conductive rainfall
176 within the total volume of precipitation at these remote sites. The captured fraction compared to
177 total volume deposited is used to gain insight into how these samplers can limit analyte dilution
178 effects and improve method detection limits, such as rejecting 50% of the total volume delivered
179 as ultrapure precipitation leading to a factor of two improvement. Chemical parameters of pH,
180 conductivity, and DOC fluxes collected according to established preservation protocols were then
181 compared to prior measurements to validate this proof-of-concept system. Measurement methods
182 for pH and conductivity of rainwater are very well-established in the literature and serve as a
183 baseline reference to ensure that the samples collected by the new devices presented in this work
184 are consistent with what is expected in samples from a remote coastal environment, given the
185 selective sampling strategy. We then move away from these well-established parameters to
186 quantify DOC fluxes using established biogeochemical preservation techniques for fresh water
187 and groundwater to demonstrate the potential of these samplers in application to automated

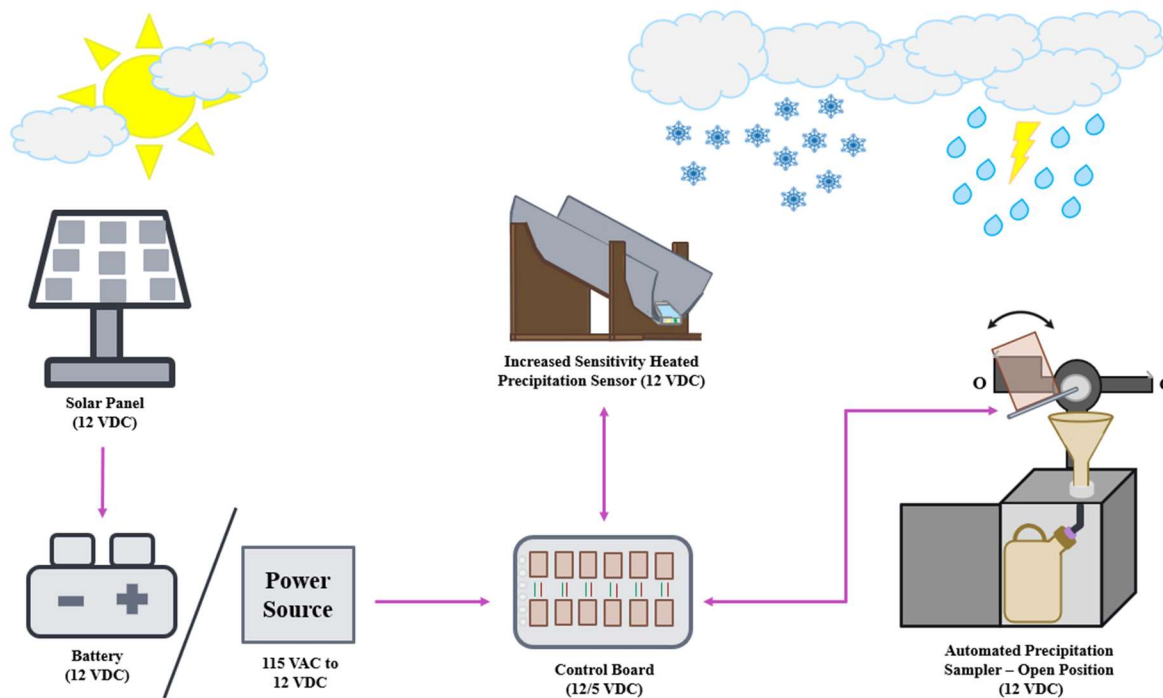
188 collection of analytes of emerging importance and interest in the remote locations of our latitudinal
189 transect.

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191 **2.0 Materials and Methods**

192 **2.1 Precipitation Sampling Array Components**

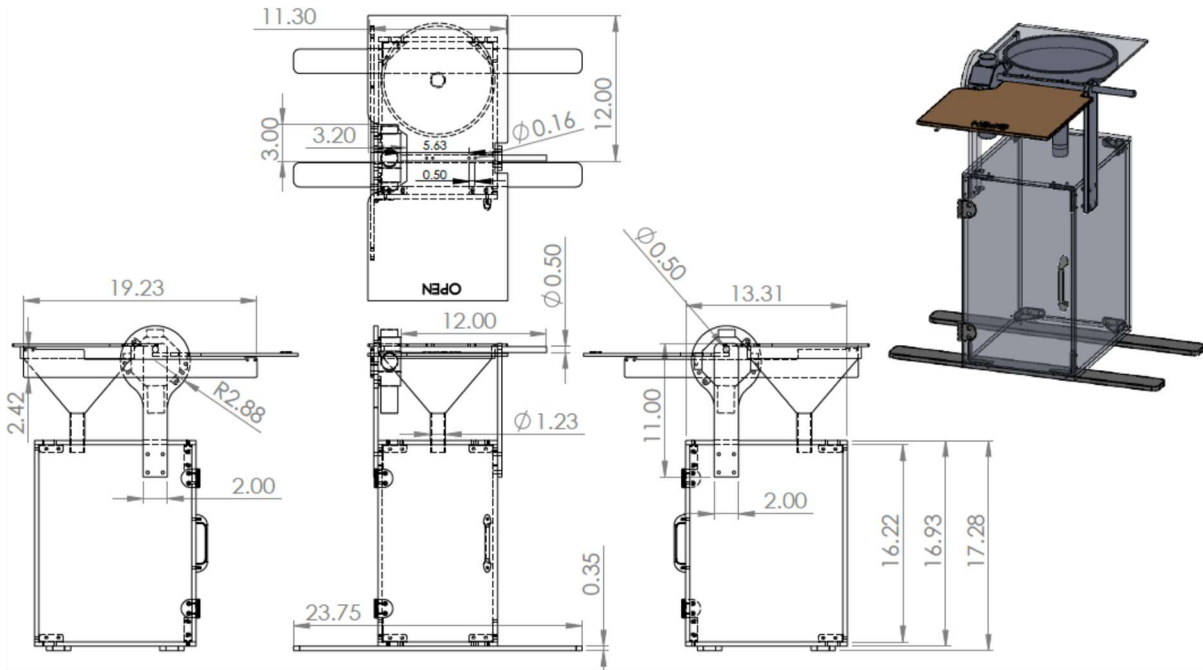
193 Each automated precipitation sampling setup can be operated as an array, here being used
194 in groups of up to six collection units (Figure 1). A collection unit is a simple opaque doored box.
195 The box protects the sample containers against exposure to direct sunlight and provides a mounting
196 location for the funnel and lid, while also facilitating easy exchange of sample containers. The
197 collection units can be fitted with stabilizing legs that allow them to be bolted to concrete or pinned
198 by retaining rods when on soil. In both cases, this prevents tipping and loss of sample during high
199 winds or wildlife-sampler interactions (e.g., Figures 2 and S1). The collection of precipitation is
200 facilitated by a funnel mounted through the top of the sampling unit. The funnel tip extends into
201 the opening of the sample collection container placed inside. The connection can be sealed to better
202 preserve volatile analytes with tubing that passes through a sealed grommet (P/N 9280K34,
203 McMaster-Carr) to enter the sample collection container and minimize evaporative losses.
204 Precipitation events are sampled selectively by modulating the position of a lid over the funnel
205 with an electric motor. The collection unit motors are operated by a digital control board, which
206 interfaces with a precipitation sensor and requires 12 volts of direct current (VDC) power supplied
207 to this system. Switches detecting the lid position ensure complete opening or closure of the funnel
208 mouth for each collection unit.



209
 210 **Figure 1.** Schematic of custom-built automated precipitation sampling array components for off-
 211 grid wet deposition collection. The pink arrows denote the direction of electrical signal and power
 212 exchanged between components. The curved black arrow indicates the rotation of a motorized lid
 213 to obtain open (O) or closed (C) sampler configurations.
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215 **2.1.1 Collection Units**

216 The collection unit materials to date have been made of both 3/8" plywood and black
 217 polyacrylate sheeting. The materials have demonstrated high durability on the order of four years
 218 under field conditions (Figures S1 and S2). Opaque materials were explicitly selected to minimize
 219 photochemical reactions and growth of photosynthetic microorganisms within the sample. The
 220 dimensions of the collection unit are detailed in Figure 2. Each can accommodate sample
 221 containers up to 20 L in volume for collection in locations with large monthly wet deposition
 222 volumes, such as in Newfoundland and Labrador (Table 1).
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224

225 **Figure 2.** Detailed collection unit schematic with all dimensions provided in inches. Further
 226 specifications for the lid dimensions can be found in Figure S3. The shaded 3D rendering depicts
 227 both open and closed states for the lid, positioning of legs to secure it to surfaces, placement of
 228 corner brackets, and the door handle and hinges.

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The box panels can be joined using hardware inserts (P/N 1556A54 and 1088A31, McMaster-Carr, Aurora, OH), 3D printed corners (Figure S4), or along the box edges with screws if using wood. The door is attached with two hinges (P/N 1549A57, McMaster-Carr) and held closed with a magnetic contact (P/N1674A61; McMaster-Carr) or hooked latch. The electric motor controlling the lid is enclosed in a standard polyvinylchloride electrical junction box, which is attached to a short paddle mounted on one side of the collection unit. Here we used an electric worm-gear motor (12 VDC, 2 revolutions per minute; TS-32GZ370-1650; Tsiny Motor Industrial Co., Dong Guan, China) mounted inside the enclosure with matching hex bolts (P/N 91251A146, McMaster-Carr) that passed through the weather-tight cover while the drive shaft protrudes through a 3/8" hole drilled in the cover. The drive shaft has a flat edge to affix the lid rod using a short set screw (Figure S5) that is cemented semi-permanently in place with thread locking compound (P/N 91458A112; Loctite Threadlocker Blue 242; McMaster-Carr). The lid rod is 3/8" aluminum machined on one end to allow connection to the motor drive shaft (Figure S5) with four threaded holes along its length to affix the lid (Figure S3). The lid rod passes through a second mounting paddle on the box that keeps the lid level and capable of isolating the funnel from the

245 atmosphere in the absence of precipitation. The lids used here were made of 1/8" Lexan
246 polycarbonate sheet.

247 Selective precipitation sampling is performed using a logic-based assessment of sensor and
248 switch states (defined in Figure S6) by the control board quadNOR gate chip (Fairchild
249 Semiconductor P/N DM74LS02) which activates the H-bridge motor driver chipset (Figure S7).
250 A 12 VDC signal drives the clockwise or counterclockwise rotation of the motor, installed in a
251 suitable port of the junction box, via a cable from the control board, which passes through a
252 weather-tight compression fitting (e.g., Home Depot SKU# 1000116446). The motor rotation
253 signal is interrupted when the lid makes contact with one of two weather-tight limit switches (P/N
254 SW1257-ND; Omron, Digi-Key Electronics, Thief River Falls, MN) mounted on opposite ends of
255 a horizontal armature connected to the vertical motor mounting paddle (Figure 2). The switches
256 controlling the lid location ensure that the funnel is completely open or covered as necessary for
257 precipitation collection. The funnels used in this work are 20 cm in diameter and made from high-
258 density polyethylene (HDPE; Dynalon, P/N 71070-020, VWR International, Mississauga, ON). A
259 7" x 5" piece of filtration mesh (P/N 9265T49, McMaster-Carr) that was tied together as a fitted
260 cone insert with Nylon thread (e.g., fishing line) to prevent large debris entering the sampler
261 containers when used, for example, in the collection of TF precipitation under a forest canopy
262 when accompanying litterfall is also expected. The exit of the funnel directs the collected
263 precipitation into the narrow-mouth opening the container inside the collection unit, such as 20 L
264 HDPE jugs or 10 L HDPE jerricans (Bel-Art Products; P/N 11215-314, VWR International).

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266 **2.1.2 Heated Precipitation Sensor**

267 The detection of rain modulates the opening and closing of the collection units by an
268 interdigitated resistive sensor (M152; Kemo Electronic GmbH, Geestland, Germany; Figures S6
269 to S8). This approach is consistent with established precipitation detection techniques used by
270 government monitoring programs (e.g., CAPMoN; Canadian Air and Precipitation Monitoring
271 Network, 1985a, 1985b). The rain sensor detects conductive deposition by the completion of a
272 conductive circuit when electrolytes bridge the connection between the interdigitated gold
273 electrodes. The sensor is supplied with 12 VDC from the power system to trigger a relay when
274 precipitation conductance above 1 M Ω ·cm conductivity is detected (determined experimentally,
275 see Section S1). This is equivalent to approximately 8 μ M sodium chloride. The sensor detection

276 limit reflects an upper limit of precipitation ion loading because the design of the rain chute leads
277 to an increase in surface area of more than a factor of 25 on which solutes can accumulate to
278 enhance the ionic content of the deposited water. An output of 12 VDC is sent to the digital control
279 board by the relay when rain is sensed, or 0 VDC in its absence, for signal processing and motor
280 control (Figure S7). When rain is sensed, the lid of each sampler in the array is simultaneously
281 opened (<5 seconds) and is dependent on the rotational rate of the lid motor. To increase the
282 sensitivity of this sensor and to extend the sampling duration when conductive atmospheric
283 constituents are completely washed out of the atmosphere, a sloped tin chute (e.g., Home Depot
284 SKU# 1001110514) was added to extend the surface of the rain sensor. The sensor was placed at
285 the end of the chute and sealed in place with caulking to allow water droplets to move easily from
286 the chute onto the sensor.

287 The angle of the chute can be adjusted to control the momentum of collected droplets so
288 that they collect on the sensor surface and only flow off it when the rate of precipitation exceeds
289 the sensor evaporation capability. When soil is available, two bent rods can be used to hold the
290 chute at the optimized angle of 10° (Figure S2). They are inserted into the soil and the chute is
291 affixed to the tops of the rods with zip ties passed through small holes drilled in the sides of the
292 chute, which are subsequently sealed with caulking. When soil is unavailable, for example in urban
293 environments, we have created a mounting frame to hold the chute at the optimized angle of 10°
294 (Figure S8). When precipitation is detected the sensor surface draws current up to 1.0 ampere (A)
295 into a heater to actively evaporate water from its surface so that it accurately detects the active
296 period of rain events. The heated sensor has undergone preliminary field tests and is also capable
297 of detecting ice and snow, provided they contain electrolytes.

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299 **2.1.3 Power Supply Systems**

300 Power for this system can be supplied from a battery at 12 VDC or using a 115 VAC to 12
301 VDC transformer power supply (P/N 285-1818-ND; TDK-Lambda Americas, Digi-Key
302 Electronics). Depending on the duration of sampling and the time of year, the battery capacity can
303 be changed to suit power needs (Section 3.2.2). To provide sufficient power density in this study,
304 over one to two month-long collection periods, the battery capacity was carefully matched; with
305 top-up options implemented when prolonged or high-frequency precipitation was expected.
306 Absorbent glass mat (AGM) marine deep cycle batteries can withstand discharge events down to

307 less than 60 % capacity and are robust under nearly all environmentally relevant temperatures (\leq
308 -20°C to 40°C). Additionally, these batteries interface easily with solar charging options as they
309 are able to accept high current input. Monthly collections in Newfoundland were powered with 76
310 amp-hour (Ah) AGM batteries (Motomaster Nautilus; Ultra XD Group 24 High-Performance
311 AGM Deep Cycle Battery, 12 VDC) topped up by a 40 W solar panel interfaced with a charge
312 controller to prevent overcharging (Coleman; Model # 51840, max current of 8 A at 14 VDC).

313 For collections made every second month in Labrador, a 120 Ah battery with the same
314 solar top-up strategy was used to ensure continuous operation. For either remote field deployment,
315 batteries and charge controllers were housed in a Pelican™ case (Model 1440, Ocean Case Co.
316 Ltd., Enfield, NS) fitted with weathertight bulkhead cord grips (P/N 7529K655, McMaster-Carr)
317 through which charging and power cables were passed (Belden, Coleman; S/N 7004608,
318 70875227, Allied Electronics, Inc., Ottawa, ON). Humidity in all weatherproof cases was
319 minimized by exchanging reusable desiccant packs (Ocean Case Co. Ltd.) when depleted batteries
320 were exchanged for fully charged replacements. Solar panels were repositioned monthly to
321 optimize orientation for solar power provision. Using either power source, the control board
322 converts and distributes the 12 VDC to the other components in the precipitation sampling array.

323

324 **2.1.4 Custom Control Board**

325 A custom control board to operate a six-collection unit array was designed based on prior
326 digital logic circuits for standalone collectors (VandenBoer, 2009). The 12 VDC battery or
327 transformer output is supplied directly to the rain sensor and relay, as well as to the motor drivers
328 for lid opening (Figure S9). Each collection unit is controlled independently to ensure lids are fully
329 opened or closed, thereby requiring six replicate motor driver control circuits that respond to their
330 independent switch signals. The remainder of the signaling and digital logic operates on 5 VDC
331 which is produced by on-board voltage regulators (Micro Commercial Co; P/N MC7805CT-BP,
332 Digi-Key Electronics). The lid switches are provided with 0 and 5 VDC to indicate collection unit
333 open or closed status (Omron Electronics; P/N D2FW-G271M(D), Digi-Key Electronics). The
334 signals from the sensor and switches connect to the board through four-conductor cable (Belden;
335 S/N 70003678, Allied Electronics Inc.) passed through weathertight bulkhead cord grips and
336 secured to screw terminals (Figure S9). The sensor and switch signal inputs interface with a quad
337 NOR GATE chipset (Texas Instruments; P/N 296-33594-5-ND, Digi-Key Electronics) to trigger

338 the motor driver (STMicroelectronics; P/N 497-1395-5-ND, Digi-Key Electronics) such that it
339 rotates or remains stationary. The additional resistors, capacitors, and diodes are necessary to
340 maintain stable signaling throughout the printed circuit board (Figure S9, Table S1).

341 The custom control board was housed in a Pelican™ case (1400 NF; Pelican Zone,
342 Mississauga, ON) fitted with cut-to-use foam inserts and a reusable desiccant pack that was also
343 exchanged alongside those for the battery cases. All collection units, sensors, and power supply
344 cables were passed through eight weathertight bulkhead cord grips and fixed to screw terminals
345 on the board. The opposing ends of the cables were fitted with weathertight Bulgin Buccaneer 400
346 or 4000 Series circular cable connectors (Table S2; Allied Electronics, Inc.) to allow easy field
347 installation with mated connectors on the cables originating from each of the previously mentioned
348 array components. Connected cables could then be buried in shallow soil trenches to reduce the
349 attention of gnawing animals, as well as potential entanglement hazards with other wildlife.
350 Precipitation events were logged from the control boards using a HOBO 4-channel analog data
351 logger (UX120-006M; Onset®, Bourne, MA) that records the sensor, switch, and motor voltages.
352 The fourth channel is reserved to monitor battery or power supply voltages over time (Section 3.2).

353

354 **2.2 Power Demand and Management Tests**

355 Power demand was calculated based on the cumulative component requirements prior to
356 the selection of batteries. This was to ensure adequate capacity to collect samples over one to two
357 month-long field deployments and are sufficient for an assumed worst-case scenario of one week
358 of constant rain without solar power charge restoration. Solar panel power production capacity
359 was determined based on the calculated energy required to recharge the battery. As a result, we
360 selected the 40 W panel which could complete charging at 14 VDC with a week of direct sunlight
361 at 8 hours per day. The power demand for a six-sampler array was measured in standby and during
362 operation with a digital power meter (Nashone PM90, Dalang Town, China) in real-time when
363 supplying 12 VDC with a transformer. Contrasting power demand tests were performed under
364 different environmental conditions and power management configurations. The first was
365 performed using the 76 Ah AGM battery with a solar top-up in an urban environment from July
366 through August 2018, while the other was performed using a 103 Ah AGM battery alone from
367 January through February 2019.

368

369 **2.3 Continuous Monthly Collection of Remote Samples at NL-BELT**

370 One array of six automated collection units (3 OF, 3 TF) were deployed within one forested
371 experimental field site located in each of the four watershed regions of the NL-BELT (24 samplers
372 in total) between 2015 and 2016. Additionally, between one to three total deposition samplers were
373 located at each of the four field sites (Table 1, Figure S10). The watersheds span 5.5° latitude from
374 the southernmost site Grand Codroy (GC), through the colocated Pynn’s Brook (PB) and Humber
375 River Camp 10 (HR) sites, to Salmon River (SR) as the highest latitude site on the island of
376 Newfoundland. The northernmost forested watershed, Eagle River (ER), is located in southern
377 Labrador and extensive details characterizing each of the four sites can be found in Ziegler et al.
378 (2017). All sampling locations are far from anthropogenic pollutant point sources, except for the
379 ubiquitous presence of marine sea spray from the nearby marine coastlines. The total deposition
380 samplers were identical to the automatic collection units except that they were not fitted with a
381 motor arm and lid, so they did not require a source of power. Three of the six automated samplers
382 were deployed in the open at a distance from the forest stand, equal to or greater than the height
383 of the trees, in line with CAPMoN and NADP guidelines. The other three automated samplers
384 were placed under the canopy to collect TF precipitation within the forest sites. These samplers
385 actively collected wet deposition into integrated monthly (Newfoundland) or two-month
386 (Labrador) samples during snow-free periods (approximately June through November). The arrays
387 were collected and stored during the winter months while total deposition samplers remained in
388 field locations year-round. It is also important to note that during the growing season, sample
389 collections were made at the same time – that is, OF and TF deposition were collected on a single
390 day at each sampling site and within a few days of each other across the transect. Collected sample
391 volumes were compared between the automated samplers and total deposition collectors for each
392 collection interval as a check on proper function (i.e., less than or equal volumes in automated
393 samples). During each site visit, the slope of the sensor was confirmed to be correct, sample
394 containers were collected and replaced with clean units, the battery and desiccant packs replaced
395 with fully recharged devices, and the entire array confirmed operational.

396

397

398

399

400 **Table 1.** NL-BELT sampling site details provide locations and identifiers, alongside those from
 401 long-term weather stations operated by Environment and Climate Change Canada (ECCC). Soil
 402 pH was determined from samples collected at the same time as precipitation. Mean annual
 403 temperature was derived from ECCC climate normals. Annual total deposition precipitation
 404 volumes were either measured for the 2015-16 period (ECCC, This Work) or calculated by the
 405 Oak Ridge National Lab DAYMET archive.
 406

Sampling Site	Sampling Site Location	Station (Climate ID)	Station Location	Soil pH ^a	MAT (°C) ^b	Average Annual Precipitation (L)		
						ECCC ^c	DAYMET ^g	This Work
Grand Codroy (GC)	47°50'43.1"N 59°16'16.0"W	Stephenville A (8403801)	48°32'29.00" N 58°33'00" W	3 to 4	5.0 ^e	53.2	58.9	45.6 (+5.17)
Pynn's Brook (PB)	49° 05' 13.20"N 57° 32' 27.60" W	South Brook Pasadena (8403693)	49°01'00" N 57°37'00" W	3 to 4	4.6 ^e	21.4	54.3	38.6 ^h
Salmon River (SR)	51°15'21.6"N 56°08'16.8"W	Plum Point (40KE88)	51°04'00" N 56°53'00" W	3 to 4	2.4 ^e	47.1	45.4	32.3
Eagle River (ER)	53°33'00.0"N 56°59'13.2"W	Cartwright A (8501100)	53°42'30" N 57°02'06" W	3 to 4	0 ^d	- ^f	56.3	25.8

407
 408 ^aSoil pH for the organic and mineral soil horizons determined by addition of 400 µL of aqueous 0.5 M CaCl₂ to a 50:50 w/w slurry
 409 of dried soil in deionised water. Note: the four remote NL-BELT sites are dominated by balsam fir trees underlain by humo-ferric
 410 podzol soil with pH ranging between 3.0 and 4.5.
 411 ^bEnvironment Canada: Canadian Climate Normals, 1981 to 2010, https://climate.weather.gc.ca/climate_normals/ (last accessed:
 412 14 July 2023).
 413 ^cAt least 20 years of measurements.
 414 ^dThe World Meteorological Organization's "3 and 5 rule" (i.e., no more than 3 consecutive and no more than 5 total missing for
 415 either temperature or precipitation).
 416 ^eAnnual precipitation averages determined using ECCC daily precipitation reports.
 417 ^fLarge quantity of missing data for this location from January 2015 to December 2016 prevents any reliable estimate.
 418 ^gEstimated deposition rates converted to volume using DAYMET (Thornton et al., 1997, 2021, 2022).
 419 ^hVolumes merged for 2015 and 2016 at PB and HR.
 420
 421

422 2.3.1 Sample Preservation

423 Four of the six sample containers (two each of OF and TF) were biologically sterilized
 424 using 1 mL of a saturated aqueous solution of mercuric chloride (HgCl₂) to preserve against
 425 biological growth and loss of bioavailable nutrients over the collection periods. Unsterilized
 426 sample containers (without HgCl₂) were used for measurements of recalcitrant species and to
 427 assess any matrix effects exerted on target analyte quantitation. The use of HgCl₂ as a sample
 428 preservation technique has been long-studied and well-established (Kirkwood, 1992; Kattner,
 429 1999); thus, additional tests to verify the preservation of collected chemical species over time were

430 not performed. The analysis of deposition collected in unsterilized and sterilized containers,
431 however, serves as a method for internal sample validation - as does our evaluation of measurement
432 outcomes in comparison to those reported within the literature. Collected sample volumes were
433 measured with a 1000 ± 10 mL graduated cylinder and aliquots were collected for chemical
434 analysis via transfer to precleaned 500- or 1000-mL HDPE containers (Nalgene; VWR
435 International). Samples were stored at 4°C before returning to the laboratory where they were
436 filtered with a 1000 mL Nalgene vacuum filtration system (P/N ZA-06730-53; ThermoFisher
437 Scientific, Waltham, MA), fitted with $0.45\ \mu\text{m}$ polyethersulfone filters (PES, P/N HPWP 04700,
438 EMD Millipore), to remove suspended solids. Filtered samples were transferred to new clean
439 HDPE containers and stored for up to two months at 4°C in a cold room until analysis. The target
440 analytes in this work are non-volatile and the described sample collection methods consider this
441 analyte property, as well as their interactions with container materials. The versatility of the system
442 design allows for the use of different collection materials, keeper solvents for volatile organics,
443 etc., so that the experimental design can be analyte specific, depending on end user needs. Sample
444 preservation approaches should thus be identified by users of this new platform based on their
445 scientific objectives and review of the literature (Galloway and Likens, 1978; Peden et al., 1986;
446 Dossett and Bowersox, 1999; Wetherbee et al., 2010). In addition to the internal validation
447 approach described here, we aim to demonstrate that the precipitation samplers in this work are
448 suitable for measuring conductive deposition on- and off-grid. Below we highlight autonomous
449 off-grid operations, determine the fraction of conductive rainfall collected from the total volume
450 of precipitation, and validate our measurements through comparison to the literature.

451

452 2.4 Cleaning and Preparation of Sample Containers

453 All sample collection and storage containers, as well as all sample handling apparatuses,
454 were made of HDPE or polypropylene for the quantitative analysis of target analytes. Prior to use
455 in handling samples, these were all acid-washed in 10 % v/v HCl (P/N BDH7417-1; VWR
456 International) followed by six sequential rinses with distilled water and ten rinses with 18.2
457 $\text{M}\Omega\cdot\text{cm}$ deionised water (DIW; EMD Millipore Corporation, Billerica, MA, USA). Containers
458 were dried by inversion on a clean benchtop protector overnight, or with protection from dust using
459 lint-free lab wipes over container openings when necessary. Field and method blanks were
460 collected through the addition of DIW to cleaned containers, and/or sample handling devices, in

461 order to quantify appropriate method detection limits and to identify any sources of systematic or
462 random contamination. Blank subtraction was applied to measurements, where appropriate.

463

464 **2.5 Measurements of pH and Conductivity**

465 The pH and conductivity of each sample was determined using a ThermoScientific™ Orion
466 Versa Star meter (ORIVSTAR52) interfaced with a pH electrode (Model: 8157BNUMD, Ultra
467 pH/ATC Triode, ROSS) and 4-electrode conductivity cell (Model: 013005MD, DuraProbe,
468 ROSS). Prior to use, the probes were calibrated daily with standard solutions specific for these
469 probes (ThermoScientific™ Orion™ conductivity standard 1413, and pH 4, 7, and 10 buffers) and
470 then stored between analyses according to manufacturer directions. Aliquots of 15 mL of
471 precipitation from archived samples were subsampled into 40 mL polypropylene Falcon tubes.
472 This was followed by immersion of a cleaned electrode for the conductivity measurement,
473 followed by the pH probe measurement to prevent conductivity bias due to potassium chloride
474 migration across the glass frit of the pH probe. Readings were recorded once signals had stabilized.

475

476 **2.6 Measurements of Dissolved Organic Carbon (DOC)**

477 Measurements of DOC were performed by catalytic combustion of samples in a platinum
478 bead-packed quartz furnace at 720°C to quantitatively produce CO₂, followed by non-dispersive
479 infrared absorption spectrophotometry using a Shimadzu Total Organic Carbon (model: TOC-V)
480 analyzer and an autosampler (model: ASI-V). Cleaning of materials prior to DOC determination
481 follows the same procedure as for the sample containers. Precipitation aliquots of at least 12 mL
482 were transferred to clean and combusted (500°C, 5 hours) 40 mL borosilicate glass vials, then
483 capped and stored at 4°C until analysis. Prior to analysis, vial caps were replaced with cleaned
484 polytetrafluoroethylene-lined septa. Inorganic dissolved carbon (e.g., H₂CO₃) was purged from
485 samples by acidification to pH 2 with HPLC grade H₃PO₄ (20 % v/v) and bubbling with an inert
486 carrier gas. Samples were analyzed in triplicate and quantified using calibrations spanning 0.1 to
487 10 or 10 to 100 ppm (mg C L⁻¹) with potassium hydrogen phthalate (KHP), depending on the
488 relative sample concentration range. Accuracy and precision were assessed using 1 and 10 ppm
489 KHP check standards analyzed every 10 injections, respectively. Calibrations were performed at
490 the beginning of every analysis day.

491

492 **3.0 Results and Discussion**

493 In addition to the general design advantages in the section that follows, we present the
494 results of various physical and chemical parameters to validate this new open source custom-built
495 modular system. The power consumption and snow-free performance testing are used to
496 demonstrate the off-grid capabilities of these samplers, as are the two-year datasets. The lower
497 power requirements are compared to existing commercial samplers and paired with solar top-up
498 to prolong the use and reduce the need to replace batteries on timescales shorter than planned
499 sampling duration (i.e., < 1 month). We then evaluate the automated wet deposition volumes, in
500 which the samplers prevent dilution during atmospheric washout events, compared to total
501 volumes collected from co-located samplers to depict the fractionation by volume as a function of
502 time. We also investigate the advantages of replicates collected across the four watersheds, using
503 deployments of triplicate samplers under field conditions. The ratio of collected TF to OF
504 replicates highlights the ability of these samplers to capture the dynamic nature of precipitation
505 interacting with forest canopies. Simple pH and conductivity measurements are then used as
506 benchmarks to situate the NL-BELT data within the established literature to emphasize the robust
507 operation of the samplers and impact of the selective sampling. Fluxes of DOC are also
508 interrogated across all four sampling sites as we demonstrate the potential of these samplers to
509 make measurements of more complex analyte pools that are of current interest to the atmospheric
510 measurement community.

511

512 **3.1 General Design Advantages**

513 While several precipitation collectors have been similarly developed to address specific
514 scientific objectives – e.g., the quantitation of dust in wet and dry deposition (Laurent et al., 2015;
515 Brahney et al., 2020), determination of ions and DOC in a tropical rainforest (Germer et al., 2007)
516 and urban environments (Audoux et al., 2023), here we present a more general design for modular
517 adaptability. When compared to other precipitation collection apparatuses, the automated
518 precipitation sampler developed in this work has several advantages. Most notable is the ability to
519 collect integrated samples at remote locations by exploiting its off-grid capabilities. Our approach
520 also maximizes the sensitivity of the rain sensor as long as electrolytes remain in the water reaching
521 it. The chute ensures that even if the precipitation contains ultra-trace analyte quantities, they are
522 still collected and quantified for an extended period when high-purity water may be deposited

523 during an atmospheric wash-out event. The chute does this by accumulating water-soluble
524 materials between rain events that require time to be completely washed off and through the release
525 of ions from the material itself, which ages under environmental conditions. As the conductivity
526 of the precipitation falls below the sensor threshold – conductive precipitation being that which
527 initially contains high solute levels that progress through trace level concentrations, the added ions
528 from the chute prolong the collection of rain past this time point. In rainfall events where extended
529 atmospheric wash-out occurs, where precipitation becomes ultrapure water, the sampler lids will
530 eventually close – preventing dilution of the sample while maintaining the collection of analytes
531 of interest. A recent study has found that rainfall events could exhibit variability and the lower
532 atmosphere can be supplied with aerosols due to specific sources, atmospheric dynamics, and
533 meteorological conditions (Audoux et al., 2023). If this occurs, the automated lid will reopen to
534 sample the polluted air masses. In application to trace pollutants, this also reduces methodological
535 sample preparation time as it decreases the extent to which additional handling steps, like solid-
536 phase extraction, are required prior to analytical determinations.

537 The six replicate measurements used in each array provide a means of assessing sampling
538 reproducibility (e.g., canopy TF has expected heterogeneity) and for multiple analyte classes to be
539 targeted. Various analytes, with different chemical properties and/or contamination considerations,
540 can be targeted by changing the materials used for the components that encounter the sample (i.e.,
541 lids, funnels, and sample holding containers). Replicate collection can also allow for selective
542 sample preservation when quantifying deposited chemical species that may be reactive, volatile,
543 or biologically transformed. The modularity of the overall system design also allows the collection
544 units to be dismantled entirely and easily reassembled on-site, minimizing logistical issues and
545 costs for transport to remote regions. Lastly, these collection units are cost-effective. We were able
546 to produce four arrays, each consisting of six collection units, at a fraction of the cost of a single
547 equivalent commercial off-grid automated precipitation sampling unit.

548 With the majority of commercial precipitation samplers requiring a source of electricity,
549 on-grid sample collection necessitates high infrastructure costs and/or samplers being positioned
550 closer than desired to point sources of anthropogenic pollution. As a result, especially in remote
551 locations, site selection becomes heavily restricted and expensive when factoring in all the
552 standard criteria, particularly with respect to the need for an easily accessible power source. Thus,
553 the off-grid capabilities of our samplers lend dexterity to these systems and makes deposition

554 sampling that follows standard siting guidelines, like those of CAPMoN or NADP but without
555 power, more accessible to the global atmospheric research community (Vet et al., 2014). To further
556 highlight and validate their capabilities, a series of fundamental performance parameters were
557 collected and are discussed in detail in the sections that follow.

558

559 **3.2 Power Consumption and Performance Testing**

560 **3.2.1 Power Consumption of Instrumental Setup**

561 The simplicity of the automated precipitation samplers allows for low power consumption
562 during operation, which is particularly important for off-grid operation. The motors operating and
563 rain sensor heating during active precipitation are the most energy-intensive elements of the system
564 (Table 2). The integrated contribution of the motor over a month-long sampling period is however
565 negligible compared to other components, since it is operational for short periods of 5 to 10
566 seconds with a current usage of only 38 mA. The continuous need to provide 5 VDC to the digital
567 logic via step-down from 12 VDC is actually the largest power consuming component of the setup
568 in the absence of rain. When the samplers are in the closed position, under rain-free conditions,
569 the power consumption of the entire array is 4.66 Watts (W) and 2.86 W for transformed 115 VAC
570 and battery 12 VDC supplies, respectively. The provision of 12 VDC to the board with a
571 transformer for the 115 VAC application results in greater total power requirements. These values
572 increase to 10.00 W and 5.04 W with the detection of a conductive liquid on the precipitation
573 sensor as it heats the sensor surface to capture the active period of the event. Based on the measured
574 power consumption, a fully charged 103 Ah AGM battery would provide at most 447 hours (or 18
575 days) in standby mode under rain free conditions and 294 hours (or 12 days) if the heated surface
576 of the sensor is in continuous use (Table 2). The lower range limit is unlikely since the sensor only
577 operates for the duration of a rain event, after which the battery is available for solar top-up again.
578 In the fieldwork conducted here, battery life was extended through the addition of 40 W solar
579 panels to the systems. The entire array was confirmed to be operational at the end of monthly (SR,
580 PB, and GC) and two month (ER) integrated sampling periods on an ongoing basis, prior to
581 exchange with a new fully charged battery, for two years.

582

583

584 **Table 2.** Measured voltage, current, and power consumption of the rain sensor and circuitry in
 585 both the idle and maximally operational state when connected to a 12 VDC battery or transformed
 586 115 VAC. Total power demand was measured for wet and dry sensor scenarios.

Parameters	Rain Sensor		AC Outlet		DC Battery		Total			
	Idle	Active	Idle	Motors	Idle	Motors	AC Outlet		DC Battery	
			Board	In-Use	Board	in-Use	Dry	Wet	Dry	Wet
Voltage (V)	12 DC	12 DC	114 AC	110 AC	12 DC	12 DC	-	-	-	-
Current (A)	0.008	0.120	0.040	0.078	0.230	0.300	-	-	-	-
Power (W)	0.10	1.44	4.56	8.58	2.76	3.60	4.66	10.00	2.86	5.04

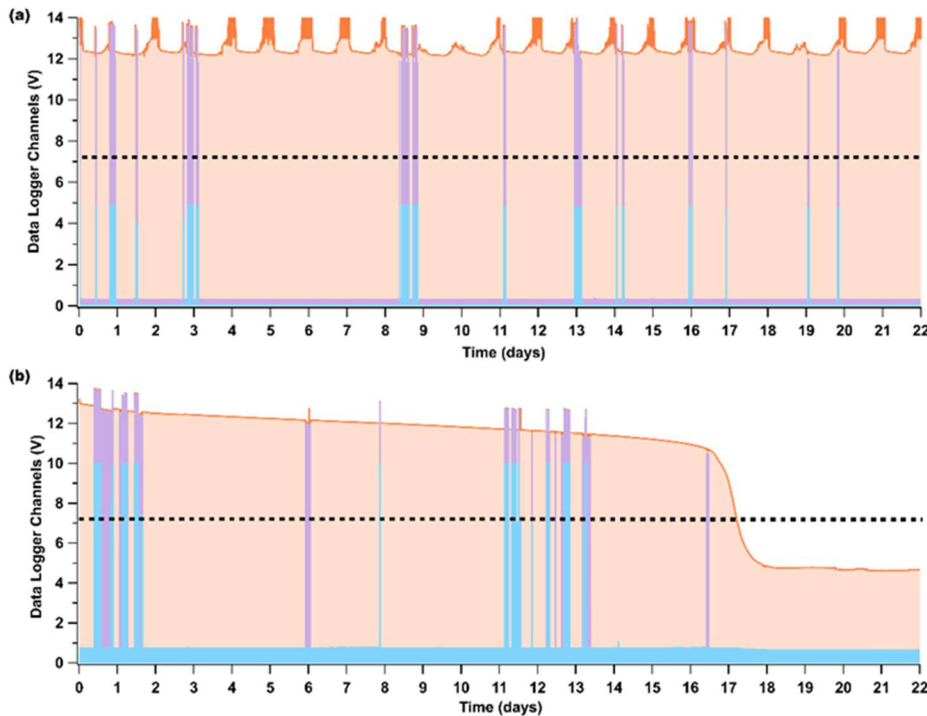
587

588 In comparison to two commercial samplers used by national monitoring networks, the
 589 power requirements of our new samplers are substantially lower. The first commercial sampler we
 590 reviewed draws a maximum of 2 A, with a ceramic heater housed within the sampler case that
 591 draws 0.8 A constantly, resulting in an upper limit power demand of 230 W (at 115 VAC) and a
 592 lower limit of 92 W. The commercial sampler can be upgraded to utilize a thermostated space
 593 heater for winter operation, drawing an additional 4.2 A (480 W), resulting in a maximum power
 594 demand of about 800 W when using a 115 VAC power supply. A second commercial precipitation
 595 sampler reviewed is used by national monitoring networks and draws approximately 5 A, resulting
 596 in a power requirement of 575 W at 115 VAC. The commercial and standard precipitation samplers
 597 for deposition monitoring programs have much higher power requirements compared to those
 598 presented in this work. The commercial samplers utilize 80 to 100 times more power. With our
 599 lower power requirements, the new automated samplers prove to be advantageous in both on- and
 600 off-grid sampling yet are disadvantaged in being unable to collect snow in the winter.

601

602 3.2.2 Precipitation Sampler Performance Tests and Data Logging

603 In addition to low power consumption during precipitation sampling, a supplied battery
 604 can obtain constant power renewal when outfitted with a solar top-up that is kept exposed to
 605 sunlight by proper orientation. At NL-BELT, adjustments were made for this during each site
 606 visit during sample collection. During the solar top-up tests below, voltages of the sensor and
 607 batteries were consistently monitored. Over a test period of 22 days, no appreciable decline in
 608 battery performance of a 76 Ah unit was observed despite the detection of more than 10 rain
 609 events during that period (Figure 3a).



611

612 **Figure 3.** Performance of off-grid precipitation samplers during sample collections from (a) 13
 613 July to 7 August 2018, using a 76 Ah battery and solar panel top-up and (b) 22 January to 13
 614 February 2019 with a 103 Ah battery and no solar panel. Battery voltage (shaded orange) is
 615 elevated above 12 VDC when charging, or decreases over time when no solar panel is used and
 616 precipitation is sensed/collected. The 12 VDC rain sensor relay signal (purple) and the open
 617 sampling lid switch voltage (blue) indicate active periods of detected precipitation. The black
 618 dashed line indicates the 60% efficiency cut off, 7.2 V, at which the battery should be recharged.
 619

620 In comparison, winter sampling with these devices is not recommended without substantial
 621 investment in a sufficient power density provided high-performance cold weather batteries. The
 622 lack of sunlight during winter at higher latitudes also negates the use of effective small scale solar
 623 top-up. Our tests show that when the samplers were deployed without a solar backup under snow-
 624 free winter conditions (temperatures ranging from -17.8 to 7°C), with a 103 Ah battery, the off-
 625 grid system only lasted for 17 days. At this point, the larger capacity battery was fully depleted by
 626 frequent snow and rainfall – probably due to the heated precipitation sensor requiring additional
 627 energy to phase change snow and ice to water and then to evaporate that water. This depletion
 628 occurred despite housing the battery in an insulated enclosure during the test. In addition, on days
 629 6 and 16, the precipitation sensor relay was activated but the lid did not rotate to the open position
 630 (Figure 3b, blue trace). This could have been because the precipitation event was not intense

631 enough for the lid to open fully and trigger the 5 V lid open switch or because of snow and ice
632 buildup around the lids resulting in them being unable to physically open. Overall, these samplers
633 may be possible to deploy during the winter if line power can be supplied. Such a deployment
634 would further necessitate that the sampling funnel be heated to render a liquid sample for collection
635 in the jugs in addition to the sensor chute to prevent snow and ice accumulation. A heated funnel
636 would also prevent snow or ice accumulation on top of the automated lids. Together, such power-
637 hungry requirements for winter operation exceed simple off-grid use with a battery package that
638 is easily transported into and out of remote field sites.

639

640 **3.3 Comparison of Sample Collection Volumes**

641 The automated samplers were colocated with total deposition samplers and deployed across
642 the experimental forests of four NL-BELT regions during the 2015 and 2016 growing seasons to
643 observe deposition trends. In addition, we compare these observations to the long-term climate
644 normals reported by ECCC and estimated deposition at 1 km x 1 km resolution from the DAYMET
645 reanalysis model (Table 1, Section S2). Three automated samplers were deployed in the open to
646 collect incident precipitation (OF) and another three under the experimental forest site canopy
647 (TF). The mean OF volumes of triplicate measurements from south to north were 1.42, 1.38, 1.31,
648 and 0.79 L, whereas the corresponding TF volumes were generally similar in magnitude at 0.96,
649 0.98, 1.02, and 1.13 L, for the 2015-16 sampling period (Figure 4). It is evident that the volume of
650 precipitation decreased as latitude increased for OF samples, whereas the opposite relationship
651 was observed in TF samplers, although the absolute volumes are more comparable in magnitude.
652 The total deposition volumes collected were as expected, decreasing from south to north in
653 agreement with the expectations from the long-term normals and comparable to the estimates from
654 the DAYMET model (Table 1), where the largest integrated volume of precipitation was collected
655 at the lowest latitude (GC) and a lower amount at the highest latitude (ER), with the intermediate
656 sites (HR and PB) having the lowest inputs overall during this observation period. Total annual
657 deposition volumes collected by our deployed samplers from south to north in 2015 were 39.5,
658 39.4, 31.9, and 17.5 L, while in 2016 they were 51.7, 37.8, 32.8, and 34.2 L. Total deposition
659 volume collected from HR was used for comparison to automated sample volumes collected at PB
660 in 2015, as they both share the same watershed. This approach had to be taken, as the HR site was
661 initially planned for full experimental use before becoming inaccessible in early 2015. The relative

662 error between the two sites for samples collected in 2016 was $\pm 15\%$ (24.6 L in PB and 32.2 L in
663 HR), comparable to the reproducibility we observe for replicates collected within a given site (see
664 below). The total deposition samplers were installed in HR in late 2014 and the automated samplers
665 were then set up at PB. Despite this, there is good agreement between the trends in predicted
666 deposition values by DAYMET with the measured values, although the absolute amounts from
667 these are systematically lower in all of our observations (Section S2). Regardless, by following the
668 recommended siting criteria from the NADP and CAPMoN as best as possible, the very strong
669 agreement of our temporal trends at both annual and monthly timescales with both comparators
670 demonstrates the suitability of the total deposition samplers and, therefore, the automated samplers
671 for use in quantifying deposited chemical species of atmospheric interest into the experimental
672 sites.

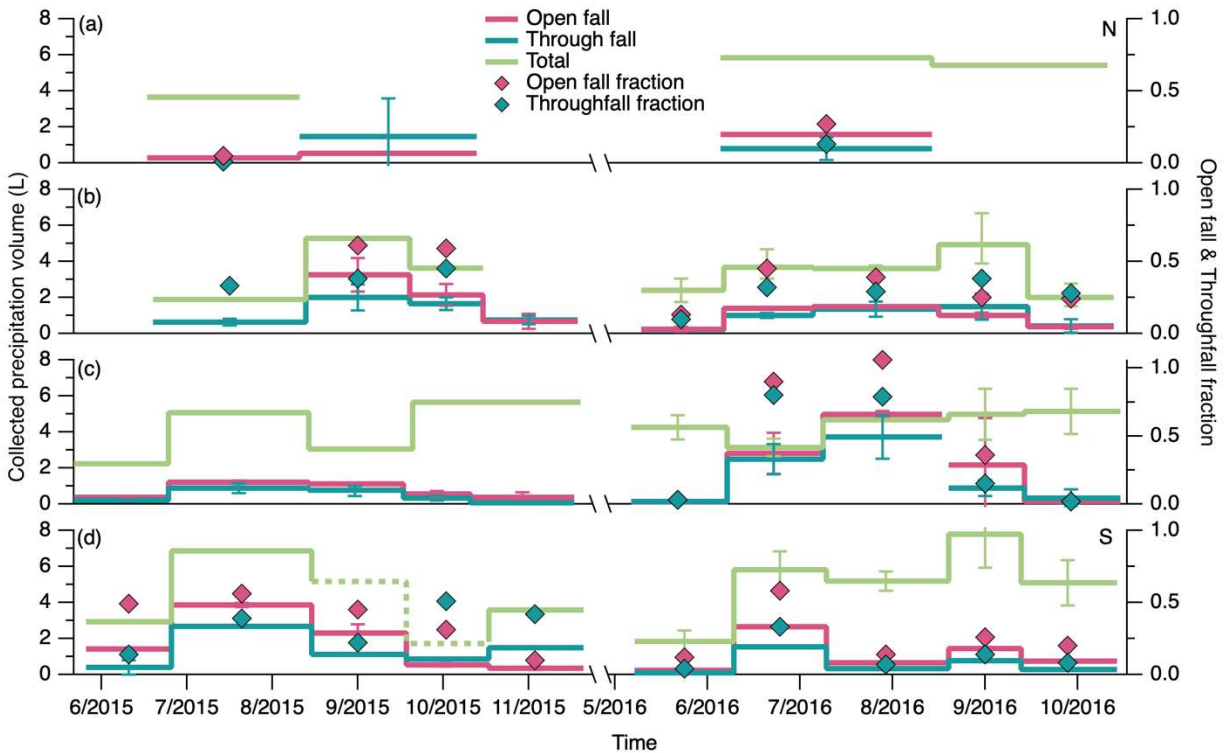
673 The wet deposition volumes collected for the snow free period using the automated
674 precipitation samplers did not follow the trends in total deposition (Figure 4), as might be expected
675 (e.g., due to pollutant loading, rainfall quantity/rate, and scavenging processes). For the 2015
676 collection period from June through October, the summed volumes of OF precipitation, from south
677 to north across the NL-BELT, were 25.4, 10.9, 20.4, and 2.2 L, while in 2016 they were 17.3, 30.4,
678 13.5, and 5.1 L. While the total and OF fractions would typically be much closer to unity in more
679 polluted regions, it would be expected in these remote NL-BELT field sites for the differences to
680 be driven by complex, non-linear processes that cannot be easily disentangled. Here we present
681 three reasons as to why the measured wet OF deposition volumes do not follow the total deposition
682 trend across the transect. First, these samplers are designed specifically to collect only conductive
683 precipitation (i.e., containing conductive atmospheric compounds) not total/bulk precipitation. As
684 a result, the OF wet deposition volume collected across the sites is mostly below 50% of total
685 volumes collected, while TF volumes are similar in magnitude or lower than that of OF (Figure
686 4). The wet deposition fraction collected was variable within and between regions, sometimes less
687 than 10%, despite large volumes collected in total and presumably due to intense atmospheric
688 washout that this region is well-known for. Second, the NL-BELT total deposition trend estimated
689 using the ECCC long-term climate normals represents a 30-year period (Bowering et al., 2022)
690 while the automated volume measurements here represent two years of targeted conductive
691 precipitation collection. The combined summed volumes of targeted conductive wet deposition
692 across the 2015 and 2016 field seasons were 42.7, 41.3, 33.9, and 7.3 L, somewhat better reflect

693 the expected precipitation trends within the transect (Table 1). Lastly, our monthly automated wet
694 deposition sample collection periods occurred from June through November and so it is temporally
695 incomplete with respect to the substantial amount of precipitation volume deposited as snow
696 delivered during the winter (Table S3). The discrepancies between the long-term trends and our
697 shorter-term observations therefore make sense as they are sensitive to interannual changes in
698 synoptic scale transport and rainwater solute loadings, as exemplified by the volumes collected in
699 SR in 2015 (Figure 4b) and PB in 2016 (Figure 4c). Overall, for the automated sampler
700 observations on a per-year basis, there is no consistent trend between site latitude and the volume
701 collected in either OF or TF. This is unsurprising as they are dependent on the conditions that drive
702 the rate of atmospheric wash-out and presence of conductive solutes.

703 The automated OF wet deposition volumes collected each year have peak values that range
704 from 1 to 4 L with an overall variability of $\pm 33\%$ for any triplicate of samples across the entire
705 dataset. Across our 33 sample collection periods, our replicate relative standard deviations (RSDs)
706 follow a log-normal distribution where volume reproducibility is typically within $\pm 12.5\%$ and
707 almost always within $\pm 31.5\%$ (Figure S11). A few outliers with higher variability skew the overall
708 view of volume precision. Out of 33 OF samples collected, 10 have RSDs greater than 40.5% with
709 2 of those 10 having RSDs greater than 100%. Those values greater than 40.5% had no systematic
710 relationship with site or time of time. Wind speeds were considered as a possible source of
711 variability. The prevailing winds over Atlantic Canada are known to be southwesterly in the
712 summer – intensifying during the autumn months – and westerly to northwesterly in the winter
713 (Bowyer, 1995; Jacob, 1999; Randall, 2015). Strong wind speeds (i.e., >100 km/hr) could occur
714 on an event basis during any time of the year and, thus, could contribute to the variability seen at
715 each field site. Wind is known to generate bias in gauge-based precipitation measurements where
716 unshielded precipitation gauges can catch less than half of the amount of a shielded gauge (Colli
717 et al., 2016). A windscreen design for obtaining rainfall rates – and thus, volumes – to be more
718 reproducible could be considered in future deployments of our developed samplers, similar to
719 recently reported innovations for smaller rainfall rate devices (Kochendorfer et al., 2023). This
720 would, however, increase costs and logistical considerations in deploying the developed devices
721 which currently operate synonymously to deposition systems employed by government monitoring
722 programs. Our siting approach is consistent with these, which often deploy a single sampler
723 without wind protection. Thus, by employing replicates, we are able to ascertain the environmental

724 variability. In addition, collection of replicate samples allows our observations to span a wider
725 physical area, reducing the impact of confounding variables such as wind speed in comparison to
726 a more typical sample size of one for many field collections. Imperfect siting and lack of shielding
727 is necessary where remote field sampling prevents the setup of such infrastructure. As a result, the
728 deployment of triplicate samplers provides researchers with a better opportunity to implement
729 quality control as they can reduce bias in the event of dynamic OF. While the effect of wind is
730 reduced, additional factors can drive variability when the samplers are placed under a forest canopy
731 for TF collection.

732 To demonstrate canopy dynamics impacting interception volumes within the sampling
733 sites, the ratio of throughfall to open fall (TF/OF) volume was compared amongst our total pool
734 of 31 samples. This group of samples encompassed the monthly average TF/OF values for each
735 set of triplicate samplers, at all four sites, from 2015 to 2016. These measurements were then split
736 into two separate populations – samples that have a TF/OF less than one ($n=24$) and those that
737 have a TF/OF greater than one ($n=7$). The samplers were positioned identically between years and
738 no single sampler was reproducibly found in the second population. In the first population, the
739 fraction collected was $56 \pm 21\%$ (ranging from 19 to 88%), likely due to the known processes of
740 canopy and stem interception (Eaton et al., 1973; Howard et al., 2022). For example, in two young
741 balsam fir-white birch mixed forest stands, the amount of precipitation intercepted by the forest
742 canopy, in similar snow-free conditions, was estimated to be $11 \pm 5\%$ (Hadiwijaya et al., 2021).
743 In mature boreal forests, 9% to 55% of rainfall can be intercepted by the canopy (Pomeroy et al.,
744 1999). Relevant to deposition of atmospheric constituents, Pomeroy et al. (1999) also reported that
745 up to 70% of intercepted rainfall may evaporate directly from the canopy, which can leave behind
746 non-volatile rainfall solutes. Wet deposition that undergoes stemflow (SF) proceeds down the
747 branches, stems, and/or trunks of a plant, transferring precipitation and nutrients from the canopy
748 to the soil at the trunk or stem base (Ciruzzi and Loheide, 2021). These known mechanisms of
749 canopy interception ultimately reduce the amount of precipitation reaching the ground as TF, and
750 thus, this explains the smaller volumes found in our samplers compared to the OF measured
751 simultaneously. In contrast, the fractions that ranged from 108% to 424%, averaging 186%,
752 demonstrates a different aspect of the highly dynamic nature of canopies where they can sometimes
753 intercept rainfall like an impermeable surface to act as a funnel, guiding large volumes of
754 precipitation on to the ground, or in this case into the TF samplers (Metzger et al., 2019).



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Figure 4. Average volume collected from replicate automated samplers deployed from June 2015 to October 2016, from north (N) to south (S), at the NL-BELT field sites: **(a)** ER, **(b)** SR, **(c)** PB, and **(d)** GC. The red trace represents open fall, teal for throughfall, and light green for total deposition (the sum of conductive and non-conductive precipitation). The total precipitation volumes depicted for PB, from July 2015 to November 2015, were collected at the nearby HR site in the same watershed since no total deposition measurements were in place at PB during this period. The missing volume for GC in 2015 was estimated from the determined ECCC station linear relationship and is presented as a broken line. The fraction of precipitation collected as open fall or throughfall, compared to the total deposition (right axis), are represented by diamonds of the corresponding color. Error bars represent the standard deviation of three measurements from replicate samples. The axis break spans the winter months when the off-grid automated samplers were stored.

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3.4 Characterizing Chemical Parameters from NL-BELT

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In addition to assessing physical parameters, chemical parameters were also evaluated in this work. Conductivity and pH are measurements commonly made on precipitation samples collected from the field and so incorporating them into our analysis is useful for instrumental validation. Additionally, with increasing recognition of their importance as a proxy for ROC estimation, and in biogeochemical carbon budget closure, DOC flux measurements were used to compare against a limited number of prior reports, each using different sampling or data interpretation strategies. These chemical measurements were also made in an underrepresented

778 part of the world in terms of atmospheric deposition sampling and are useful additions to the
779 overarching study of precipitation chemistry.

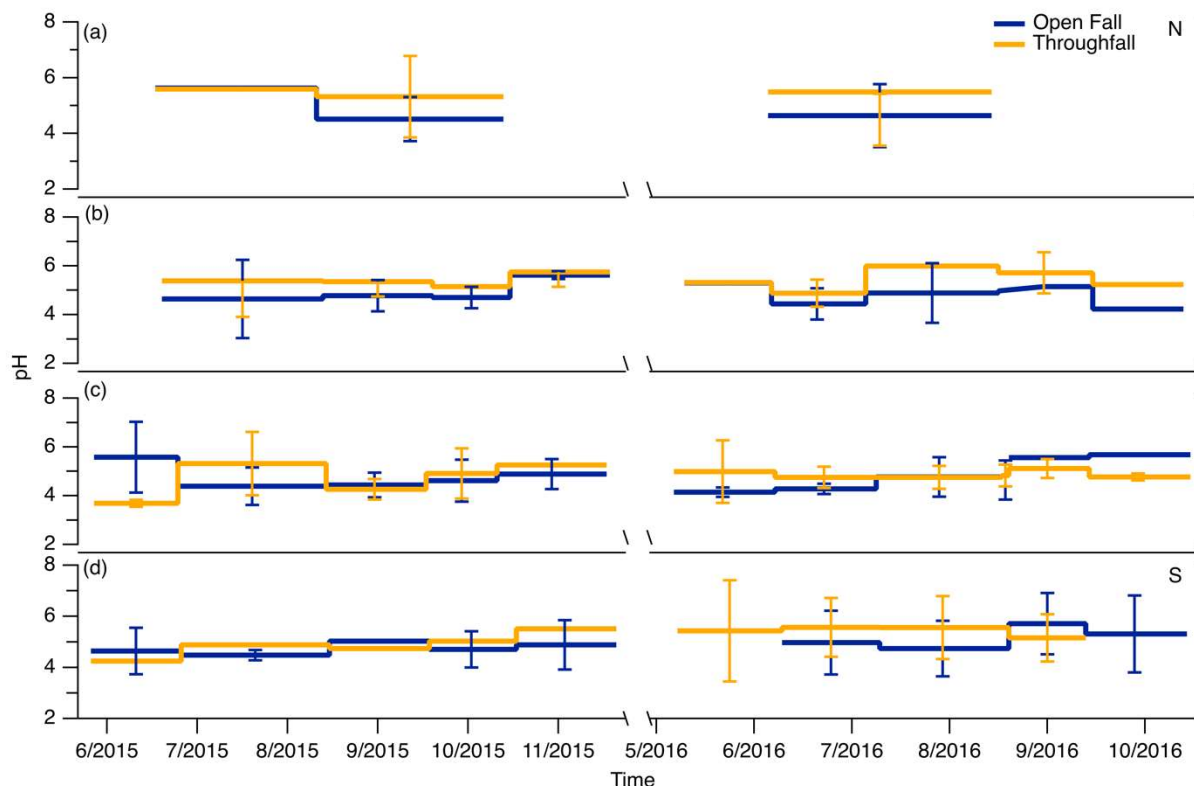
780

781 **3.4.1 Precipitation pH**

782 The deposition of atmospherically persistent pollutants and biogeochemically relevant species
783 to the Earth's surface, or even NO_3^- and SO_4^{2-} historically, can affect the environmental health of
784 soil, air, and water. With the pH range of natural rainwater in equilibrium with atmospheric CO_2
785 expected to be between 5.0 to 5.6, acid rain is defined by values lower than this (Han et al., 2019).
786 Traditionally, the extent of acidity depended on the intercepted atmospheric concentrations of
787 HNO_3 and H_2SO_4 . In any case, monitoring acidity and deposition is especially relevant in remote
788 regions, where major uncertainties and gaps in deposition measurements and global ion
789 concentrations exist (Escarré et al., 1999; Vet et al., 2014). A change in pH can modify the
790 chemical state of many pollutants, altering their transport, bioavailability, and solubility (Guinotte
791 and Fabry, 2008). For example, this can increase exposure and toxicity of metals and nutrients in
792 marine habitats which can go undetected for longer periods in remote areas.

793 Most TF samples were observed to have slightly higher pH than those from OF which had pH
794 values ranging from 4.14 to 5.71 (Figure 5, Table 1). The TF precipitation pH, on average, ranged
795 from 4.74 to 5.99 with rare exceptions falling outside of that range (e.g., July and September 2015
796 PB pH of 3.69 and 4.26, respectively, and the July 2015 GC with pH of 4.12). Excluding these
797 exceptions, there are no major variations observed spatially between the four sites, or temporally
798 between seasons or years (Figure 5). The pH values reported at each of the NL-BELT field sites
799 are comparable to recent OF measurements made at CAPMoN sites in Nova Scotia and
800 Newfoundland and Labrador, where the reported pH of precipitation ranged from 4.44 to 5.19
801 (Houle et al., 2022). The more basic TF overall is expected, as it has been found that up to 90% of
802 H_3O^+ in precipitation can be absorbed by leaves while passing through the canopy (Cappellato et
803 al., 1993). Foliar leaching, the release of ions from leaves, has been commonly reported for base
804 cations such as Mg^{2+} , K^+ , and Ca^{2+} while being minimally observed for other ions such as Cl^- and
805 SO_4^{2-} (Carlson et al., 2003). Mechanisms for foliar leaching include passive cation exchange of
806 H_3O^+ with, for example, cells in the interior of the leaf (Burkhardt and Drechsel, 1997).
807 Additionally, alkaline dust – deposited on the leaves of the canopy, can decrease the acidity of TF
808 precipitation. Such dust can accumulate on leaf surfaces as a result of anthropogenic (i.e., industrial

809 processes) or natural (i.e., wind erosion) sources (Csavina et al., 2012), so that precipitation
 810 passing through the canopy can interact with it (e.g., CaCO_3); thus, neutralizing acidic species and
 811 increasing the TF pH observed in our automated samplers.
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 814 **Figure 5.** Average pH values from replicate samples collected at each NL-BELT field site, from
 815 north (N) to south (S), at (a) ER, (b) SR, (c) PB, and (d) GC, from June 2015 to August 2016.
 816 Open fall collections are represented using the solid blue trace whereas the orange trace is the pH
 817 of the precipitation collected as throughfall under the balsam fir canopy.

818
 819 The pH of the collected precipitation appears to be similar in both TF and OF as a function of
 820 time – despite the potential for foliar leaching and dust dissolution in the canopy. The same
 821 chemical components may be setting the pH, as these measurements do not vary much seasonally,
 822 geographically, or temporally. As pH is a long-studied measurement, its purpose in this work was
 823 to validate the sample quality from our described collection approach, rather than drive any
 824 scientific objective. Nevertheless, while the NL-BELT measurements demonstrate a recovery
 825 compared to rainwater pH in 1980s eastern North America – prior to NO_x and SO_2 regulation (pH
 826 from 4.1 to 5.0; Barrie and Hales, 1984), the present-day pH remains lower than expected for
 827 natural rainwater (~5.6; Boyd, 2020). Keeping in mind the successful environmental policies
 828 limiting SO_2 and NO_x , leading to considerable decreases in atmospheric concentrations of H_2SO_4

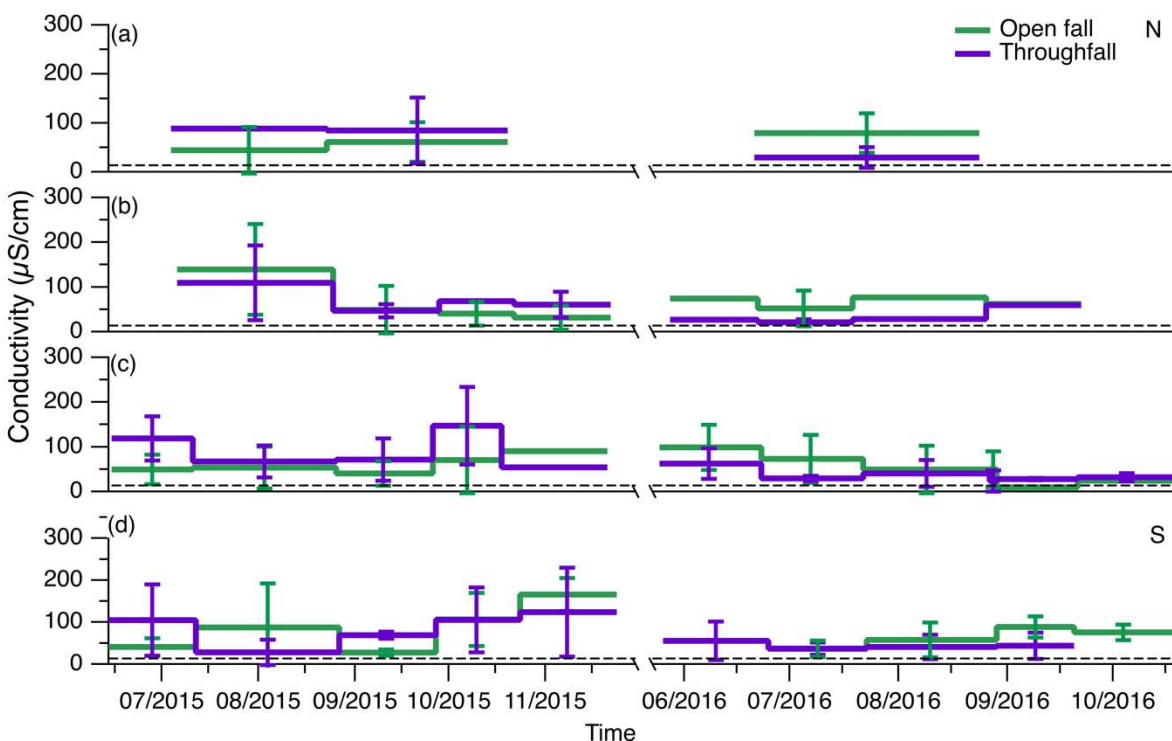
829 and HNO₃, a modern view on the trajectory of continental U.S. cloud water composition and pH
830 has recently been reported (Lawrence et al., 2023). Across the U.S. and eastern Canada,
831 measurements of anion molar charge equivalents have been lower than cations – a potential
832 explanation being an increase in the presence of weak organic acids which commonly have pKa
833 values near 4 (Feng et al., 2021), an outcome we have also observed in aerosol sample chemical
834 composition from Atlantic Canada (Di Lorenzo et al., 2018). With the frequency of acid rain
835 having a pH < 5 decreasing over the past 20 years, these recently reported measurements depict
836 deposition composition shifting away from a ‘linear’ chemical regime dominated by H₃O⁺ and
837 SO₄²⁻ towards a ‘non-linear’ regime designated by low acidity, moderate to high conductivity,
838 potentially weak acid-base buffer systems, and increasing base cation and TOC concentrations
839 (Lawrence et al., 2023). It would seem the evolving chemical contributors to global rainwater pH
840 remain an open line of investigation.

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842 **3.4.2 Precipitation Conductivity**

843 In all the collected OF and TF precipitation samples, across all four NL-BELT sites, the
844 average measured conductivity values ranged from 21 to 166 μS/cm following no apparent
845 seasonal or temporal trend (Figure 6). Additionally, the conductivity in both OF and TF also
846 appear to vary across the field sites - only within the 2016 TF samples does the conductivity appear
847 to increase with decreasing latitude. Yet, with the typical conductivity of surface and drinking
848 waters being between 1 to 1000 μS/cm (Lin et al., 2017), and typically below 200 μS/cm in stream
849 water measurements within the watersheds of each of the NL-BELT sites, our observations are
850 comparable and fall within the expected range. Our field blanks – encompassing a variety of
851 materials and apparatuses, and our cleaning procedures, routinely produced conductivities of 9 ±
852 5 μS/cm. The conductivity of saturated HgCl₂ in water (at 0.1% vol/vol) was 13.6 ± 0.4 μS/cm,
853 which is also comparable to but statistically higher than our field blanks (p = 0.0015; unpaired t
854 test) and less than what was observed for our samples (p < 2 × 10⁻⁶ for each site considered
855 separately and also across all sites; unpaired t-test). Even with this background correction applied,
856 the conductivity values presented here are expected to be similar to or higher than what would
857 typically be found in rainwater (4 to 150 μS/cm; Beverland et al., 1997) as the rain sensor
858 deliberately selects for precipitation containing ionic chemical components with conductivity
859 greater than 1.0 μS/cm, while excluding pure water during atmospheric washout, which would

860 dilute the dissolved solutes in the wet deposition sample and lower the resulting conductivity
 861 values. The overall comparability between our range and those previously reported, where the
 862 lower limit is slightly higher in our dataset, demonstrates that the principle of operation of our
 863 instrument is robust. It decisively collects precipitation with the property of conductance indicating
 864 dissolved ionic solutes of interest to atmospheric chemical processes.
 865



866
 867 **Figure 6.** Average conductivity measured from replicate automated samplers at the NL-BELT
 868 field sites, from north (N) to south (S), at (a) ER, (b) SR, (c) PB, and (d) GC, from June 2015 to
 869 October 2016. The green trace represents open fall samplers whereas the purple trace represents
 870 throughfall samples. The error bar represents the standard deviation between replicate
 871 measurements. The dashed black line represents the upper threshold of conductivity (13.6 $\mu\text{S}/\text{cm}$)
 872 that arises due when an addition of saturated aqueous HgCl_2 is made to microbially sterilize
 873 samples. Note that all samples have conductivities equivalent to or higher than 13.6 $\mu\text{S}/\text{cm}$.
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875 3.4.3. Wet Deposition of Dissolved Organic Carbon (DOC) at NL-BELT

876 The concentration of DOC in OF and TF precipitation, across all four sites, ranged from 3
 877 to 46 mg L^{-1} and 5 to 65 mg L^{-1} with averages of $16 \pm 10 \text{ mg L}^{-1}$ and $22 \pm 12 \text{ mg L}^{-1}$, respectively
 878 (Table 3). Concentrations are influenced by the volume collected and are not useful when
 879 discerning deposition trends and/or mechanisms. The concentrations were converted to elemental
 880 fluxes using the volume of precipitation collected, the area of the funnel and the number of

881 sampling days of each sampling period (Figure 7). The total flux for each sample period was
882 summed and reported as an equivalent annual flux with the following units: $\text{mg C m}^{-2} \text{ a}^{-1}$. Annual
883 fluxes ranged from 600 to 4200 $\text{mg C m}^{-2} \text{ a}^{-1}$ across the study sites for the snow free period (Table
884 S4).

885 The TF DOC fluxes were enhanced compared to the corresponding OF samples as
886 precipitation was intercepted by the forest canopy, with fluxes higher in TF samples by 600, 400,
887 and 400 $\text{mg C m}^{-2} \text{ a}^{-1}$ at GC, SR, and ER, respectively (Table S5). The accumulation of water-
888 soluble organics on forest canopies that increases DOC detected in TF could originate in part from
889 organic carbon-containing compounds aged through oxidation reactions in the atmosphere, which
890 increases their water solubility and propensity for surface interactions. In periods without
891 substantial rain, these oxidized organics deposit effectively to the high surface area of forest
892 canopies, contributing to the elevated DOC measured in TF. Additionally, non-volatile organics
893 left behind from evaporated precipitation intercepted by the canopy could also contribute.
894 Conversely, other mechanisms within the forest could result in enhanced DOC in TF. Recently,
895 Cha et al. (2023) utilized a mass balance approach to determine whether DOC deposition is driven
896 by canopy leaching (i.e., soluble tree resin, leaf exudates, internal tissues and microbes) or
897 dissolution of dry deposited gases and $\text{PM}_{2.5}$ on plant foliage into rainwater. It was found that
898 canopy leaching is the major contributor to TF DOC, accounting for ~83% of throughfall DOC.
899 Whereas, $\text{PM}_{2.5}$ and rainwater only accounted for ~3 and 14%, respectively, while dry deposited
900 gases were not considered. This suggests that internal cycling of DOC within the forest could be
901 an important source of DOC to the throughfall soil interface (Cha et al., 2023). It is possible that a
902 similar mechanism may be responsible for the elevated levels of DOC in TF at the NL-BELT sites,
903 but we cannot explicitly distinguish between internal cycling versus external deposition in the
904 current study.

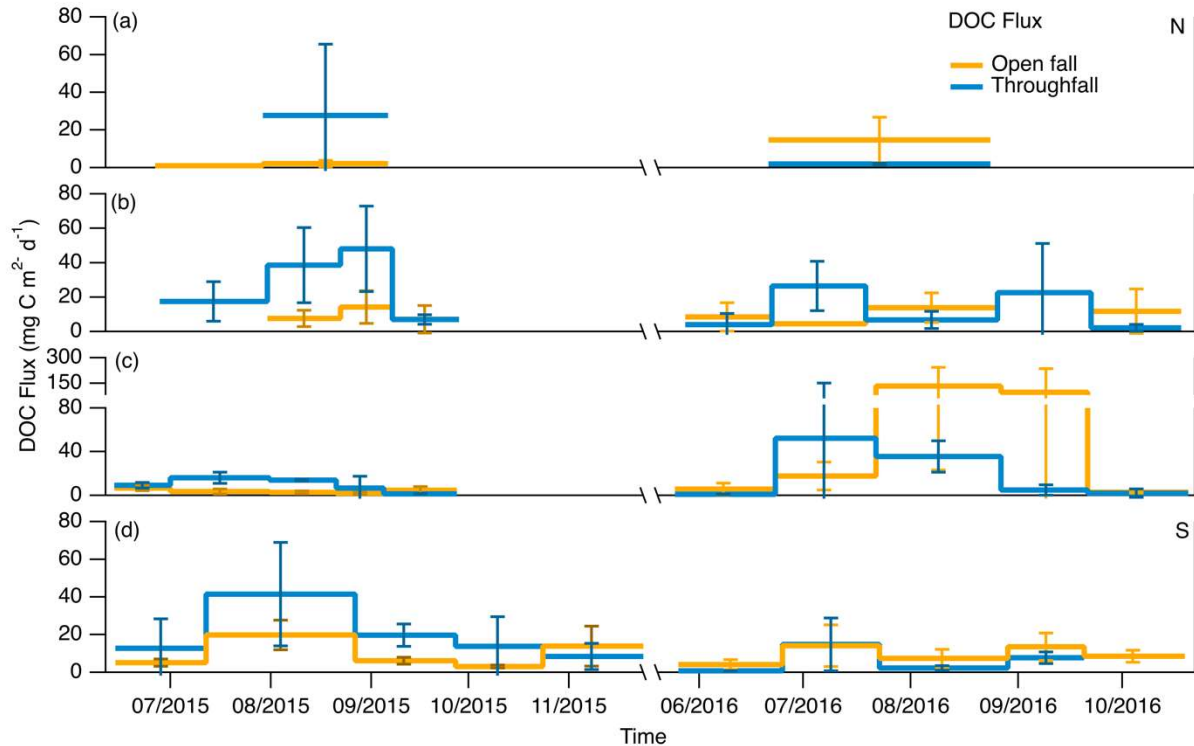
905 A notable exception was observed at PB, where the DOC fluxes in the open fall sample
906 were enhanced up to 1800 $\text{mg C m}^{-2} \text{ a}^{-1}$ when compared to the TF in 2016. This may be attributed
907 to a difference in forest type within this NL-BELT region being black spruce (*Picea mariana*)
908 instead of balsam fir (Bowering et al., 2023). Some studies have suggested that forest type could be
909 a major factor affecting DOC variability (Arisci et al., 2012; Sleutel et al., 2009). Specific
910 differences in canopy height, leaf area index, canopy structure and the shape of leaves and needles
911 could drive DOC differences between forest types (Smith, 1981; Erisman and Draaijers, 2003;

912 Sleutel et al., 2009). The elevated levels in OF samples relative to TF within PB are consistent
913 with idea of uptake and/or leaching of canopy DOC in the internal cycling of DOC, while the
914 enhanced TF at the rest of the sites is more difficult to observational constrain the participating
915 processes.

916 Episodic events, such as polluted air masses from wildfires could also result in elevated
917 deposition of DOC. It is estimated that $\sim 116 - 385 \text{ Tg C a}^{-1}$ is produced globally due to the
918 incomplete combustion of biomass during landscape fires (Santín et al., 2016; Coward et al., 2022).
919 Several studies have associated enhanced DOC levels with wildfires (Gao et al., 2003; Moore,
920 2003; Wonaschütz et al., 2011; Myers-Pigg et al., 2015). More recently, Coward et al. (2022)
921 measured DOC in Pacific surface waters along the California coastline and observed 100 to 400
922 % increases in DOC concentration, when compared to pre-wildfire conditions. It is possible that a
923 similar biomass burning plume that underwent atmospheric washout, could be responsible for the
924 enhancement in the observed DOC at NL-BELT, overlaid on a background more typical of
925 seasonal oxidation of biogenic DOC. This also coincides with the seasonal variability observed in
926 OF samples from the same summer where elevated levels of DOC were measured. For instance,
927 the DOC deposition at PB for August 2016 was 4800 mg C m^{-2} , whereas the total deposition for
928 the same year was $7800 \text{ mg C m}^{-2} \text{ a}^{-1}$. This single period accounts for 62% of the total DOC
929 deposition at this site. This underscores the pivotal role that episodic transport may play in
930 influencing the dynamics of DOC deposition, particularly with a warming future where wildfires
931 are more prevalent.

932 The deposition trend observed in the current study also highlights the complexity of the
933 varied drivers of atmospheric ROC, in which some months have more DOC in TF versus OF and
934 occasionally the opposite is observed. Generally, we observed similar fluxes in both samples –
935 suggesting that the amount of deposited carbon is comparable. Although the volume of
936 precipitation captured in TF samplers are generally lower when compared to the corresponding
937 OF samplers, the deposition flux of DOC is greater in TF samplers. With DOC enhanced in TF
938 samples, the values reported here could be an underestimation of the amount of carbon reaching
939 the forest floor during precipitation events due to competing processes within the canopy. One
940 such process is stemflow (SF), where a fraction of precipitation intercepted by the forest canopy
941 is funneled over the bark of the tree surface to the base of the tree stem (Oka et al., 2021). Although,
942 SF was not measured in the current study, several studies have demonstrated that DOC

943 concentrations are enhanced in SF when compared to the corresponding TF and bulk precipitation
 944 samples (Stubbins et al., 2017; Van Stan and Stubbins, 2018; Ryan et al., 2021). Additionally, we
 945 cannot rule out that the chemical speciation differs between OF, TF, and SF even if the DOC values
 946 are similar, but such insights require more selective instrumentation for chemical analysis (e.g.,
 947 high resolution mass spectrometry).
 948



949 **Figure 7.** Average DOC fluxes ($\text{mg C m}^{-2} \text{d}^{-1}$) in replicate samples collected at the NL-BELT field sites, from north (N) to south (S), at (a) ER, (b) SR, (c) PB, and (d) GC, from June 2015 to August
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957 The ability to accurately determine DOC in OF and TF precipitation demonstrates the
 958 capability of the automated deposition samplers. To validate our measurements, we compared our
 959 observed fluxes to other studies in different forest types. Mean annual DOC fluxes were generally
 960 similar to those reported in some other boreal forests (Table 3). These include Finland, with work
 961 in stands that consisted mainly of Scots pine (*Pinus sylvestris L.*) (mean OF 2.32; TF, 4.35 $\text{g C m}^{-2} \text{a}^{-1}$;
 962 Pumpanen et al., 2014), as well as in Mont St. Hilarie, Québec (mean OF 0.49; TF 2.05 $\text{g C m}^{-2} \text{a}^{-1}$;
 963 Dalva and Moore, 1991), which also consisted of a variety of tree species such as yellow

964 birch (*Betula allenghanien*), red maple (*Acer rubrum*), and sugar maple (*Acer saccharum*).
965 Conversely, the annual fluxes were orders of magnitude lower than measurements made at the
966 University of Georgia (23 to 48 g C m⁻² a⁻¹) which has a subtropic climate consisting mainly of
967 southern live oak (*Quercus virginiana Mill.*) and eastern red cedar (*Juniperus virginiana L.*)
968 occasionally hosting dense epiphytes (Van Stan et al., 2017). This highlights the potential
969 variability to expect when measuring DOC in different forest systems, as the annual DOC fluxes
970 vary depending on factors such as climate, tree species composition, and environmental conditions.

971 These results underscore the pivotal role the off-grid custom-built automated deposition
972 samplers can play in advancing scientific research, particularly in precipitation monitoring and
973 analysis. The automated system enabled long term continuous sample collection in remote
974 locations, which was previously challenging to attain due to the need for frequent human
975 intervention and resources required to regularly access these experimental forest stands. These
976 samplers also allowed us to compare DOC through replicate measurements in TF and OF samples
977 which sheds light on the potentially different DOC deposition chemistries within the NL-BELT
978 region. The automated system better maintains the integrity of DOC in the samples. This was
979 achieved by following standard procedures for biogeochemical sample preservation (i.e., adding
980 HgCl₂) (Argentino et al., 2023), employing a rigorous cleaning procedure, and preventative design
981 against the intrusion of forest litter which could result in a positive bias for DOC in the collected
982 precipitation. The use of replicates also results in more robust scientific conclusions and broader
983 applicability of the results, and they can be obtained for a fraction of the cost of a commercial
984 equivalent, highlighting the contribution these automated systems are capable of when applied to
985 current precipitation monitoring. As a result, these samplers show promise in the quantification of
986 biogeochemical and anthropogenic chemical species of interest, which will be visited in future
987 manuscripts drawing from the samples presented in this dataset, and others since obtained, but are
988 beyond the scope of this manuscript in demonstrating the performance of this new instrumentation.

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996 **Table 3.** Concentrations (mg C L^{-1}) and annual fluxes ($\text{g C m}^{-2} \text{a}^{-1}$) of DOC in precipitation (P),
 997 open fall (OF), throughfall (TF), and stemflow (SF) collected at forested sites. Where volumes are
 998 not available for other studies, fluxes are not possible to calculate. The values reported in the
 999 current study are the estimated DOC flux for the wet deposition sampling period (~June through
 1000 October) for each year and therefore represents the lower limit of DOC deposition, as the dataset
 1001 excludes snow.

Site	Type	Mean Concentration (mg C L^{-1})	Annual Flux ($\text{g C m}^{-2} \text{a}^{-1}$)	References
Grand Codroy, NL, Canada (2015 to 2016)	OF	12.83	1.56	This study
	TF	23.40	2.20	
Pynn's Brook, NL, Canada (2015 to 2016)	OF	19.98	4.21	
	TF	21.24	2.44	
Salmon River, NL, Canada (2015 to 2016)	OF	16.14	1.33	
	TF	21.00	2.65	
Eagle River, NL, Canada (2015 to 2016)	OF	11.59	0.53	
	TF	28.26	0.86	
Mont St. Hilaire, QC, Canada (1987)	P	2.00	0.49	Dalva and Moore, 1991
	TF	12.13	2.05	
	SF	40.10	0.10	
Northern China (2007 to 2008)	P	2.4 to 3.9	1.4 to 2.7	Pan et al., 2010
Coulissenhieb, Northeast Bavaria (1995 to 1997)	P	2.70	-	Michalzik and Matzner, 1999
	TF	15.20	-	
Hobcaw Barony, South Carolina, USA (2014 to 2015)	P	1.20	-	Chen et al., 2019
	Pine TF	26.00	-	
	Oak TF	38.8	-	
University of Georgia, USA 2015 to 2016	TF Epiphyte Oak	17	23**	Van Stan et al., 2017
	TF Bare Cedar	20	32**	
	TF Epiphyte Cedar	54	48**	
SMEARII Site, Southern Finland (1998 to 2012)	P	3.24	2.32	Pumpanen et al., 2014
	TF	10.10	4.35	

** Estimated DOC yield for 2016 ($\text{g C m}^{-2} \text{a}^{-1}$) where sampled storms values (g C event^{-1}) were scaled to an annual deposition value using meteorological data and a linear rainfall-DOC yield relationship.

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1006 **4.0 Conclusions and Future Directions**

1007 This paper presents a cost-effective automated deposition sampler for continuous
1008 collection of precipitation. An open-source procedure and schematics for building these samplers
1009 is provided alongside the rationale for selecting the materials in the current study to target analytes
1010 of scientific interest in wet deposition samples. These low-power systems are demonstrated in
1011 being capable of continuous off-grid use for sample collection over two years at the NL-BELT
1012 experimental sites, with replacement of battery power packs monthly or bimonthly, with on-grid
1013 performance also provided for comparison. The resulting systems enhance the accessibility of
1014 automated wet deposition samplers to scientists globally and this work highlights their robust
1015 performance in collecting and preserving rainwater conductivity and pH, alongside providing
1016 measurements of DOC from this understudied region that builds a broader picture of the
1017 atmosphere-surface exchange of this biogeochemical pool across the NL-BELT. Comparability
1018 and complementarity of our results to well-established and current measurements of interest like
1019 DOC, demonstrate their efficacy and potential application to the study of processes such as
1020 canopy-precipitation interactions through the collection of open fall and throughfall replicates. The
1021 capacity to autonomously collect wet deposition, in addition to traditional bulk deposition samples
1022 can shed light on competing wet and dry deposition processes. Should on-grid capacity suit
1023 scientific objectives, these samplers are anticipated to be possible for use year-round when paired
1024 with more power-intensive strategies to facilitate solid to liquid phase transfer for detected and
1025 collected precipitation in the winter.

1026 For the broader deposition-motivated community, the instrument design also allows for
1027 easy cost-effective modification of the number of replicate samplers, the material composition of
1028 all surfaces the aqueous samples interact with, as well as preservation strategies - depending on
1029 the analyte of interest. For example, the lack of organic nitrogen measurements within universally
1030 established sampling and measurement procedures serves as a general example of the substantial
1031 knowledge gaps that may result when translating limited data sets to the wider global picture. This
1032 includes incomplete speciation and quantification across precipitation, aerosol, and gas phases.
1033 Monitoring systems that support U.S. deposition assessments (e.g., the NADP) only characterize
1034 the inorganic fraction of wet deposition. Additionally, modern emerging issues that require the
1035 continuation of existing deposition measurements or expansion of observation programs revolve
1036 around identifying and quantifying compound classes of concern, such as persistent organic

1037 pollutants. As reported in the literature, the deposition of these types of pollutants (e.g.,
1038 polychlorinated biphenyls, polycyclic aromatic hydrocarbons, etc.) can be monitored using
1039 suitable collectors made of amber-coloured glass or stainless steel (Fingler et al., 1994; Amodio et
1040 al., 2014) - modifications which can be applied to the sample design detailed here. The samples
1041 collected in this work from this new instrumentation, specifically, are expected to be used further
1042 in several upcoming complementary and novel environmental monitoring studies. Not only will
1043 this future work extend our biogeochemical analysis, but it will also assist in our studying of the
1044 transport of other anthropogenic pollutants of emerging interest which are beyond the scope of
1045 describing this new platform.

1046

1047 *Data availability.* The data are available from the corresponding author (TV) on request.

1048

1049 *Author contributions.* AC, DP, and ML performed the data analysis. AC and DP wrote the
1050 manuscript with contributions from all authors. Sampler design and construction were led by TV,
1051 with assistance from BP and RH for initial prototypes, DP and ML for the revised iteration, and
1052 AC for the final modular control boards. Sample collection and associated characterization
1053 measurements were performed by BP and TV. Conceptualization and conduct of the sampling
1054 experiments were made by TV, CY, KE, and SZ. All authors were involved in examining and
1055 reviewing the results. All authors were involved in editing the paper.

1056

1057 *Competing interests.* The contact author has declared that none of the authors has any competing
1058 interests.

1059

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1082

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