

1 **Title**

2 PolyWAG: Autonomous filtered water sampling for eDNA

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13 **Abstract**

14 Environmental DNA (eDNA) is an ideal way of researching aquatic environments and determining what
15 species are present in an area the biodiversity of an area, and if any invasive or endangered species are
16 present. Traditional sampling of eDNA consists of manually filtering water, which is labor and cost-intensive
17 for remote locations. Furthermore, commercialized solutions are either expensive or require a field operator to
18 function. We have built an eDNA sampler capable of autonomous multi-sampling for a greatly reduced price
19 compared to existing technologies. Our PolyWAG eDNA sampler system is a water sampling device that
20 collects DNA samples via 47mm filter and provides a non-invasive, safe and autonomous means of eDNA
21 collection. The sampler can hold 24 filters and is designed to be easily replaced and reusable. A browser
22 application is used for real-time monitoring, scheduling tasks, and data logging for time, pressure, flow, and
23 filtered volume. Additionally, the sampler design is openly published, modular and is constantly being tested
24 to help us optimize our software and hardware to give us the best results. The 13-step sampling sequence
25 helps reduce cross contamination significantly. Our machine can be deployed for an extended period. It is
26 completely autonomous and costs around \$3800.

27 **Keywords**

28 Environmental DNA * Sampling * Arduino * Data Logging

Hardware name	<i>PolyWAG</i>
Subject area	<i>Environmental, planetary and agricultural sciences</i>
Hardware type	<i>Field measurements and sensors</i>
Closest commercial analog	<i>Dartmouth Ocean Technologies' eDNA Sampler</i>
Open source license	CERN Open Hardware Licence Version 2 - Strongly Reciprocal GNU AFFERO GENERAL PUBLIC LICENSE Version 3
Cost of hardware	<i>\$3800 (Cost of just components)</i> <i>\$6000 (Cost with labor included)</i>
Source file repository	<i>If you've uploaded your source files to an approved repository (OSF, Mendeley Data or Zenodo) write the DOI URL here. For example: http://doi.org/10.17605/OSF.IO/WGK7Q</i>

Table 1:

1. Hardware in context

Environmental DNA (eDNA) is DNA derived from mucus, feces, gametes, and carcasses [1]. Many things can be learned once this DNA is put through sequencing. eDNA can be used to determine what species are present in an area, the biodiversity of an area, and if any invasive or endangered species are present [2]. eDNA sampling provides scientists and researchers a non-invasive, rapid, cost-effective and sensitive way to detect and quantify species in many environments.

Traditional sampling of environmental DNA consists of manually filtering water, often requiring one or more researchers to be on location for days or weeks [3]. The filtration process varies depending on the researcher, but it is common to pull a sample of water with a bottle and pour that water into a funnel containing a filter. This can be connected to a vacuum pump to expedite the filtering process. After the sampling process is completed, the filters need to be preserved and the setup cleaned to avoid cross contamination [3]. This process is labor intensive, cost intensive, and can be dangerous, especially for remote locations. While commercialized solutions to this problem exist, they either still require an operator to be on location or are very expensive. Smithroot’s commercial solution offers a simplified process with additional data collection such as GPS location for a fair price, around \$8,000 [4]. A disadvantage of this solution is that it is not fully autonomous, still requiring an operator to be on location to use the device [4]. An alternative is the DOT Sampler which is a fully autonomous solution that is capable of multiple samples (20+ samples) and is also submersible but comes at a cost of ~\$55,000 [5].

The solution designed by the OPEnS Lab is the middle ground of these two solutions. While it is not submersible (limiting its potential sampling environments), it is capable of autonomous, multi-sample operations for extended periods of time (approximately one month) for the cost of \$6,000. The two core priorities for our design are its autonomous function and the cross-contamination. The autonomous function of the sampler is important for a handful of reasons. An autonomous system requires less researcher hours spent in the field. This has cost benefits from the reduced hours worked and safety benefits when sampling in hazardous environments.

2. Hardware description

The eDNA sampler we have developed is an autonomous multi-sampling device that collects eDNA samples from water via 47mm filter holders and provides a non-invasive, safe, and autonomous means of DNA collection. The sampler can hold 24 filter housing which are designed to be easily replaced and reusable. The sampler is

60 controlled by a custom logic board with an Adafruit M0 Feather Wi-Fi microcontroller loaded with a webserver
61 to act as the interface for the sampler's operations. This webserver hosts a browser application which is used
62 for real-time monitoring, scheduling tasks, and data logging for time, pressure, temperature, flow, and sample
63 volume. The data is located stored onto an SD Card for later data analysis.

64

65 The basic function of the sampler can be split into five main sections: Hydraulics, Sampling Procedure,
66 Utilities, Electronics, and User Interface. The [Hydraulics](#) section describes how the sampler is physically
67 arranged, the different devices connected to the system, and provides a general idea on how liquids flow through
68 the system. This section is supported by the [Sampling Procedure](#) Section which covers what the PolyWAG
69 sampler is doing during each stage of the process. The Sampling Procedure Section refers to the devices
70 described in the Hydraulics section in order to describe what is happening in each state as well as the purpose
71 of that state. The [Utilities](#) section is similar to the Sampling Procedure section, but covers the four main utility
72 functions of the sampler. These are functions that are used by an operator during cleaning or setup of the
73 sampler. The [Electronics](#) section gives a brief overview of how the custom control board functions. Finally, the
74 [User Interface](#) sections gives a brief overview on what the can be found and one within in the UI. More specific
75 details on the User Interface can be found in the [Operation Instructions](#) section

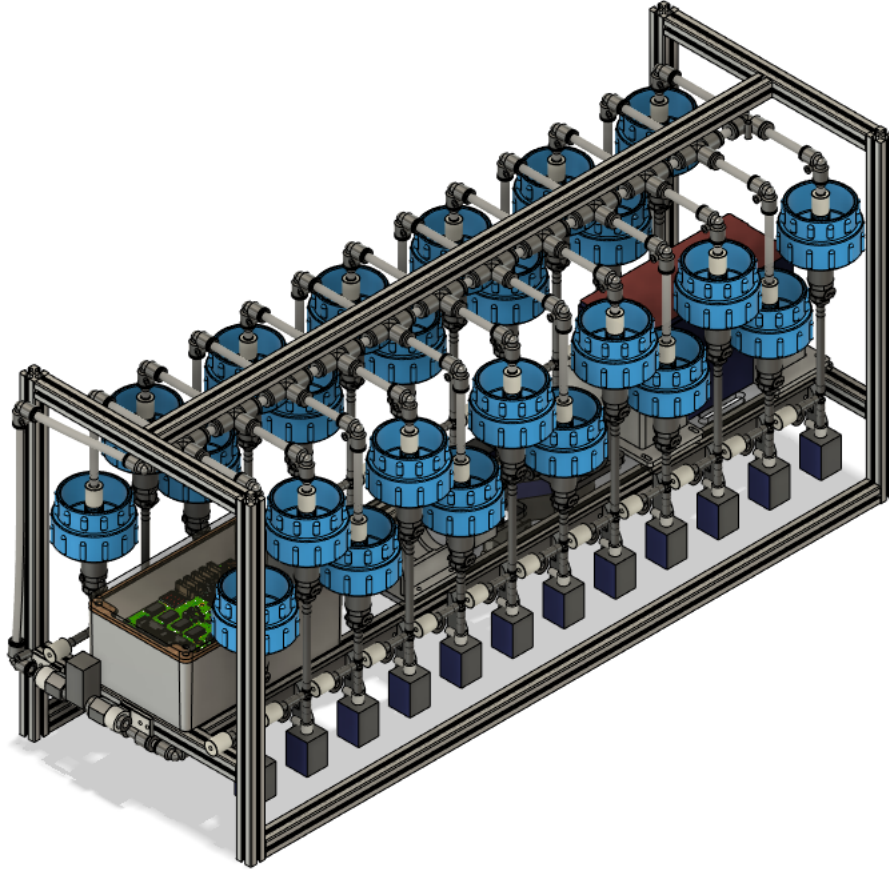


Figure 1: CAD Image of the complete sampler assembly

2.1 Hydraulics

The hydraulics of the sampler can be roughly split into the following sections:

- The Pump and Inputs
- The Lower Hydraulics
- The Filter Housings
- The Upper Hydraulics and Outputs

2.1.1 The Pump and Inputs

There are three inputs into the sampler: one for air, one for preservative, and one for water. The preservative input is connected to a hydration bladder where the preservative of choice can be stored. The sample water input has a prefilter at the front end of the tube to prevent debris from entering the sampler. Three valves are used to control the flow from these inputs with the air and preservative being regulated by solenoid valves and

87 the water being controlled by a ball valve. These three valves connect into a single tube connected to the input
 88 of the peristaltic pump. The pump is capable of 400mL/min of flow under ideal conditions. The output of the
 89 pump connects directly into the Lower Hydraulic Rail.

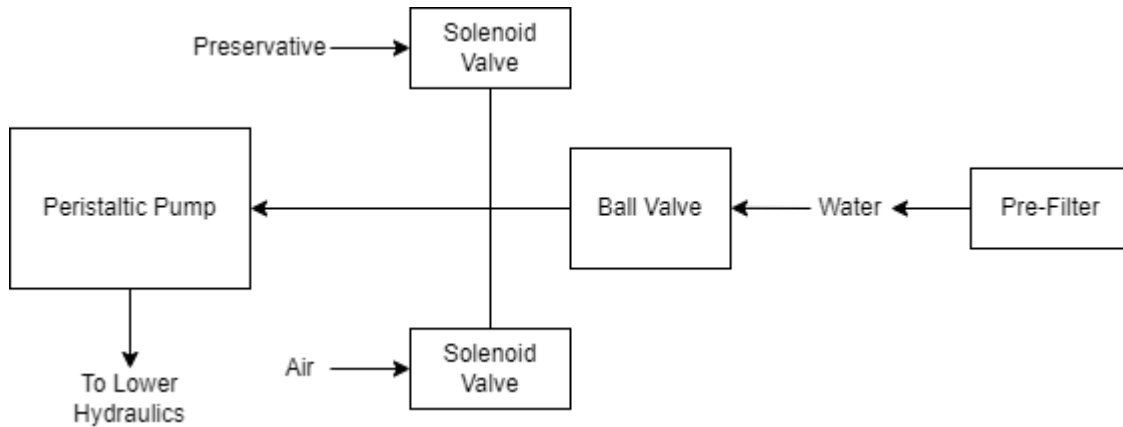


Figure 2: Pump and Input Hydraulics

90 2.1.2 The Lower Hydraulics

91 The Lower Hydraulic Rail consists of 24 solenoid valves connected parallel to each other and controls which
 92 filter liquid flows through. These filter valves are split into two sets, one on each side of the sampler. In between
 93 these two sets of valves is a M32JM-000105-100PG pressure and temperature sensor. The temperature is logged
 94 for later use and the pressure is used for monitoring, stopping an operation if the pressure exceeds a certain
 95 margin. At the end of the Lower Hydraulic Rail is another solenoid valve which allows for the lower hydraulics
 96 to be purged of their current contents when necessary.

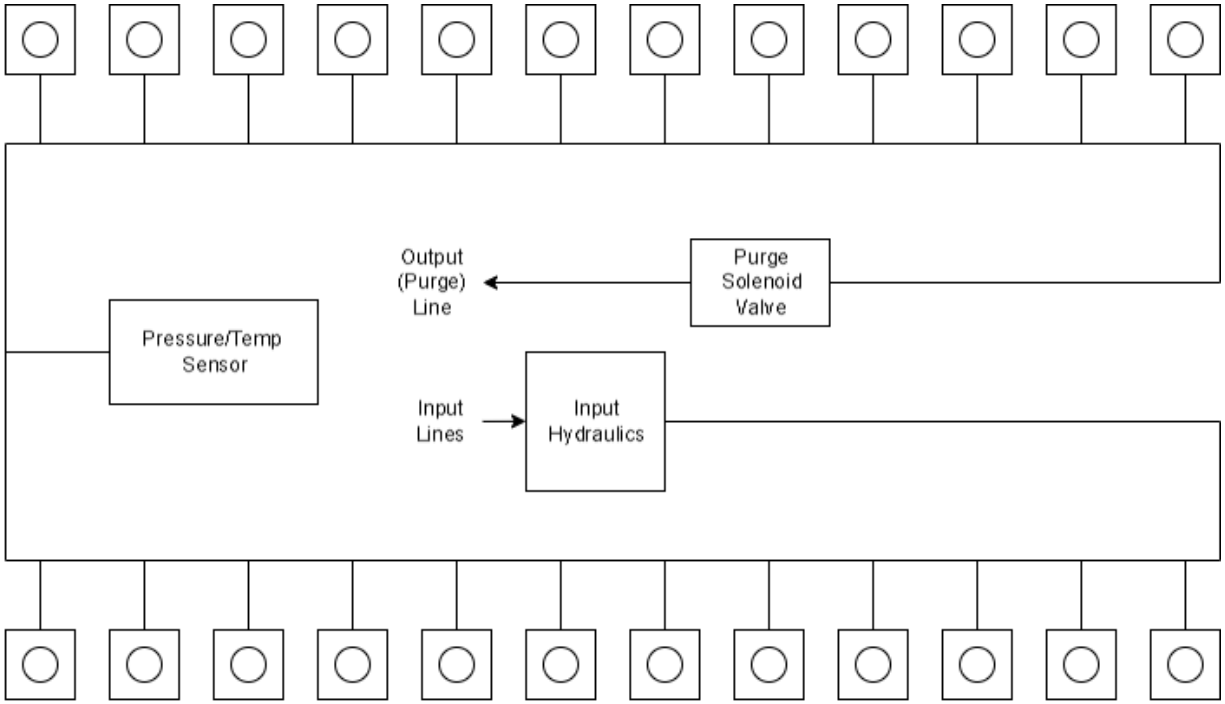


Figure 3: Lower Hydraulic System

97 2.1.3 The Filter Housings

98 Downstream of each filter solenoid valve there is a tee connection that goes to a one-way check valve and a
 99 modified Advantec filter. The one-way check valve allows air into the solenoid valve that opens when the pump
 100 runs backwards. The Advantec filter is modified with a CPC quick disconnect and a one-way check valve. The
 101 one-way check valve is connected to the Upper Hydraulics and is used to prevent liquid from going backwards
 102 through the filter. The Upper Hydraulics simply connects the output of all the filters to one central line that
 103 goes through a flow meter and out of the sampler.

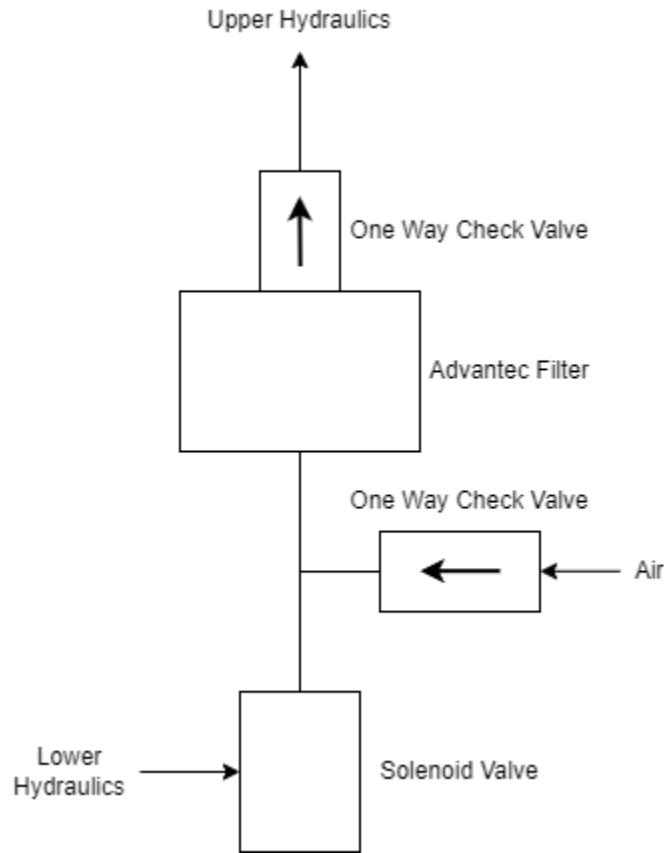


Figure 4: Filter Hydraulics

2.1.4 The Upper Hydraulics and Outputs

The output of the filters connect into a single output hydraulic line. This line is the main output of the filters, any water that goes through the filters will end up going through this line. This allows for a single flow meter to be added that can measure the flow going through any of the filters. This flow meter is crucial as this is how the sampler knows how much water has gone through a filter. After the flow meter the output line goes outside of the sampler and lets liquids flow back into the environment.

2.2 Sampling Procedure

Having worked on multiple iterations of the sampler, we have decided to go with a 13-step sampling sequence that helps reduce cross contamination significantly. This sequence can be split into 9 unique steps: Idle, Prefilter Clear, Flush, Offshoot Clean, De-pressure, Sample, Preservative Flush, Preservative, Air Flush, and End.

1. Idle
2. Prefilter Clear
3. Flush

- 117 4. Offshoot Clean
- 118 5. Flush
- 119 6. Sample
- 120 7. De-pressure
- 121 8. Preservative Flush
- 122 9. Preservative
- 123 10. Flush
- 124 11. Offshoot Clean
- 125 12. Air Flush
- 126 13. End

127 **2.2.1 The Idle State**

128 Idle is the default state of the sampler. The sampler waits for a signal from the RTC to move to the first/next
129 state of the Sampling Sequence. If the sampler is not in sleep mode, this is when a client would interact with
130 the UI to do a handful of tasks such as setting up a Sampling Schedule or using the other task utilities. If the
131 sampler is in sleep mode, then only the RTC and supporting circuits are powered. This means there is no way
132 to interact with the sampler without exiting sleep mode.

133 **2.2.2 The Prefilter Clear**

134 Once the RTC sends the signal to start a sample procedure, the sampler enters the Prefilter Clear (PC) state.
135 The purge and input ball valve are opened, and the pump is run in the backwards direction. This will allow for
136 air to flow from the purge and out the input line. This is used to clear the prefilter of anything that might be
137 clogging it, such as accumulated debris. This state runs for X seconds, before moving onto the next state.

138 **2.2.3 The Flush State**

139 The Flush state prepares the lower hydraulics before the next state. The Flush state starts with the purge
140 valve and the ball valving opening, then the motor starts to run in the forward direction. This fills the lower
141 hydraulics with sample liquid and clears out/dilutes and liquid that remained from previous sample. The Flush

142 state runs for the time specified when the Sampling Schedule is created. We recommend a Flush time of 6
143 minutes.

144 **2.2.4 The Offshoot Clean**

145 The OC state closes the purge valve and opens the filter valve for the filter which is about to be used. The
146 pump runs backwards for a few seconds. This clears anything that might be in the tube between the valve and
147 the filter (what we refer to as the offshoot). The Flush state is run one more time before moving to the Sample
148 state.

149 **2.2.5 The Sample State**

150 In the Sample state, the system pushes the sample water through the filter. This is done by opening the filter
151 solenoid valve and Ball Valve and running the pump in the forward direction. The system moves to the next
152 state when the target Sample Volume is reached. This volume is measured by a Flow Meter on the filter output
153 line. There is an additional condition that will end the Sample state, Sample Time. This time cutoff was added
154 since the filter clogs, rapidly decreasing the flow rate during the sample process. To prevent the sample state
155 running for too long, the time limit was implemented. Both conditions are set during task scheduling. Since
156 the pressure greatly increases due to the clogged filter, the de-pressure state is used to reduce the pressure in
157 the lower hydraulics to ensure that the valves can operate consistently.

158 **2.2.6 The Preservative Flush**

159 The Preservative Flush state is nearly identical to the Flush state except the Preservative input valve is used
160 instead of the ball valve. The lower hydraulics are saturated with preservative, preventing additional sample
161 water that may have been stored in the lower hydraulics from going through the filter. If this water was allowed
162 through the filter, then the Sample Volume would be inaccurate by the end of the sequence.

163 **2.2.7 The Preservative State**

164 The Preservative state is like the Sample state except preservative is the input fluid instead of sample water.
165 This state runs for a time specified by the user during scheduling.

166 **2.2.8 The Air Flush**

167 Before the Air Flush, another sequence of Flush and Offshoot Clean states are run to purge the leftover
168 preservative in the lower hydraulics. After these two states, an Air Flush (AF) state is run which is identical

169 to the Flush and PF states but uses the air valve as the input instead of the other two inputs. This ensures
170 that any liquid that is in the lower hydraulics is purged.

171 2.2.9 The End State

172 In the End state, the system sets an RTC alarm for the time of the next sample. The system then moves into
173 Idle and if the system was in sleep mode, then the system will go into its low power state.

174 2.3 Utilities

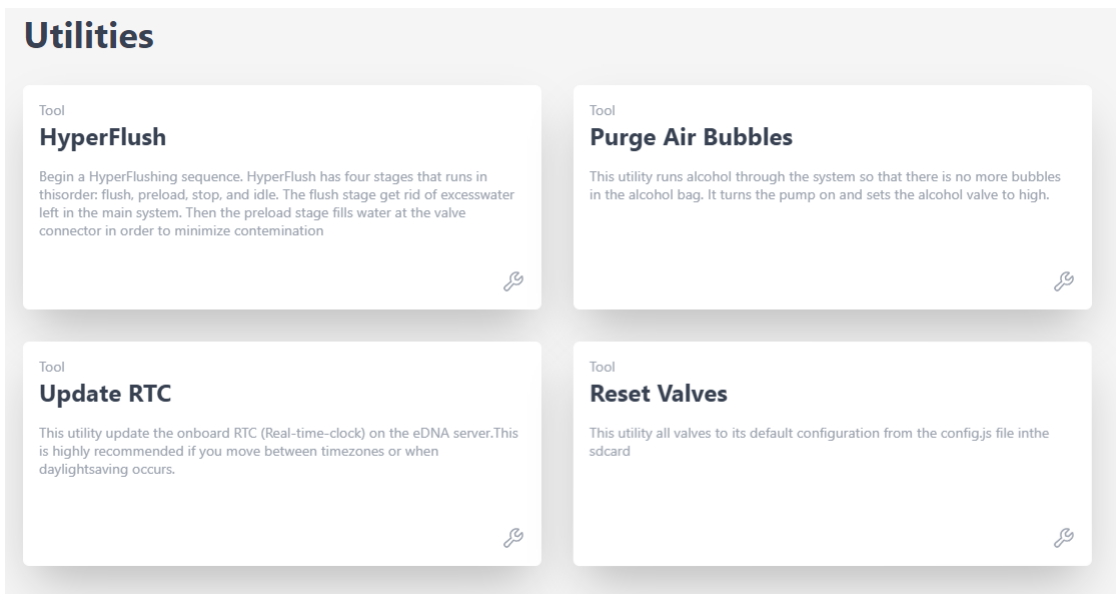


Figure 5: The utilities page in the PolyWAG sampler user interface

175 The HyperFlush utility runs water through every filter sequentially for a few seconds per filter. This is mainly
176 used for cleaning out the system after a sample task (i.e., a set of 24 samples) to prevent any unwanted cross
177 contamination. This utility can also be used to test the basic functionality of the sampler, as nearly every
178 component is activated during this sequence.

179

180 The Preservative Air Purge (PAP) utility turns the pump on and opens the alcohol valve for 10 seconds. This
181 runs some alcohol through the system and removes air bubbles from the alcohol bag. Often it helps to use this
182 utility multiple times and to tilt the Preservative Bladder so that the air is near the port.

183

184 The Update RTC utility is needed to make sure that the time on the sampler matches your local time, so
185 scheduling a task will remain accurate. Whenever the system is fully depowered (ie the battery is removed), or

186 when new code is uploaded to the microcontroller, the RTC will need to be updated. It is also recommended
187 that the RTC is updated when there is a daylight-saving change, or when you move between time zones.

188

189 The Reset Valves Utility is used when valves have been sampled that you want to be sampled again. This is
190 required since the system ‘locks’ the filter valves when they have been used in a sample, this prevents samples
191 from being corrupted accidentally. The code does not let you sample a valve multiple times without being reset
192 to prevent messing up a sample. It is important to note that this utility will reset all valves, not a specific one.

193 2.4 Electronics

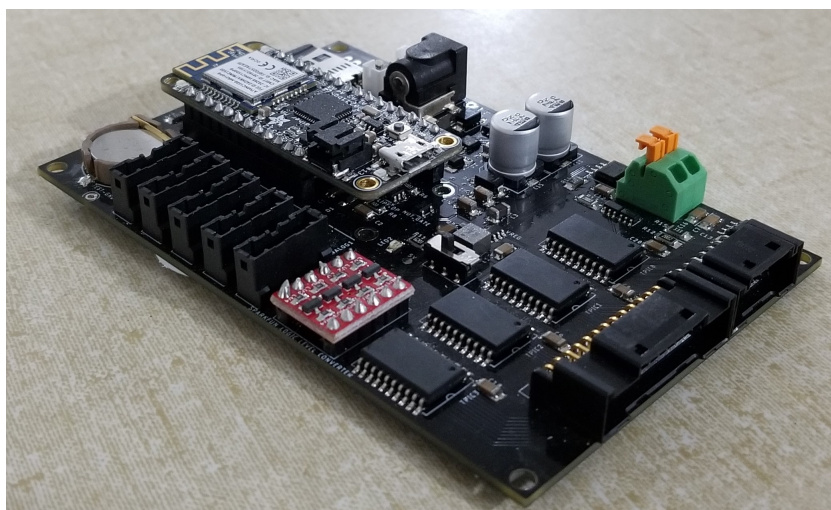


Figure 6: The main electronics control board for the PolyWAG sampler

194 The PolyWAG Sampler is designed with a custom electronics control board that can be split into 8-10 blocks
195 with an Adafruit Feather M0 at its core. These blocks consist of the microcontroller/Wifi Block, Power, RTC,
196 and sleep control blocks, and the output blocks consisting of the Shift Register, Pump, and Ball-Valve H-Bridge
197 Blocks.

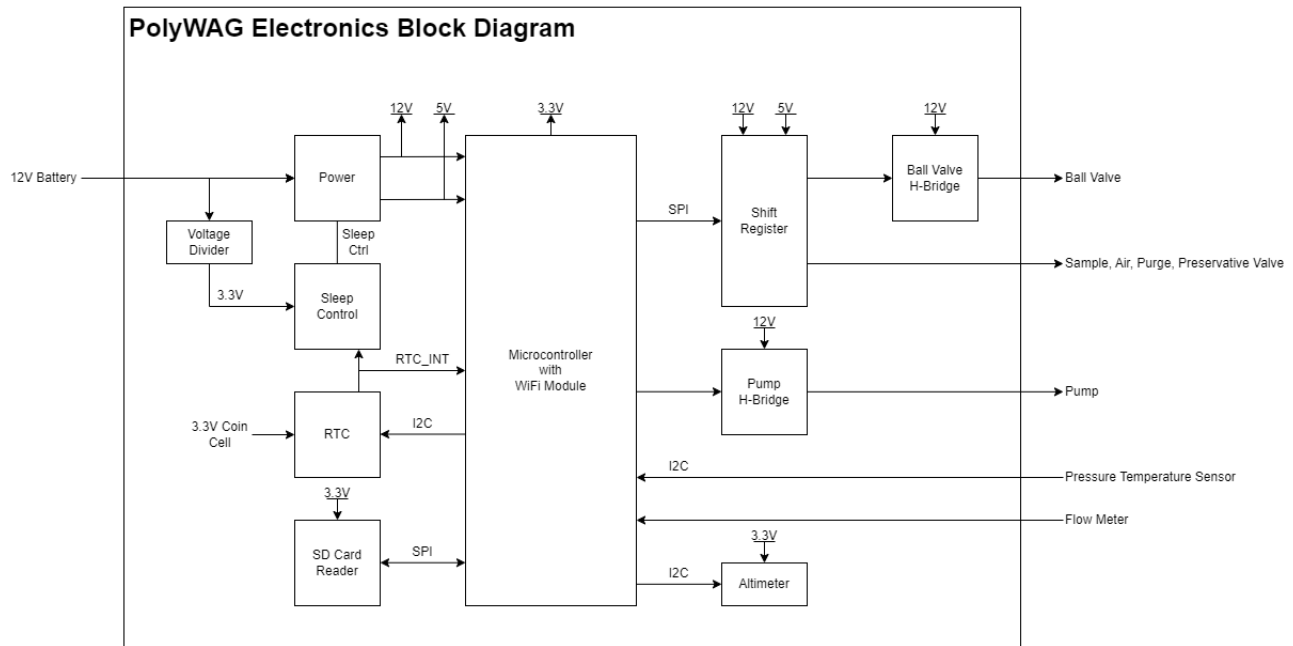


Figure 7: Block diagram of the main electronics control board

198 The power block consists of a reverse polarity current (RPC) circuit and a voltage regulator circuit. The
 199 RPC Circuit was added to protect the 12V battery from current flowing backwards through the system. While
 200 the battery has its own protection circuits, they lock the battery in the case of a short and need to be reset
 201 using the battery charger. The RPC circuit was added to prevent any “permanent” power loss while in the field.
 202 The voltage regulator circuit is a 12V to 5V regulator with an enable pin that connects to the sleep control
 203 circuit. This is used to save power during long term deployments.

204

205 The RTC and sleep control circuit are used to keep track of time and to save power respectively. The sleep
 206 control circuit controls the output of the power circuit and is constantly being power by a simple voltage divider
 207 circuit. It is basically a Flip Flop circuit that is reset when the RTC triggers an interrupt. The RTC circuit is
 208 used to keep track of the time between samples and is powered by a coin cell while power is off. This allows
 209 it to keep accurate track of time and signals an interrupt when its internal alarm is triggered. This interrupt
 210 is used to both turn power back on and to inform the microcontroller that it is time for a sample. If noise
 211 causes the sleep control circuit to reactivate power, the microcontroller will see that the RTC did not trigger
 212 the interrupt and will fall back into power saving mode.

213

214 The shift register circuit consists of 4 8-bit shift registers connect to the microcontroller via SPI. The shift
 215 registers are pull-down style shift registers where the ‘output’ pins are pulled to ground. This allows the shift
 216 registers to control devices that use higher logic voltages. This allows us to control the 27 12V solenoid valves

217 with a 5V IC. The shift registers are also used to control the H-bridge for the Ball valve. The H-bridge for the
 218 pump is controlled directly by the micro-controller itself.

219

220 The board contains an SD Card circuit for data logging purposes. The data is logged every second and includes
 221 the current state, time, and data from the sensors. The sensors include an in-line pressure temperature sensors
 222 for monitoring the lower hydraulic line and a flow meter out the output for measuring volume.

223

224 The micro-controller of choice is an Adafruit Feather M0 WiFi. The WiFi version of the Feather M0 was chosen
 225 as the user interface requires the feather to host a web-server.

226 2.5 User Interface

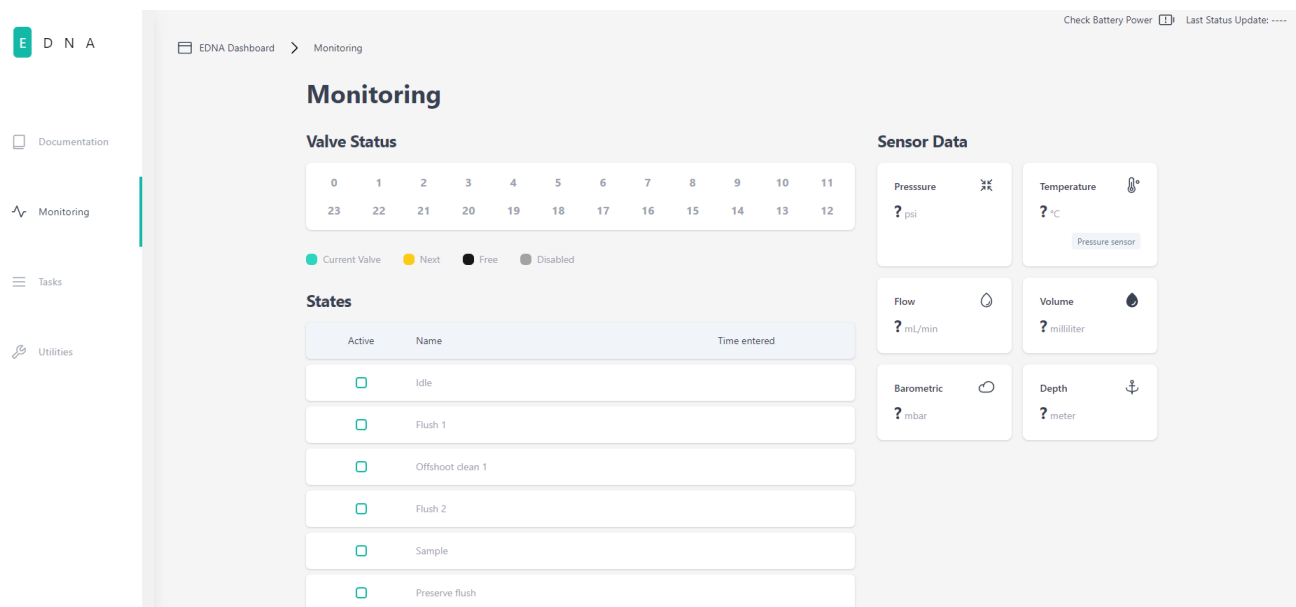


Figure 8: The monitoring page in the PolyWAG sampler user interface

227 PolyWAG Sampler hosts a webserver that can be connected to via a browser. This acts as the user interface
 228 for the system. There are three main sections that make up the user interface: monitoring, tasks, and utilities.
 229 The monitoring page displays the data from the sensors, the current state of the sampling procedure, and
 230 information on the sampling valves such as the current valve being sampled, and which valves are locked and
 231 unlocked.

232

233 The utilities page is used to activate the utilities mentioned earlier. The tasks page is where sampling tasks
 234 are created. Multiple tasks can be created, and each task is saved in memory for later modification and use.

235 This page is also where tasks can be scheduled for sampling. Each task contains the information on which
 236 valves are being used as well as for how long each state occurs.

237 3. Design Files Summary

Design filename	File type	Open source license	Location of the file
CAD Assembly	CAD file	CERN-OHL-S 2.0	...
Battery Bracket	CAD file	CERN-OHL-S 2.0	...
Sample Valve Mount	CAD file	CERN-OHL-S 2.0	...
Preservative Valve Mount	CAD file	CERN-OHL-S 2.0	...
Flow Meter Mount	CAD file	CERN-OHL-S 2.0	...
Tube Guide	CAD file	CERN-OHL-S 2.0	...
Central Assembly Mount	CAD file	CERN-OHL-S 2.0	...
Control Board Mount	CAD file	CERN-OHL-S 2.0	...
Electronics Box Lid	CAD file	CERN-OHL-S 2.0	...
Control Board Schematic	EDA file	CERN-OHL-S 2.0	...
Control Board PCB	EDA file	CERN-OHL-S 2.0	...
Switch Breakout Schematic	EDA file	CERN-OHL-S 2.0	...
Switch Breakout PCB	EDA file	CERN-OHL-S 2.0	...
UI Code	Software	GPL 3.0	...
Device (Server) Code	Software	GPL 3.0	...

Table 2:

- 238 • The CAD Assembly is a CAD file with every major components. The tubing and minor things such as
239 zip ties for cable routing are not included.
- 240 • The Battery Bracket is a 3D-Printed component to hold down the battery during transit. Paired with a
241 Velcro strap, the battery does not move.
- 242 • The Sample Valve Mount is a 3D-printed bracket that holds four solenoid valves. There are six brackets
243 in the sampler and each valve corresponds with a filter.
- 244 • The Preservative Valve Mount is a 3D-printed component that holds the preservative valve in place.
- 245 • The Flow Meter Mount is a 3D-printed bracket that holds the flow sensor to the frame.
- 246 • The Tube Guide is a 3D-printed components that helps hold the input and output tubes in place.
- 247 • The Central Assembly Mount is a laser-cut acrylic components that all of the "central" components mount
248 to. This includes the pump, input control valves, and the battery.
- 249 • The Central Board mount is a laser-cut acrylic components that the allows the main control board to
250 mount inside the electronics box.
- 251 • The electronics box lid is a CNCed Acrylic components that replaces the metal lid but maintains the
252 groove for the gasket.
- 253 • The Control Board Schematic and PCB are EDA files in the Autodesk EAGLE format for the sampler's
254 main control board.
- 255 • The Switch Breakout Schematic and PCB are EDA files in the Autodesk EAGLE format for the sampler's
256 main power switch.
- 257 • The UI Code is the codebase for the web application. The built UI files are stored in the sampler's SD
258 Card.
- 259 • The device (server) code is the codebase for the sampler that is uploaded to the microcontroller. It handles
260 all of the sampler's functions.

261 **4. Bill of materials**

262 Given the number of materials required to build a PolyWAG Sampler, The BOM will be located in an external
263 file and can be found [here](#)

264 **5. Build instructions**

265 Given the complexity of the PolyWAG Sampler, a Build Instructions section of sufficient detail would be near
266 a hundred pages long. This is why an external build guide document will be linked for those who are interested
267 in knowing how one of these samplers are assembled. This file can be located [here](#)

268 **6. Operation instructions**

269 6.1 User Interface Setup & Browser Configuration

270 The codebase for the user interface (UI) is designed to be compiled into files that are put into the SD card so
271 that the sampler machine can read. As a npm package, Node.js, specifically Node.js 10, and npm are required
272 to run the scripts to compile. Information for installing both is available [here](#). After downloading the codebase
273 repository, it can be installed using the npm CLI commands [install](#) and compiled using the build script with
274 the command [run-script](#). The compiled files are created in the /dist subdirectory of the repository.

275

276 Note that the sampling machine uses the HTTP protocol, and not HTTPS. This means that depending on
277 your browser, you will have to ensure that it does not fallback to HTTPS when trying to load the UI from
278 the sampler. The WiFi network name and password are set within the configuration.hpp file of the sampler
279 codebase, and the UI is always hosted on the IP 192.168.1.1 on that network.

280

281 Instructions beyond a technical description are available [here](#).

282 6.2 Sampler Code Upload

283 The codebase for the sampler machine uses PlatformIO, an IDE for Microsoft Visual Studio Code. Download-
284 ing the codebase also requires downloading another repository, the framework repository, that is used as a Git
285 submodule in the /lib subdirectory. Information on how to use PlatformIO is available [here](#).

286

287 Instructions beyond a technical description are available [here](#).

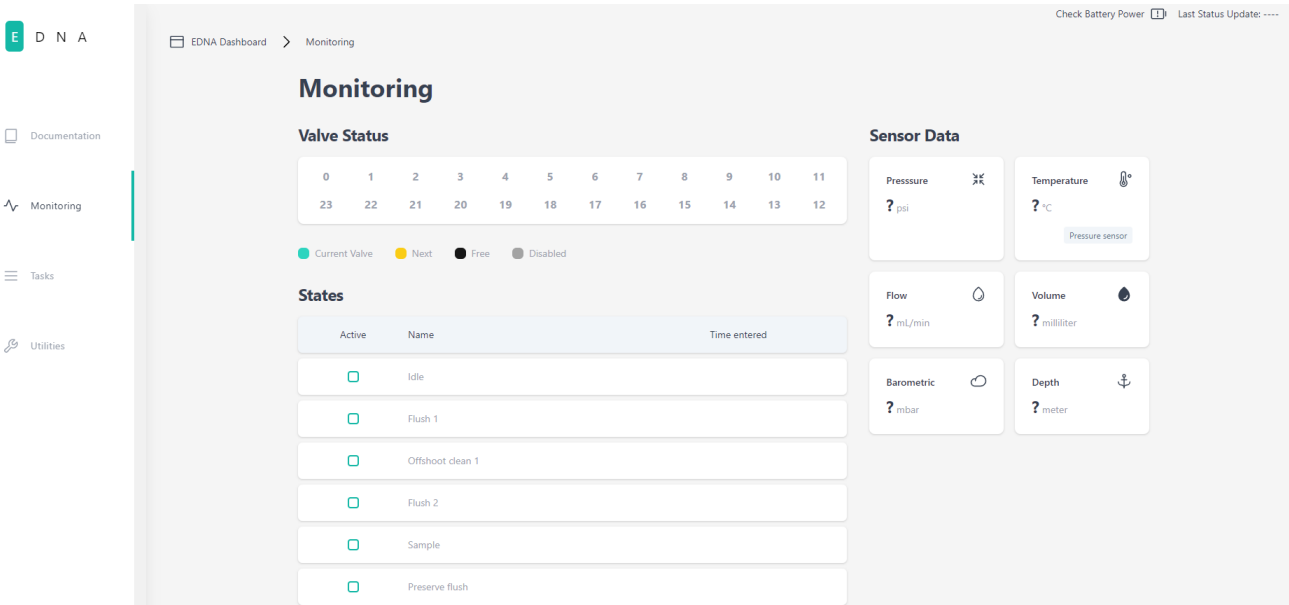


Figure 9: Screenshot of User Interface on Utilities Page

289 To ensure as little contamination as possible, the Hyperflush procedure can be used to clean the sampler. The
290 procedure is available by connecting to the UI under the utilities tab. To clean the sampler, put all inputs and
291 outputs into a container of bleach, and run the Hyperflush procedure completely five times. Afterwards, put
292 the intake line into Ascorbic Acid and run the HyperFlush procedure five times. Finally, put the intake line
293 into a DI water source and all other connections into a disposal system or sink, and again run Hyperflush five
294 times.

295 6.4 Sampler/Filter Setup

296 Filter housings have a quick-connect plug on the bottom of their assembly, which allows for them to be quickly
297 switched out with other housings as needed. The filter housings themselves are designed to be hand tightened
298 and loosened to access the membrane filter inside. The valve layout that tasks use follows the diagram below:

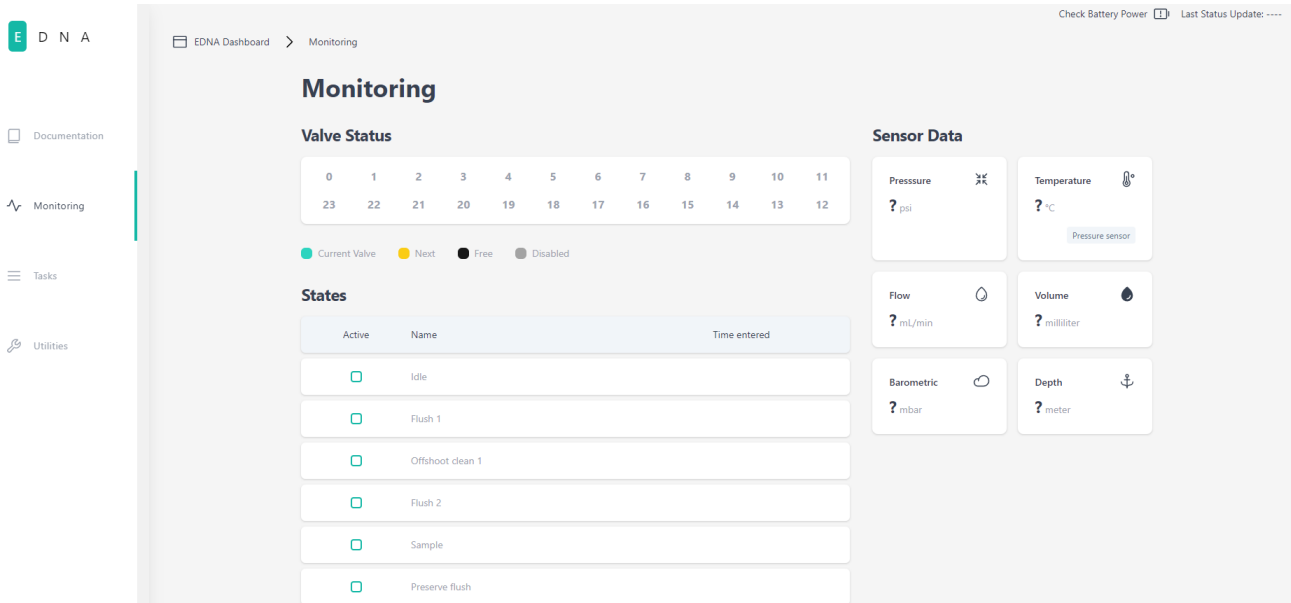


Figure 10: Visual indicator of filter labels

299 6.5 Task Configuration

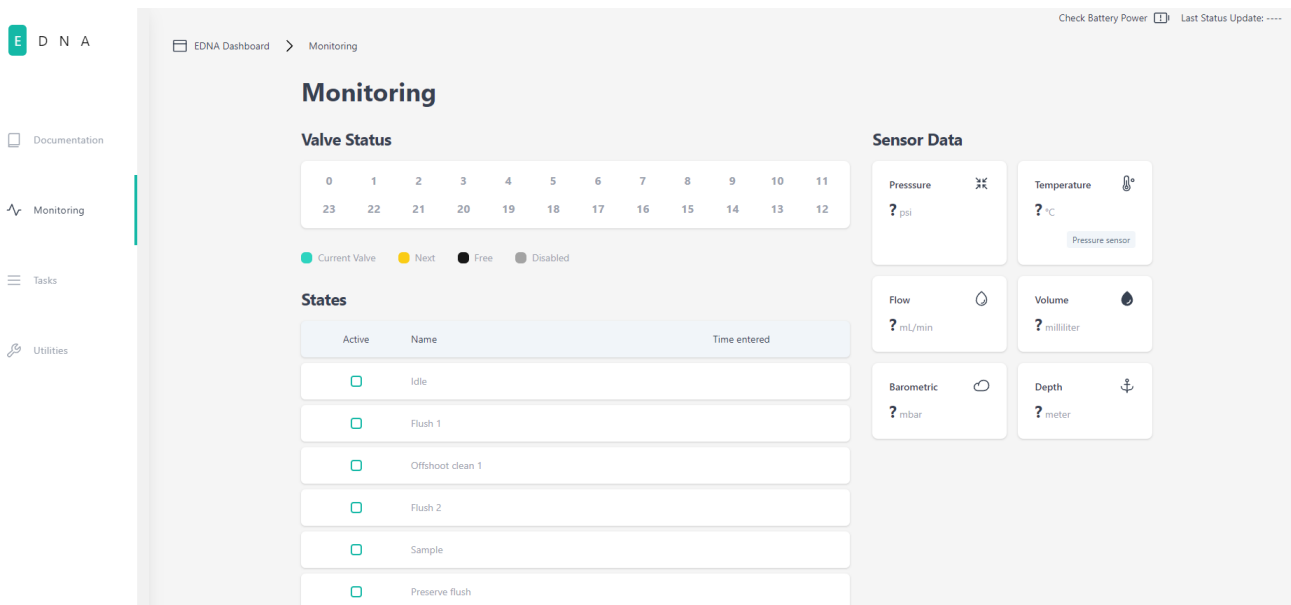


Figure 11: Screenshot of User Interface on Task Page

300 Scheduling a task is done with the UI by navigating to the tasks tab, and either creating a new task or modifying
 301 an old one. Tasks are sorted on the task tab by whether they are scheduled or inactive. Clicking on a task will
 302 bring you to the task configuration page, where you can set different parameters for the task. Scheduled tasks
 303 need to be unscheduled if they are going to be modified, otherwise they will be executed on the scheduled time.
 304 If a task has any parameters change, they must be saved before scheduled.

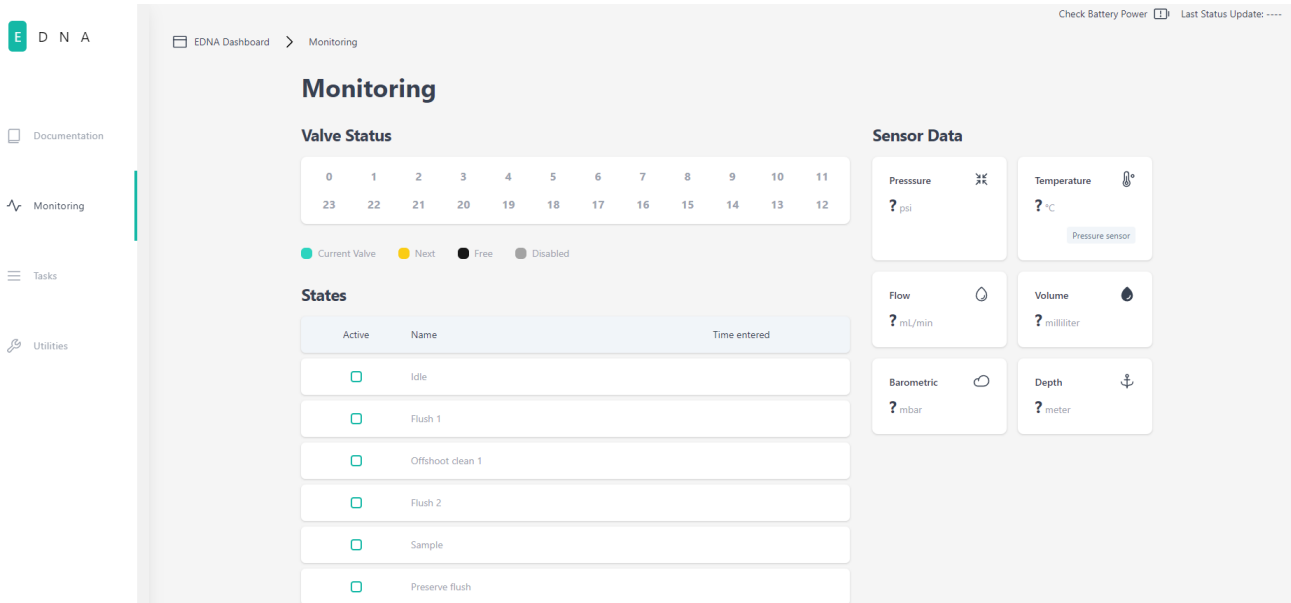


Figure 12: Screenshot of User Interface of a Task Configuration Page

Tasks can sample with multiple filters, with the ability to set the time between the ending of sampling one to the start of another sampling. Additionally, some states of the sampling procedure have a variable time that can be set by the user: flush, sample, dry, and preserve states. The sample state is unique in that the state will be considered complete depending on the time in the state, the volume of water sampled, and the maximum pressure reached while sampling. All three parameters can be set by the user depending on their use case.

7. Validation and characterization

The PolyWAG Sampler was tested by the Openly Published Environmental Sensing Lab at Oregon State University. It was characterized by a blend of isolated in-lab testing procedures as well as field tests on the Willamette River and the Irish Bend Covered Bridge In Corvallis, Oregon. To assess the capabilities of the sampler, the following evaluations were conducted: to establish the viability of field deployment at running water sources in Alaska.

7.1 Cross-Contamination Testing

The primary use case of the PolyWAG eDNA sampler is for use in water sampling to capture existing trace biological information in the form of DNA by means of filtration. In gauging the use case of the sampler for eDNA collection and characterization, we subjected the sampler to a lengthy cross-contamination study for detecting residual DNA in subsequent sample steps. Prior to sampling, a cleaning and sterilization process was conducted to eliminate any sources of contamination from previous testing. After running the sampling

322 procedure, the results were evaluated by means of Polymerase-chain reaction (PCR).

323

324 The basis for these observations is the use of Alaskan Sockeye Salmon DNA. To obtain Alaskan Sockeye Salmon
325 DNA, 1 filet of salmon (~500g) is placed in an 8 gallon container of DI Water and left for 24 hours. During this
326 time, biological material diffuses into the bulk water. After 24 hours the remaining salmon mass is removed to
327 ensure constant salmon DNA concentration across trials.

328

329 The sample cleaning procedure makes use of the sampler's HyperFlush utility to sterilize both the filter hous-
330 ings and the sampler hydraulic lines. The HyperFlush utility flushes each valve of the system sequentially with
331 any bulk liquid connected to the input line. The cleaning procedure makes use of three HyperFlush cycles in
332 the following liquid order: DI water, 6% bleach, 5% ascorbic acid. These cycles are repeated three times and
333 followed by three subsequent HyperFlush cycles of DI water to purge all residual chemicals

334

335 The sampling procedure for cross-contamination characterization makes use of a sample of DNA followed by
336 subsequent samples of DI Water. To establish experimental controls, the first sample of the study is a negative
337 control containing DI water. This negative control gauges the success of the cleaning protocol, ensuring no prior
338 contamination. Following this control, the process consists of three subsequent samples: Fish water (Positive
339 Control) and two samples of DI water. Separate inlet lines are used for DI water and fish water to isolate the
340 source of cross-contamination to the sampler itself, the lines being switched manually between samples. The
341 data collected during this study consists of four trials performed using the following task settings on the eDNA
342 UI: 1000 mL max sample volume, 4-minute max sample time, 10-second preservative flush, and 12-minute flush
343 time (~5L). After sampling, the samples were individually packaged and sent for PCR analysis.

344

345 For PCR analysis, the cellulose nitrate filters containing the salmon DNA are dissolved releasing the trapped
346 DNA into solution. Sockeye Salmon primers were added to the DNA solution. During PCR the DNA is ampli-
347 fied on a scale of 109 times the original DNA concentration. From here, immuno-fluorescence is used to quantify
348 the amount of DNA retrieved by measuring the intensity of light emitted by the sockeye-salmon DNA. These
349 values are interpreted in terms of a CT score. CT scores between 0-40 CT indicates the presence of Salmon
350 DNA, while a CT score >40 indicates no Salmon DNA is detected.

351

352 Using these processes 13 samples were taken using the sampler. This included 1 Negative control, and 4 sample
353 trials.

Sample Name	CT	Results
Negative Control	39.65583038	Positive-Low Quantity(1/2, high Ct)
Negative Control	Undetermined	
Positive Control	29.15957832	
Positive Control	29.47208214	Positive
Trial 1: Sample 1	Undetermined	
Trial 1: Sample 1	Undetermined	Negative
Trial 1: Sample 2	Undetermined	
Trial 1: Sample 2	Undetermined	Negative
Positive Control	30.94010544	Positive
Positive Control	30.6325016	Positive
Trial 2: Sample 1	Undetermined	
Trial 2: Sample 1	39.86541748	Positive-Low Quantity(1/2, high Ct)
Trial 2: Sample 2	Undetermined	Negative
Trial 2: Sample 2	Undetermined	Negative
Positive Control	29.15957832	
Positive Control	29.47208214	Positive
Trial 3: Sample 1	Undetermined	
Trial 3: Sample 1	39.46964645	Positive-Low Quantity(1/2, high Ct)
Trial 3: Sample 2	Undetermined	
Trial 3: Sample 2	Undetermined	Negative
Positive Control	29.15957832	
Positive Control	29.47208214	Positive
Trial 4: Sample 1	Undetermined	
Trial 4: Sample 1	Undetermined	Negative
Trial 4: Sample 2	Undetermined	
Trial 4: Sample 2	Undetermined	Negative

Table 3:

354 The 4 trials conducted came to varying degrees of success. In trials one and four, the CT scores of the
355 samples following the positive controls were undetectable. This indicates that there is no detectable DNA in

the PCR-amplified sample. Samples two and three contained CT scores <40 ; however, in very low detectable quantities. In both cases, the sample immediately following the control are the samples in question. In the duplicate amplification one trial contained a CT score between 39 and 40 while the other was undetectable. This indicates a very faint presence of fish DNA between samples. The cause of this can be explained by two reasons. One is that the sampler is unable to prevent cross-contamination between trials. The other is that the cleaning cycle is not adequate for ridding all DNA in the sampler. The latter of the two seems more likely because of the negative control. Like the two barely detectable samples the negative control indicated a faint positive signal. This means one of two things: contaminated sample water or insufficient cleaning cycle. Based on these results the sampler does not justify evidence that the sampler prevents cross-contamination between samples. That being said there is evidence that the flushes between samples do minimize contamination. This is supported because sample two in all trials were identified as negative. Therefore, increasing the flush and cleaning times shows promise for mitigating cross-contamination issues.

7.1.1 Turbidity

To ensure the effective use of the sampler in non-ideal conditions, the sampler was analyzed by means of a flow analysis across a range of turbidity levels. The purpose of this test is to ensure adequate flow through the filter before it clogs. For eDNA purposes a minimum of 500 mL of filtered volume is required to accurately analyze the DNA concentration of the input stream. For testing, a 5.0 μm filter and 75 μm pre-filter are evaluated.

Based on water turbidity data for Alaskan streams in April, the turbidity of the water ranges from 30-80 NTU. To reproduce this range, a 2000 NTU formazin based turbidity standard was diluted to produce 3L of the following NTU standards: 50, 100, 250 NTU. These turbidity standards were mixed within a beaker with a 2-inch stir bar at the lowest speed that a vortex was observed. The 75 μm system pre-filter was suspended two inches from the bottom of the beaker while sampling. Sampling was conducted and stopped once the flow rate dropped below 60 mL/min. This drop in flow rate indicates a filter clogging.

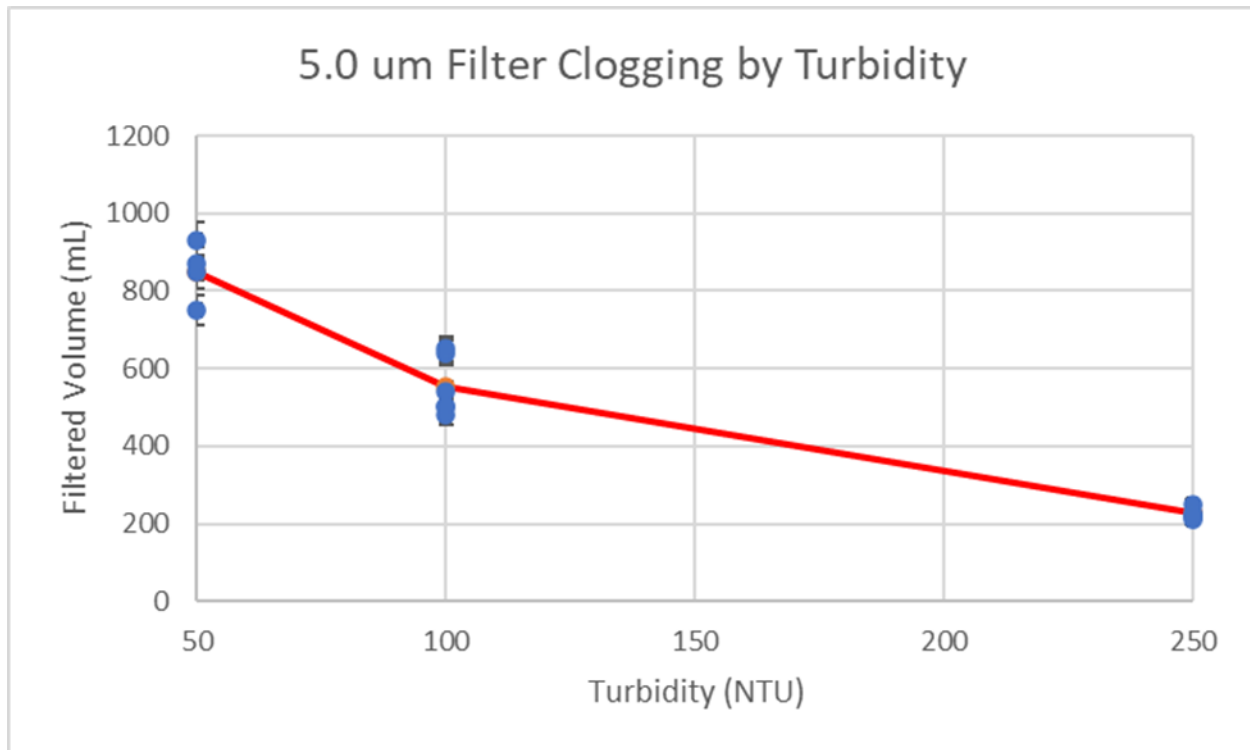


Figure 13:

380 To gauge the effectiveness of the sampler in the different turbidities the values are compared to the 500 mL
 381 minimum filtered volume. Greater than 500 mL of filtered volume indicates a successful sampler run at that
 382 water quality. As a result of this testing, the sampler is found to successfully operate in the range of 30-80
 383 NTU. The data indicates the sampler can run in water qualities up to 100 NTU before samples under 500 mL
 384 are observed. Subsequent testing at 250 NTU indicates for lower-quality water streams, the sampler's function
 385 decreases substantially, only filtering 200-250 mL per filter. This indicates that the sampler's peristaltic pump
 386 is not able to build a high enough pressure gradient across the filter. Therefore for environments with lower
 387 water quality an increase in pump power is needed.

388

389 After conducting idealized laboratory trials, the sampler was placed in the field to test the flow for real-world
 390 applications. The sampler was placed in two locations in Corvallis, Oregon: the Irish Bend Covered Bridge on
 391 Oak Creek, and the Willamette Boat Landing on the Willamette River. In both locations, the sampler was
 392 deployed on the shore of the river with an inlet hose that ran to faster-moving water. For Oak Creek, the
 393 sampler's prefilter was staked 4 inches between the water level in the middle of the stream. For the Willamette
 394 River, the inlet hose was immobilized at the end of the Willamette landing boat launch dock. The purpose of
 395 this trial is to serve as a proof of concept of the device in real world water streams. In both Oak Creek and
 396 the Willamette River, three trials were conducted in series to investigate the effects of non-idealized real world

397 conditions on the sampler. The success of these trials is based on the ability to achieve greater than 500 mL of
 398 filtered volume per sample. The two rivers provide variable conditions on the quality of the water. Oak Creek
 399 contains low river flow and high water quality, while the Willamette river was experiencing flooding with very
 400 murky, low water quality. Without a turbidity sensor, the relative turbidity of these streams were determined
 401 relative to the observed turbidity standards used for the lab turbidity testing.

Filtered Volume Field Testing			
Trial	Location	Volume (mL)	Turbidity (NTU)
1	Oak Creek	523	<100
2	Oak Creek	750	<100
3	Oak Creek	810	<100
4	Willamette River	270	>1000
5	Willamette River	300	>1000
6	Willamette River	340	>1000

Table 4:

402 As a result of these trials, the sampler performed in much the same way as the lab turbidity trials. For the
 403 lower turbidity water sources the sampler had no issues reaching over 500mL of filtered volume. For the higher
 404 turbidity sources the 500 mL filtered volume minimum was not met; however, this was expected of the trial.
 405 Altogether, these results confirm the use case of the sampler in low turbidity environments.

406 7.1.2 Flow Accuracy Testing

407 The kinematic viscosity of water varies with temperature. This indicates that the sampler will read different
 408 values for flow rate for different temperatures. To ensure the sampler the flowmeter provides accurate flow
 409 readings for the upcoming Alaskan trials, the sampler was calibrated to reach accurate flow readings between
 410 3-5°C. For accurate eDNA measurements the accuracy of the filtered volume must be within 10% of the true
 411 filtered volume. Using the 10% value as an end condition, several trials were conducted to determine the optimal
 412 volume constant for the flow meter pulses. In these trials, the outlet of the sampler is manually measured with
 413 a graduated cylinder and compared to the sampler's calculated value. The percent error was then calculated
 414 between the two quantities. To ensure accuracy across the sampler's operating range, the filtered volume was
 415 measured and compared at 250, 500, and 1000 mL for each volume constant. By averaging these values at each
 416 volume constant the average error per volume constant was discovered.

Percent error vs. Constant

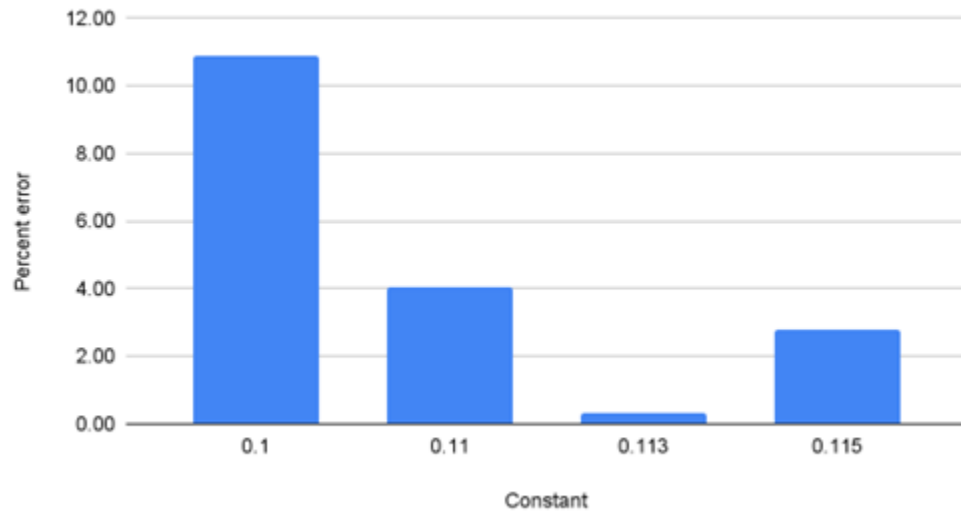


Figure 14:

417 The results determined from this process indicate the constant that is centered within the error. In other words,
418 the smaller the bar the more the positive and negative errors negate each other. This essentially centers the
419 solution in the error domain. The minimum error indicates an ideal volume constant at or near 0.113. This
420 alone does not demonstrate the accuracy of the volume measurement. Rather this indicates the most normalized
421 variability between the trials. When looking at the max error between trials the hard limit of 10% places these
422 values in context.

Max Error vs. Constant

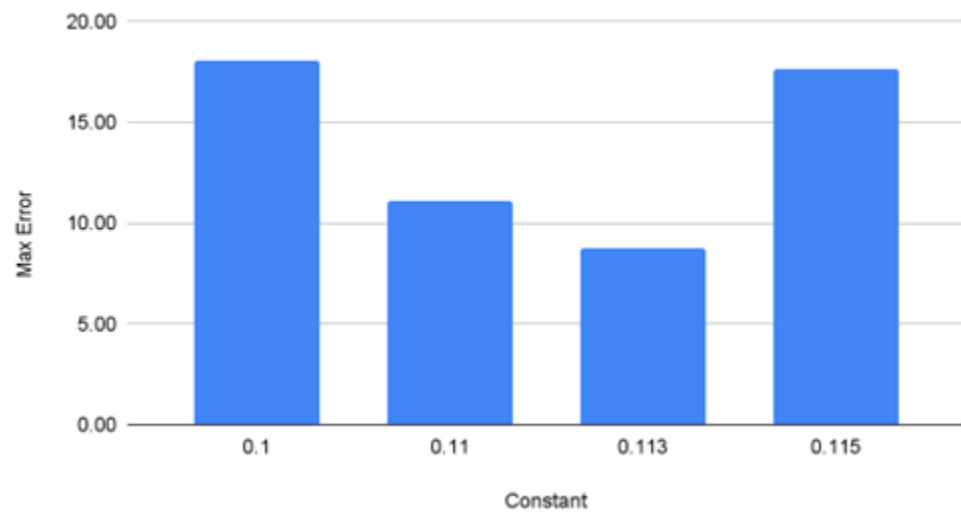


Figure 15:

423 When viewing the max magnitude of error, a convex solution with a minimum error between real and observed
424 volumes. In this context, the minimum absolute error is also easily determined to be at volume constant 0.113.
425 The max error at this measurement falls below the 10% threshold, indicating the sampler's viability for eDNA
426 collection purposes.

427 **7.1.3 Battery Life Testing**

428 Battery life varies with the ambient temperature of the surrounding environment [6]. To gauge the battery
429 life of the sampler in a cold Alaskan environment the battery was placed in a freezer at 0°C. The battery was
430 left for two weeks to simulate the two-week field deployment time in Alaska. After two weeks, the battery line
431 was plugged into the sampler while remaining in the freezer. A full 24-sample procedure was conducted. To
432 ensure normal operating conditions the full 24-sample procedure was used using the default sampling procedure.
433 Because the battery did not die during this time the sampler was determined to have enough battery life for
434 normal operation of the sampler.

435 **8. Conclusion**

436 The PolyWAG Sampler shows a lot of promise when it comes to eDNA Sampling applications. This combined
437 with the low cost of \$3800 for the materials makes the PolyWAG a potentially alternative to existing eDNA
438 Samplers. Though the cross-contamination tests need to be redone to fully verify the sampler, even the existing
439 results show a lot of promise.

440

441 There are other improvements that can be made to the PolyWAG system outside of redoing some of the
442 validation tests. This include new or improved features such as an increased the max sample volume and
443 turbidity capability, Automatic Calibration for Water Viscosity based on Temperature, as well as upgrading
444 certain components to raise the specifications of the sampler, such as the flow meter, solenoid valves, and
445 battery.

446

447 **Ethics statements**

448 The work does not use any human or animal subjects.

449

450 **Bios**

451

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455 *In addition, please list any funding sources in this section. List funding sources in this standard way to facilitate*
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464 *This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit*
 465 *sectors.*

466

467 **References**

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