A review of open source ventilators for COVID-19 and future pandemics [version 2; peer review: 3 approved]

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Any reports and responses or comments on the article can be found at the end of the article.

Abstract
Coronavirus Disease 2019 (COVID-19) threatens to overwhelm our medical infrastructure at the regional level causing spikes in mortality rates because of shortages of critical equipment, like ventilators. Fortunately, with the recent development and widespread deployment of small-scale manufacturing technologies like RepRap-class 3-D printers and open source microcontrollers, mass distributed manufacturing of ventilators has the potential to overcome medical supply shortages. In this study, after providing a background on ventilators, the academic literature is reviewed to find the existing and already openly-published, vetted designs for ventilators systems. These articles are analyzed to determine if the designs are open source both in spirit (license) as well as practical details (e.g. possessing accessible design source files, bill of materials, assembly instructions, wiring diagrams, firmware and software as well as operation and calibration instructions). Next, the existing Internet and gray literature are reviewed for open source ventilator projects and designs. The results of this review found that the tested and peer-reviewed systems lacked complete documentation and the open systems that were documented were either at the very early stages of design (sometimes without even a prototype) and were essentially only basically tested (if at all). With the considerably larger motivation of an ongoing pandemic, it is assumed these projects will garner greater attention and resources to make significant progress to reach a functional and easily-replicated system. There is a large amount of future work needed to move open source ventilators up to the level considered scientific-grade equipment, and even further work needed to reach medical-grade hardware. Future work is needed to achieve the potential of this approach by developing policies, updating regulations, and securing funding mechanisms for the development and testing of open source ventilators for both the current COVID-19 pandemic as well as for future pandemics and for everyday use in low-resource settings.
Keywords
ventilator, pandemic, ventilation, influenza pandemic, open source, open hardware, COVID-19, medical hardware

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Amendments from Version 1

The manuscript was updated in several ways:

Introduction
Based on recent news about the Italian patent infringement lawsuit this example was removed entirely.
Enhanced considerably the description of distributed manufacturing in this context.
Expanded considerably what files are for in open hardware to help better introduce those outside of the open hardware community to the concept as well as make it clearer why they need to be included.

Analysis of the literature
Added a much more explicit section detailing the meaning of open source as well as provide examples of it being misused.

Existing peer reviewed literature
Added detailed analysis of new study, which is the first fully documented ventilator in the peer-reviewed literature.
Increased discussion about need for diversity of solutions needed in a pandemic.
Updated grey literature review, added the Read review, many of the other major teams and projects, and listed the approaches.
Stressed the need to share plans

Future work
Added a section on best practices for sharing a design
Included points about the need for multi-disciplinary collaboration, the need to involve medical personnel, and to aim to publish in the medical literature.
Discussed the use of sharing incomplete designs and status tags.
Added a paragraph on the need and current status for ventilator testing.
Deleted earlier claim about easier replication in developing communities and replaced it with a call to streamline regulation while maintaining standards.

Analysis of the literature
Added a paragraph on the need and current status for ventilator testing.

Additional discussion of need for transparent quality control, standards and qualifications.

Added concept of expanding Good Samaritan laws.

Qualifed conclusions that are changing rapidly.

Added references throughout text and fixed minor typos.

For more granular details of changes see individual responses to reviewers.

Any further responses from the reviewers can be found at the end of the article

Introduction
Coronavirus disease 2019 (COVID-19), caused by a novel coronavirus (SARS-CoV-2), is in part so dangerous because it threatens to overwhelm our medical infrastructure at the regional level, causing spikes in mortality rates[2–4]. Within the medical infrastructure, there are critical technologies that are generally available, but simply do not exist in a high enough density to handle the excessive volume of patients associated with pandemics[5]. Thus, people die unnecessarily throughout the world because of a combination of COVID-19 infections and the lack of access to some of these technologies[6]. Ventilators are an example of technologies that are currently in critical short supply[7]. Mechanical ventilators are essential for treating both influenza and COVID-19 patients in severe acute respiratory failure[8,9]. Past studies have shown that intensive care units (ICUs) will not have sufficient resources to treat all patients requiring ventilator support during a massive pandemic[10–13], and ethically challenging triage[4,14] would need to be used to decrease mortality over first-come first-served basis for ventilator allocation among patients. Some work has shown promise for using a single ventilator to support multiple patients during a disaster surge[15–18]. In addition, it has already been shown that 3-D printed manifolds can assist with rapidly deploying this solution and there are open source designs[19]. This is not necessarily straightforward[20]. Although some countries, like the United States, have stockpiles of ventilators[21], there is consensus that there is not enough supply for serious pandemics[22–25] and that rationing would be needed[26]. The current medical system relies exclusively on specialized, proprietary, mass-manufactured ventilators from a small selection of suppliers. This supply model clearly fails when there is a sudden surge in demand for a relatively low-volume specialty product such as ventilators in a pandemic as analyzed here. The vast majority of medical equipment is heavily patented by a few specially medical firms that sell small volumes because during ‘normal’ times, a medium-sized hospital only needs a handful. These firms have historically aggressively protected their intellectual monopolies[27,28] to the detriment of human lives. In addition, non-practicing entities continue to attempt to actively prevent medical treatments from being deployed, even during the current COVID-19 pandemic[29]. Putting aside the absurdity of patenting and then obstructing others from using obvious inventions in normal times[30–32], in the wake of a pandemic where millions of lives are at stake, it is intuitively obvious that this type of greed is no longer acceptable.

Fortunately, with the recent development and widespread deployment of open source small-scale manufacturing technologies[33,34], there is now another way – mass distributed manufacturing[35–38]. In this new model, designs are developed and then shared with open source licenses freely on the Internet so that others can simply download and replicate the design on their own equipment, even at the household scale[39]. There has been tremendous and ongoing success of open source scientific hardware proliferation[40–45], where lower-cost and superior-functioning custom equipment as compared to proprietary scientific tools[46–49]. Based on such scientific hardware results, there appears to be a significant opportunity to apply open source design principles[50] and mass-scale collaborative distributed manufacturing technologies to make medical equipment[41–44]. In the current situation, this would at least partially overcome medical supply shortages[46–46] in general, and specifically for ventilators[51–61].

Of these enabling technologies, the most advanced is the fused filament fabrication (FFF)-class of desktop 3-D printers that have spawned from the self-replicating rapid prototyper (RepRap) project[45]. With the distributed manufacturing model, designs are downloaded even in remote areas and are manufactured on demand as needed[47] from readily available (and possibly recycled[48–50]) materials. These 3-D printers are, in general, not particularly fast when making products, but with tens of
there are, however, many other means of open hardware-based digital distributed manufacturing approaches including CNC mills, laser cutters, engravers, and etchers and other digitally controlled fabrication tools. As pointed out by Mohammed et al. many of these tools would overcome limitations of 3-D printing (e.g. speed of replication for flat parts are more easily cut from stock with a CNC tool using subtractive manufacturing than 3-D printing based additive manufacturing).

In this study, after providing a background on ventilators, the academic literature will be reviewed to find the existing and already openly published vetted designs for ventilator systems. These articles will be analyzed to determine if the designs are open source both in spirit (license) as well as required practical details. To be open source a ventilator project needs to include:

1. the design source files (e.g. computer aided design or CAD), which are needed to iterate on the design mechanically;
2. as well as production files (e.g. STL files which are used by 3-D printers to make mechanical components);
3. printed circuit board (PCB) layouts and other electronics design files to allow production as well as design evolution of the electronics;
4. bill of materials (BOM), which is needed to allow reviewers to evaluate the components employed as well as more easily find alternatives;
5. list of tools required, which are needed to determine if a device can be fabricated in a specific facility;
6. wiring diagrams, which are used to assemble the device with electronics;
7. firmware and software, which are needed to run the actual device;
8. instructions for the assembly, so makers can fabricate the device when the parts are made or acquired;
9. instructions for calibration as in many cases ventilator designs demand fine tuning to achieve adequate performance;
10. instructions for operation, so the end users can use and maintain the device.

Next, the existing Internet and gray literature will be reviewed for open source ventilator projects and designs. Lastly, as this is a rapidly evolving area, future work will be described to enable wide-spread mass distributed manufacturing of open source ventilators to fight against the current COVID19 pandemic as well as for future pandemics and to provide the devices to low-resource regions of the world that are underserved even in normal times.

Analysis of literature
Oxygen therapy coupled with mechanical ventilation is meant to support patients so that an adequate oxygen saturation (>88%) in arterial blood is maintained. The mechanical repository cycle has four parts: 1) inspiration, where the exhalation valve of the ventilator is closed and the ventilator uses pressured air to cause gas to flow into the lungs; 2) cycling, where change-over from inspiration to expiration occurs; 3) expiration, where the main ventilatory flow is interrupted and the exhalation valve opened to allow gas to escape from the lungs, and 4) triggering, where the changeover from expiration to inspiration occurs. According to Andreoli et al., mechanical ventilators are classified on what factor terminates inspiratory flow, as follows: 1) pressure-cycled ventilators terminate flow when preset pressures are reached in airways; 2) volume-cycled ventilators provide a set volume of gas to the patient over a range of pressures (but a maximum pressure is set to avoid damage to the patient’s lungs during delivery of the set tidal volume); 3) time-cycled ventilators set tidal volume by setting the inspiratory time and flow rate; and 4) flow cycled ventilators, where the inspiratory flow is terminated when the inspiratory flow rate drops below a specific level. The most common commercial modes of mechanical ventilation both provide a specified number of breaths per minute (BPM) and are 1) synchronized intermittent mandatory ventilation (SIMV) where patients can take additional breaths over the set rate and 2) assist control (AC) that uses triggering so that if the patient makes an effort to breathe, it helps them, and if not, it maintains the set rate. These modes can be used alone or in concert with 1) continuous positive airway pressure (CPAP), which uses a high-pressure reservoir and constant flow of gas that exceeds the patient’s needs; 2) positive end-expiratory pressure (PEEP), which increases the residual reserve capacity and allows for many alveoli and small airways to remain open that would otherwise close off; or 3) pressure support ventilation (PSV), which adjusts the pressure on the fly as the patient breathes to maintain a preset inspiratory pressure. For those designing open source ventilators using any of those modes and methods, there is a good base of established literature to draw upon. The classic background is available in Hess, et al.’s 1996. Essentials of mechanical ventilation, Tobin’s 2010 Principles and practice of mechanical ventilation, and Owens’ 2018. The Ventilator Book. In addition, Chapter 4 in the openly accessible book Equipment in Anaesthesia and Critical Care: A complete guide for the FRCA, provides a good starting point to help makers understand existing designs and terminology for ventilators. Texts are available for the use of a ventilator for the standard of care of patients with acute
There exists some confusion on the meaning of the term ‘open source’, which must be clarified to understand how the ventilator designs are evaluated in this review. Ventilators are hardware and thus to be an ‘open source ventilator’, a device must meet the principle and definition provided by the Open Source Hardware Association (OSHW A), specifically:

“Open source hardware is hardware whose design is made publicly available so that anyone can study, modify, distribute, make, and sell the design or hardware based on that design. The hardware’s source, the design from which it is made, is available in the preferred format for making modifications to it. Ideally, open source hardware uses readily-available components and materials, standard processes, open infrastructure, unrestricted content, and open-source design tools to maximize the ability of individuals to make and use hardware. Open source hardware gives people the freedom to control their technology while sharing knowledge and encouraging commerce through the open exchange of designs.”

Thus, a ventilator (or any other hardware) is not ‘open’ unless it both provides all of the source (as detailed above) to replicate it as well as shares it with a license that protect others’ freedoms to make or use it. There are some flawed uses of this term from two types of designers. The first type consists of designers claiming they have open source projects before they have shared the code. This is the most rampant in the current ventilator design community with many pretty renderings and high-production value videos with nothing of technical value behind them (i.e. there is no source to replicate the machine available). Most of these designers may have good intentions but the source code may never materialize. Perhaps the most highly publicized case with a good ending was of Medtronic, a large commercial ventilator company, which first announced an open ventilator project on 3-29-2020, but did not release the CAD, BOM, software, etc. to actually fabricate it. Medtronic has now released these documents under a permissive license for their Puritan Bennett 560 ventilator, which already has been commercialized ($10,000 and first introduced 10 years ago) and received FDA approval. Although these design files have been accessed over 90,000 times, this system is designed for mass manufacturing and will likely only be manufactured in that context. All ventilators made from the designs must be labeled with a warning noting that it was built in response to COVID-19, and is only to be used to address this pandemic. Thus, it should be stressed that thisa permissive license is not an open source license. The license only covers addressing the current global coronavirus pandemic, and its term ends either when the World Health Organization’s official Public Health Emergency of International Concern (PHEIC) is declared over, or on October 1, 2024, whichever comes first.

The second type of designers who misuse the term open source, have shared their code, call it ‘open’ but do not actually provide open hardware licenses or they specifically restrict the freedom of others from using it. This confusion is observed throughout the community working on COVID-19-related designs. An example of this confusion is with the ‘make the masks’ website that hosts a 3-D printable mask. They state: ‘These designs have provisional patents in place, and are intended for this goodwill campaign during the course of COVID-19. If you choose to pursue injection molding, it must be a not-for-profit venture that operates at cost to serve your local community. No license agreements will be awarded to for-profit ventures working to manufacture and distribute this product.” Specifically, in their FAQ it states “…we would like to stress the fact that these files have provisional patents and are for open source use only.” This is not what open source means and there is no ‘fair use’ provision for patents as there is with copyright.

In addition, although the STL files are available for replication one must email them for the CAD. This Montana Mask/Billings project is thus not open source by the OSHWA definition. Although it is unquestionably doing some good for the global community because some of the files have been released for distributed manufacturing, it is clearly restricting the end use. The masks take over 3 hours to print on standard 3-D printer and demand for such personal protective equipment (PPE) in some communities outstrips their local supply of 3-D printers. If manufacturers wanted to injection mold them at scale and sell them for a profit while increasing their accessibility and helping people, they are explicitly denied the freedom to do so.

The application of both of these fundamental misunderstandings of what open hardware are have been termed ‘open washing’ or ‘fauxpen’. There has thus been a call for open source hardware standardization of practices in a way that legally prevents such misuse of the term.

Existing peer-reviewed literature

The peer-reviewed literature itself is currently limited, but there has been some research on low-cost ventilation, even if the source is not available. First, a field portable ventilator system for domestic and military emergency medical response has been conceptually designed, but does not include enough information to construct it (e.g. the software was written in assembly language and not shared). This article does contain design considerations that may be useful for open source designers.

A new, compact and low-cost mask respirator concept has been developed and prototyped successfully. The blower unit was able to provide adequate ventilation to the test lungs. In addition, the integrated sensor for airway pressure was able to detect airway occlusion and leakages. It is a relatively low-power device and could be operated wirelessly with batteries. It provides a cross-sectional view of the blower unit and some details, but...
again, not enough to be considered full open hardware or to be easily replicated. It should be noted, however, that many of the components are within RepRap-class 3-D printing capabilities.

In addition, research has been undertaken on a pre-stage public access ventilator (PAV)\(^\text{105}\). The PAV is made up of several low-cost technologies including a self-designed turbine and a range of sensors for differential pressure, flow, \(F_106\), \(F_107\), and three-axis acceleration measurements. The PAV was tested under three conditions to show that it was adequate for an automatic emergency system: 1) pressure-controlled ventilation (PCV), 2) PCV with controlled leakage and 3) PCV with simulated airway occlusion. The PAV was tested for and showed effective ventilation for tidal volume, breathing frequency and inspiratory pressure. Similarly, there has been a proposal to replace artificial manual breathing unit (AMBU) bags with electric blowers to act as emergency ventilators\(^\text{106}\).

In contrast, another approach is to build a low-cost ventilator utilizing an AMBU bag that is not based on constant blower use\(^\text{107}\). The study by Mukaram Shahid showed the AMBU setup was able to perform all the functions of a conventional commercial ventilator for a far lower cost (<$100US excluding labor). The automated AMBU device was able to adjust the breathing rate and the volume of the air, which is comparable to older ventilators. However, it was also able to regulate the inspiration to expiration ratio and PEEP rate. Shahid’s system comes with two modes: 1) mandatory ventilation (as in older models) and 2) assisted ventilation (as with most current systems). Thus, the medical personnel can choose to use either the built-in triggering mechanism (assist boosted mode), which alters the inspiration pattern once it detects a change in air pressure, or set a time interval for the respiration pattern. The article contains pictures, an electric schematic, a control loop diagram, and very basic results. Again, this can be used as starting point, but there is not enough shared to replicate in the open hardware fashion.

Next, a low-cost ($420 prototype) portable mechanical ventilator was designed and prototyped that delivers breaths by compressing a conventional bag-valve mask (BVM) with a pivoting cam-actuated arm pushed by an electric motor\(^\text{108}\). This eliminates the need for a person pushing on the BVM, which is generally viewed as only a short-term solution. This system uses knobs to determine the tidal volume appropriate to the patient (usually 6–8 mL/kg of ideal body weight), adjustable BPM of 5–30, and inhalation to exhalation time ratio options of 1:2, 1:3 and 1:4 and a minimum respiratory rate\(^\text{109}\). This design is run with an open source Arduino micro-controller\(^\text{110}\) and the article provides enough details to be used as a guide for others to build a similar device, but not the full plans, code, etc. needed to qualify as an open source hardware device.

One of the most relevant designs is a pneumatic ventilator specifically designed for pandemics, which has a low oxygen consumption\(^\text{111}\). In this study by Williams et al., they describe and test three simple, pneumatically powered, low oxygen-consumption ventilators. The three designs were tested for different lung compliances (i.e. different ventilator workloads) on the delivered \(F_100\) and oxygen consumption. They used a commercial mechanical test lung for these tests (Vent Aid; Michigan Instruments Inc., Grand Rapids, MI, USA). The results of this study support the potential for mass distributed production of a low-cost, gas-powered, volume-controlled ventilator with a low oxygen consumption (anywhere with oxygen at 2–4 bar). The designs could alternatively be operated on hospital compressed air. The single use, self-inflating bellows system prevents cross contamination among patients. In addition, the system possessed one-way and safety overpressure valves, which could be incorporated into other designs. The designs are in part supplied including basic principle schematics, an example BOM, but falls far short of what is expected for a complete open hardware design.

A large multidisciplinary and international team has just published (currently accepted, available in pre typesetting form) in a study on a low-cost, easy-to-build non-invasive pressure support ventilator meant for under-resourced regions\(^\text{112}\). The design is based upon using off-the-shelf components and is comprised of an open source Arduino Nano for control, high pressure blower and two pressure transducers. It was bench-marked against commercial systems. Their supplementary material also covers the testing with healthy volunteers, but more importantly, has the basic layout of the device, PCB and circuit schematics including source files, a BOM, STLs for the 3-D printable case, description of the algorithm and the Arduino ino file, and a user manual. This device’s source is available and would represent a method to fabricate a ventilator for <$75, which has already been vetted by medical professionals. There are several interesting points about the approach used\(^\text{113}\). First, Garmendia et al. took the non-invasive medical approach, which is particularly well suited for both low-income countries\(^\text{114}\) and also perhaps during pandemics where even the wealthiest nation’s medical systems are strained. By focusing on off-the-shelf components their design could be easily replicated. In ‘normal times’, this approach is second only to systems that can be completely digitally fabricated with local resources. In pandemic situations, it exposes why it is important to have many such designs, as the global supply chains have been disrupted\(^\text{115,116}\). Normally, in the U.S. to replicate Garmendia et al.’s design based on the documentation provided would only be expected to take a few days. With the disruption, numerous makers have been having trouble sourcing supplies in the U.S., and the lowest-cost blower following the Garmendia et al. design has an estimated shipping of 8–18 days on 4-28-2020 in the U.S. There are alternatives for providing this function (both suppliers and devices), which is why it is important to have a ‘diversity of solutions’\(^\text{117}\) with as many alternative suppliers, components and possibly even digitally manufacturable parts as possible (e.g. there are already several 3-D printable centrifugal blowers developed, which would demand future work for this application). Lastly, this design did not appear to have a license associated with it being a purely medical science publication. Even the Arduino code, which did have an author information for help, did not contain any license. This could hamper rapid
deployment in some contexts as not explicitly indicating a license declares an implicit copyright without explaining how others could use the code\textsuperscript{11,12}.

There are also completely different approaches to the design of a ventilator, such as the high-frequency oscillatory ventilator\textsuperscript{13}, but only basic design schematics and preliminary testing is provided. Thus, within the peer-reviewed literature, most of the quasi-appropriate ventilator devices use a standard ventilation bag that is cyclically compressed by either an electromechanic or pneumatic setup and controlled by a microcontroller. Fortunately, the most complicated part of these designs is the controls, which is made accessible by the maturation of Arduino-based microcontrollers that can actuate and sense over a wide array of accessible and already-developed technologies (e.g. code libraries are available). It should be noted that most of the low-cost options in the literature used the bag approach, but that modern commercial ventilators are generally not manufactured with bags, bellows or pistons due to performance concerns. These concerns may be overcome by the nature of a pandemic, as well as by replacing low-cost components during failure, but this does indicate failure detection is warranted and certainly preferred in an open source ventilator design.

**Open source ventilator designs shared on the web**

There are a number of proprietary commercial low-cost products like the Pumani bubbleCPAP for infants, D-box or One Breath Ventilators (not yet for sale), which could be used to relieve some of the demand for conventional ventilators. Rather than attempting to conduct a market review of such devices, however, because presumably hospitals facing a shortage of ventilators would already consider all commercially-available and regulated/approved systems, this section will investigate the growing body of knowledge to help makers develop open source ventilators as well as the preliminary designs. This section was largely supported by information gathering of the rapidly evolving open source Internet communities such as Project Open Air, which is a group of “Helpful Engineers” on the platform Just One Giant Lab. They have congregated to help in the COVID-19 pandemic by developing open source solutions and of most relevance to this study, on a project specifically on the development of open source ventilators. Their documentation and information is freely available. Although just starting, as of 17 March 2020, they have over 2,500 registered volunteers and over 9,000 on their Slack team and by the beginning of April numbered over 15,000. In addition to an offset ventilator, in their first round of project proposals, they have prioritized oxygen concentrators and PPE as their top priority projects. In addition, their future work will focus on tube connectors and building a database for local manufacturers able to produce hardware with high score in reviews.

There are other teams working on the development of open source ventilators. Their progress is rapid and there appears to be more groups being formed and joining regularly to address needs in their communities. Robert Read et al., have been attempting to stay on top of these in a COVID-19 Ventilator Projects and Resources with FAQs available on GitHub. This resource contains a color-coded spreadsheet of the various projects and scores them on openness, buildability, community support, functional testing, reliability, COVID-19 suitability, and clinical friendliness and then ranks them by their average score. One can argue with the ranking, but the value of the resource is clear and all projects when they have obtained a reasonable level of development should ask to be evaluated. In addition, the spreadsheet has projects broken down into modular components whenever possible including drivers, monitors, flow sensors, display, oxygen blending and valves.

There are many other teams including those organizing around the open source wiki Appropedia for an open source ventilator, the Air Collective in Bulgaria, EndCoronaVirus.org, the Ventilators Collaboration Network, The Pak Innovation Club, the Oxford-based Ventilator Crowd, 1 Million Ventilators, 1,000,000 Ventilators in 100 days, #EngineersAssemble, and the Ventilator Project. YouTube has over 80 ventilator videos compiled in a list by Kramer. Facebook has an Open Source COVID19 Medical Supplies Group. There is a long-going Pandemic Ventilator Project that hosts their designs on Instructables. The RepRap community is starting on an open-source oxygen concentrator, which can be used alone or in tandem with an open source ventilator. Hackaday recently called for a medical hackathon to design and deploy an open source ventilator\textsuperscript{14}. Other communities are crowd-sourcing information about COVID-19 medical technologies and developing a Coronavirus Technology Handbook. Some resources for makers are appearing as basic specification provided by Botta. In addition, The Center for Safety, Simulation, and Advanced Learning Technologies (CSSALT) at the University of Florida has started an open source ventilator project based on hardware store components on the assumptions that the FDA will waive clearance for the bare-bones design if there is a massive shortage. The CSSALT system is one of the most professionally documented, with full files available for each sub-system and published engineering specifications for the ventilator that could be useful even for open hardware designers using completely different approaches. They are maintaining their documentation on GitHub. Many of the sub-modules, however, have not yet been developed, nor have a team working on them. Other projects are also using GitHub, like Jackson’s Open Respirator project, but are at the very beginning stages of development as of this writing. To assist these efforts the UK government has issued guidelines.

There are several approaches being attempted in the open ventilator community including pumps, pressure regulators, bellows, pneumatic systems, screw compressors, servo gas modules, fans, blowers, fluid based, cuirass (negative pressure/iron lung), and pistons. The most favored by both the academic literature as well as the maker community is just to use manual ventilators – BVMs/AMBU bags. There are many commercial suppliers available and there is very preliminary documentation for open source manual ventilation for the developing world\textsuperscript{15,16}. Although, in theory, purely manual ventilation could work to provide ventilation for patients over long periods, there is a real concern of both the availability of the needed man-power, as well as the continued exposure of the laborer. In addition, using a bag-valve mask may increase aerosolization of virus, and in general medical staff are not supposed to bag mask before intubation due to that risk. Many of the open source designs rely on this BVMs/AMBU bags approach where one automates the manual
squeezing. It only needs an exhaust system and PEEP valve. Students at Rice University have also created an automated bag-valve mask device that fits around a normal BVM using a dual rack and pinion design with a servo motor that continuously operates (open/close) squeezing the bag a specific amount to supply air. Rice provides a full non-peer-reviewed report, that is considerably richer in details than most of the others. It offers their design strategy, a partial BOM, basic testing, the source code as well as a summary of the standards and regulations necessary to go to market. Unfortunately, in their preliminary testing, the servo motor failed after only 11 hours of service and Rice is withholding the full CAD designs and results. To overcome the limitations of both the MIT and Rice designs, a group in Ireland formed and is moving along with full open source documentation of OpenLung on GitLab. The German language DIY-Beatmungsgerät project They are on their fifth iteration as of this writing based on the surrounding low-cost BVM/AMBU bag concept discussed above. Another project building off the MIT design is DIY Ventilators. Finally, the open hardware OxyGEN project is also using automated AMBU approach and although at the preliminary stages their 3-D and MATLAB design files are hosted openly on GitHub. The OxyGEN current system is under production in Spain.

Makers are also considering other types of non-invasive ventilators (NIV) such as those based CPAP (an alternative to PEEP), which is a form of positive airway pressure ventilator that applies mild air pressure on a continuous basis. A 3-D printed CPAP fan has been designed and tested as a blower and the design files (AutoDesk Fusion 360) and STLs are freely available. Another approach is to turn a commercial CPAP machine into a ventilator currently under development on GitHub by Lee. Lee built the system around an Arduino nano and has performed very basic tests to it that show that it provides enough pressure for a ventilator used on COVID-19 patients; however, there is not nearly enough information to recommend it for medical use. In addition, there are bi-level positive airway pressure (BiPAP) machines that are commonly used at home to treat sleep apnea and lung diseases as they decrease the effort of breathing by changing the pressure for inhalation and exhalation. Home-use BiPAPs could be used in place of hospital NIVs, but care would need to be taken because poor interfaces could generate viral aerosols. Negative pressure ventilation (iron lung) overcomes this problem, helping lung function by pulling from the outside (there has been some development on Appropedia). It provides a full BOM, but insufficient details for replication or complete open source documentation.

A unique design currently under development is the ARMEE Ventilator based off of painstakingly recreated Army design. Without any moving parts the system controls air flow so that its output alternates between two pressure levels. With careful design the pressures, flow rates and cycle times can mimic the output of a mechanical ventilator. This design available on GitHub being licensed a CERN Open Hardware License appears to be particularly amenable to digital fabrication. Other military units have worked on this problem in the past and are doing so currently. An example of another notable design is the AmboVent developed by the Israeli Air Force and now a non-profit. The AmboVent approach is to use a BVM system and aim at mass manufacturing for <$1,000, but their full designs are available on GitHub. Another project meant for mass manufacturing but open source licensed is the Mechanical Ventilator Milano.

In addition, several makers have developed pandemic ventilators, such as John Strupat, some time ago, but unfortunately, in addition to the lack of testing, the source does not appear accessible. Another approach is to use a blower, as in the Pandemic Pressure Control Ventilator being developed openly on Hackaday.io by Frank. Other open source projects are at their beginnings, like the TogRespirator project housed on GitHub developed for a Science Hackaday Dublin 2020, DIY and open source respiratory and a project to build an open source ventilator on GoFundMe.

In the review of Internet-reported ventilators, it is somewhat disappointing that many of the most promising designs do not share their source code. Designers that do not share their source making their projects functionally non-replicable. A current representative example would be the Utah-Stanford Ventil4us, which although looking promising and using an innovative linear actuator-driven pinch valve-based implementation has only indicated they will release their designs in the open source domain, but has not (as of 4-28-2020), despite preliminary evaluation and submission to peer review. In fact, in many cases, little more than a picture or video are available (e.g. Drexel University’s Dragon Ventilator Project as of 4-28-2020). The newer projects do tend to be following better documentation protocols. Unfortunately, despite the many promising approaches in the maker community, the one problem that the vast majority of the current partial designs have in common is that there is not nearly enough information available about their performance to recommend them for medical use.

Future work needed
It is clear from this review of the peer-reviewed, gray and open web literature on open source ventilators, that there is considerably more work to do. The tested and peer-reviewed systems lacked complete documentation and the open systems that were documented appropriately were either at the very early stages of design (sometimes without even a prototype) and were essentially only basically tested (and some were not tested at all). With the considerably larger motivation of an ongoing pandemic, it is assumed that these projects will garner more resources and members (as is happening with the Open Air Project) to reach a critical mass to make significant progress to reach a functional and replicable system. Although the motivation of working during a pandemic on a device that may save your life is high, the access to resources, however, is far from optimal. Already, many locations throughout the world are essentially forcing citizens to shelter-in-place, which restricts access to government and university labs, as well as to makerspaces and fab labs. In addition, some areas of the world are suffering from supply disruptions and shipping challenges. This perhaps underscores the importance of developing open source hardware for disasters.
before the disaster strikes. Future work is needed to develop policies and funding mechanisms for such work as it appears rational to make a small investment in developing and sharing the designs for any critical hardware.

For those planning to work on (or who are already working on) the development of open source ventilators one of the primary challenges is to determine when to share your designs. People are literally dying from lack of ventilators and it is hard for designers not to feel responsible if they are reasonably confident a preliminary device design would possibly prevent those deaths if shared. Many makers follow this belief and often aggressively share their content before there is any evidence that it works. At the same time, well intentioned engineers and designers can have their work mischaracterized and promoted before it is documented by overly aggressive public relations outfits at both companies and universities, which has greatly added to the clutter in this space. On the other hand, as these are medical devices, which literally can mean life and death for a patient, it is reasonable to want to follow the conventional hardware developers’ method: wait to release it until it has been fully tested. In addition, the effort and time it takes to do full documentation correctly may also appear to be lower priority than the making, prototyping and testing of the device itself. However, as Bowman[12] points out the “the intent to share a design in the future misses the myriad benefits of open hardware - in terms of scrutiny, feedback, and improvements from the community. It also stifles the development of a community around the design, and there are many cases of promised openness never materialising.” It appears clear that if a project is announced as open source it should include all of the code that is available.

A best practice for open sourcing a design is simply to maintain full documentation of your project as you are working on it in an open platform like GitLab or the Open Science Framework. Designers can start with their designs private, but as parts of the device become ready for ‘show time’ they can click share and everyone can get to the source code immediately. When this is done open announcements can be made to recruit feedback from the community. As ventilators are complex devices made up of mechanical components, electronics, sensors, and firmware/software it is challenging to have a complete device ready (Ideally the systems could reach a maturity in which the subcomponents could be made modular[13]). These particular devices are also specifically targeted at medical doctors therefore as Farre has pointed out “it is important to make an effort to actively involve health professionals for both design and testing, trying to publish contributions in medical journals”[14]. The diverse multidisciplinary skill set needed to develop a successful open source ventilator is therefore extremely challenging and explains in part why there is a lack of completed designs. Academic authors can still operate normally with publishing even if it is a hardware related publication. If anything, aggressive sharing before formal peer-review in this way protects precedence and scooping from unethical actors.

To overcome this challenge of partially completed design release, the method that Appropedia uses for status tagging projects can be borrowed to make it clear what level of development each of the subcomponents is currently at. Appropedia is a website primarily dedicated to developing open source appropriate technology (OSAT)[15] for the developing world in a massive collaborative fashion akin to Wikipedia[16,17]. Thus, because of the enormous relative investment someone in a developing community must make to fabricate a device a clear designation of the status is provided to readers. The Appropedia Status tags are color code as follows: red for designed, i) orange for modeled, iii) yellow for prototyped, iv) light green for verified by a specific organization, and v) dark green for deployed listing organizations and numbers of replicants at specific locations in the world. Status tags are placed at the top of a wiki page for a project so that users quickly know what level of risk they must accept to fabricate the project OSAT. For those doing ventilator development on a MediaWiki wiki, the source code from Appropedia for these tag templates can be borrowed immediately as it is under CC-BY-SA, but the concept can be easily adapted to any repository. Following this methodology at the initial writing of this article many of the grey literature devices would be in the red, but as of this update have moved into yellow and presumably will make it to light green and have functional designs verified by someone with an artificial lung shortly (Michigan Tech’s Open Sustainability Technology lab, for example, has offered this for anyone that has developed and shared a fully open source device). The status of the subcomponent and the project as a whole should be clearly visible on the project pages.

It is clear, that open source ventilators would ideally be modular so development can occur in parallel more easily, completely open (by the OSHWA open source hardware definition) with a transparent design and testable with a fully transparent and open validation methodology ideally itself done on vetted open hardware. To assist to democratize these testing steps further there are several efforts to develop an open source artificial lung specifically to test ventilators underway including the VentMon project documented openly on Github and currently at version 0.1 and the Ventilator Inline Sensor Package (VISP) on Hackaday.

This review article uncovered other limitations to this approach. First, due to 1) potential legal issues challenging an open source ventilator design. It is important that open source ventilators (and other open medical hardware) meet the high-standard levels approved for use in developed countries. The ethics of saving lives, however, require regulation to be more streamlined so that lives are not unnecessarily lost during delays for approval[18]. Second, because of the general lack of useful technical information in patents (the average Instructable generally has more useful information for constructing a device than a patent despite that being a requirement of obtaining a patent), patents were not included in this review. It should be noted, however, that there are currently over 277 inactive patents in addition to those that have expired covering ventilators. Researchers can obtain this list with direct links to these patents using the Michigan Tech Free Inactive Patent Search[19]. There may be useful information contained in those documents that could help open source ventilator designers.
Another challenge with this approach is maintaining a proper level of sterility of devices fabricated using distributed means. Specifically, for the FFF-based 3-D printing parts, it has been reported that the prints are sterile at the time of print\textsuperscript{12}. If not kept in a sterile environment, however, they could quickly become biologically contaminated. One approach to deal with this is to use washing or a chemical bath. A relatively complete analysis of the chemical compatibility of commercial 3-D printed plastics is available\textsuperscript{10}. If a specific polymer is needed that cannot be 3-D printed easily, it is possible to make molds in high-temperature plastics, such as polycarbonate, and then use lower temperature plastics to make disposable single use plastic parts\textsuperscript{13}. Similarly, silicone molds can be made from a 3-D printed reverse mold and used in the same way\textsuperscript{12}.

Even when more mature open source ventilator designs are available and can be safely manufactured by a distributed means, another area of critical future work is validation of these designs. In the medical sciences, open source devices like syringe pumps\textsuperscript{13,14} are already established\textsuperscript{15-18} and have been developed into sophisticated devices\textsuperscript{19-22}. However, these devices are used in labs in general and not on people continually. For medical professionals to use an open source ventilator, they first must be convinced it will do no harm to them (or others) as well as to the patient. As COVID-19 was reported to spread via droplets, contact and natural aerosols from human-to-human, there has been a concern that high-risk aerosol-producing procedures may put medical personnel at high risk of nosocomial infections, which is a concern for some designs reviewed here\textsuperscript{23}. During the airway management, enhanced droplet/airborne PPE is needed and the study by Zuo et al. provides a list of other recommendations to overcome this challenge. There have been some developments in 3-D printing some of these PPE\textsuperscript{24,25}. Similarly, for designs that could aerosolize the virus, a negative pressure room would be necessary and future work is needed to design an open source approach to creating such rooms. Likewise, the greatest concern for untested open source ventilator designs is that they harm the lungs of the patients; there is significant literature in this area of ventilator-induced lung injury\textsuperscript{14-15}. There are, however, solutions for preventing this, like controlling the tidal volume\textsuperscript{16,17}. Thus, the designers of open source ventilators must ensure that their designs have safety features to prevent ventilator-induced lung injury, as well as having basic testing of the prototypes to ensure that the designs themselves are thoroughly vetted. This must be done using yet-to-be-developed transparent process controls and quality assurances as well as tests. There has been some preliminary efforts in this direction for minimal qualifications based on the UK guidelines and the more than 70 COVID-19 Ventilator Validation Tests under development by Public Invention. More work, is needed however, to make freely accessible standards and protocols for testing, quality assurance and use of ventilators. Today such information is behind a complex web of paywalls from standards vendors, equipment suppliers, and copyright holders from publishers, universities and medical vendors and in the worst case are simply proprietary trade secrets. A future review is needed to find pathways to accessible sources of reliable information in each of these areas to accelerate the development of low-cost and free and open source medical hardware of every kind including ventilators.

Within the open source scientific equipment community, such procedures are relatively well established and have been working reasonably well through normal peer review of hardware-based articles like those published by HardwareX, the Journal of Open Hardware, and PLOS One. For medical equipment that could be all that stands between life and death, this vetting is even more important and open calls for papers for a Special Issue on Open-Source COVID19 Medical Hardware are attempting to address this.

However, technical validation may not be enough. Medical hardware used on humans is also more complicated, as any studies involving humans needed to verify its functionality on people, need institutional review board approval and, if in regulated areas like the U.S., such a study would need an Investigational Device Exemption to allow for a non-FDA approved device to be used as part of a study. This is only a temporary approval and the full device would need actual FDA approval for legal deployment unless the laws are changed (or were temporarily suspended during a pandemic). These same regulatory roadblocks are in place in other nations, which has conventional ventilator manufacturers skeptical that even conventional manufacturers of other products (e.g. vacuum cleaner and automobile manufacturers are doing this now in the UK) could switch over to produce ventilators\textsuperscript{26}. Clearly, this process is a problem during a pandemic. Both for the current situation and during potential future situations, there is a need to limit liability on the part of the designers, makers and users of such open source medical hardware\textsuperscript{27}. One approach is for ‘Good Samaritan’ laws to be expanded to protect both the makers and designers of open source medical hardware\textsuperscript{28}. Substantial future work is needed in this area. Finally, it should be pointed out that personnel and training can become limitations to deploying mass medical efforts, even if open source ventilators are available. So, future work is needed to create training materials and translate it into the languages spoken throughout the world as well.

Conclusions
There is clear technical potential for alleviating ventilator shortages during this and future pandemics using open source ventilator designs that can be rapidly fabricated using distributed manufacturing. The results of this review, however, found that the tested and peer-reviewed ventilator systems lacked complete documentation (with one recent exception) and that the current open systems that were documented were either at the very early stages of design or had undergone only early and rudimentary testing (although this is changing rapidly). With the considerably larger motivation of an ongoing pandemic, it is assumed these projects will garner greater attention and resources to make significant progress to reach a functional and easily replicated open
source ventilator system. There is a large amount of technical future work needed to move open source ventilators up to the level considered adequate for scientific-grade equipment and further work still to reach medical-grade hardware. Future work is needed to achieve the potential of this approach not only on the technical side, but also by developing policies, updating regulations and securing funding mechanisms for the development and testing of open source ventilators for both the current COVID19 pandemic, as well as for future pandemics and for everyday use in low-resource settings.

Data availability
No data are associated with this article.

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Upon reviewing the comprehensive modifications made by the author, I am deeply satisfied the present manuscript should be accepted for indexing.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Medical Devices, Bioengineering, Product Design, Sustainable Technologies and Materials, Open Source Hardware

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 01 May 2020

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No more comments

Competing Interests: No competing interests were disclosed.
**Reviewer Expertise:** Respiratory mechanics, Respiratory medical devices, Biomedical engineering.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Version 1**

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**Introduction**

The manuscript presents a review of open sources ventilators with particular emphasis towards applications in the COVID-19 pandemic during the early phases of 2020. Given the submission date of the manuscript, which has coincided with the emergence of the COVID-19 pandemic, the review is very topical, in addition to being thought provoking and insightful. COVID-19 disease is caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), where the presenting symptoms of a patient are predominantly based upon respiratory ailments, requiring ventilation based equipment to treat the most severe of cases. As the virus infiltrates a greater frequency of the general population, the availability of ventilation based products has rapidly become one of the primary causes for concern not only in developing, but in developed economics. Given the disruption of standard medical device supply chains during a pandemic, a viable alternative to meet the demand for ventilation equipment from healthcare providers is to turn to communities of designers, engineers, industrial specialists and knowledgeable maker enthusiasts to develop easily accessible, low-cost and open source alternatives to traditional devices. The manuscript is therefore a vital piece of documentation to assist researchers in global efforts to create ventilator alternatives. Equally, the manuscript does an exceptional job at highlighting the current state of the art in this area with balanced and considered conclusions throughout. **I would therefore very much recommend this article for indexing, which will be of great interest to the scientific community and those seeking to develop their own open source solutions.**

**Recommendations and Thoughts**

It is noted that the article is based on the premise of discussing ventilator technology which are deemed open source. As review manuscripts generally attract a wider target audience that scientific bodies of work, it would seem appropriate for the author to define what is meant by open source, for the benefit of unfamiliar readers. Indeed, one may see general scientific publications are open source information by virtue of the information being in the public domain. However, it is clear that the authors perception of this would require a deeper level of ‘disclosure’
of the innovations presented by various research groups to allow for ready duplication and adoption of such systems. This is starkly evident later in the manuscript during discussions of existing literature. Therefore, the distinction of what constitutes open source in this context should be explicitly defined, ideally within the introduction.

In light of the rapidly evolving nature of the pandemic and the volume of initiatives that are attempting to provide viable, often, open source solutions, there is likely to be some developments that have not been addressed in the review. Generally, the author has done an exemplary job of drawing the readers attention to many of the most topical and noteworthy examples. However, one would imagine as we approach the end of the pandemic period that there is likely to be a wealth of additional technologies which will have surfaced and so perhaps a follow up review may be justified, ideally inviting multiple authors involved in such projects to contribute. It is however noted that certain information within the manuscript has since evolved, specifically relating to the comment in the first paragraph of the introduction, where it is mentioned that ‘……for a recent example, consider the fact that a manufacturer threatened to sue a maker for 3-D printing life-saving valves in Italy for patent infringement….’ Upon further development of this story, the company had made official statement to clarify that they did not attempt to sue the party which made ventilator valve parts but had primarily withheld designs based upon medical device regulation. Although withholding designs during this particular circumstance when supply chain needs could not be met resulting in potential mortality of patients, there is an argument to disclose such information despite legal implications. However, this is very much different to the notion of the company suing the Italian firm. At the time of writing the article this would not have been known to the author, but given the controversial nature of the comment, I would request the author to reword this sentence to reflect the final outcome of this case study.

During the introduction when discussing the very many technologies available to the open source community, the discussion preferentially revolves around the use of rep-rap 3D printing, under the notion of digital fabrication technologies which have distributed manufacturing potential. This is a very important point to make by the author as distinctions are made as to why this approach would provide added value within the context of a pandemic. In particular, designs may be shared both at a national and international level using internet based data transfer, while leveraging manufacturing and technical capacity closer to the point of use. Such capacity has long been utilised by the open source community, providing strong resilience in instances when typical supply chains are disrupted, as would be the case during a pandemic. However, this discussion appears to be somewhat incomplete for readers who are unfamiliar with this approach of manufacturing. More specifically, it may be useful for future readers to hear some mention of other digitally driven distributed technologies, such as milling/CNC machining, laser engraving/etching and other digitally controlled tools. Indeed, several of these technologies would serve to reduce or eliminate the perceived limitations of 3D printing both in terms of manufacturable materials and speed of manufacturing. Please could the author include some additional discussion here to contextualise available options to the open source community with respect to digital and distributed manufacturing.

The author presents an eloquent attempt to discuss both the academic and non-academic ventilation systems, citing many interesting studies and raising most of the key facets of each respective technology. Following from the previous comments regarding the open source nature of a given study, it is highlighted that sadly many academic studies, though presenting some remarkable feats of engineering, simply do not provide sufficient information to allow for other
researchers and experts to duplicate a respective ventilation system. This is indeed a shame and highlights perhaps a necessity for researchers to adopt a more open framework of reporting academic findings and equally for reviewers to encourage and accept such ways of reporting. In light of major global health catastrophes such as pandemics, such openness within the literature may in fact prove advantageous to hasten innovations to tackle the detrimental effects on a given population.

The discussion on the current open source efforts appears to cover several of the major projects to the awareness of this reviewer, but more crucially provides a good cross section of important developmental aspects to inform the reader. As highlighted previously, given the rapidly evolving nature of the pandemic and the willingness and passion of researchers and technical experts, new attempts to create ventilation products are arising on a weekly, if not daily basis from groups around the world. Therefore, in this reviewer's opinion, to cover all such attempts would be impractical but also unnecessary given that the overlap of technical development with existing projects. As with the previous section, it would have been good to present annotated diagrams and a summary chart/table of the systems broken down into the primary attributes that fulfill the requirement for ventilation. In reading this section it was difficult to surmise how these efforts were truly moving towards a functional ventilation device. One of the more noteworthy examples from Rice University, with credible data to back the developments to date was sadly hindered by component failure after only a 11-hour evaluation period, which falls considerably short of a functional ventilator. Clearly, several if not all examples highlighted by the author raise several notes of caution toward open source design, namely the clarity and robustness of evaluation, which lacks the rigour of typical academic scrutiny in addition to the lack of standardization of components which are suitable for purpose. Arguably, the author to varying degrees' highlights this by stating the case studies in question are either in the very preliminary stages of investigation, show a lack of relevant performance data or a Bill of Materials (BOM) which enables scrutiny of the components employed. I believe there is a missed opportunity of discussing these elements in greater depth, which is vitally required if open source ventilators are to truly enter mainstream acceptance and use. I would very much welcome further discussion on limitations with appropriate recommendations, which both do not stifle the breadth of design ideas by the community, but also provide substantiative guidance to direct those involved to be mindful of critical milestones and 'codes of best practise' during the journey from inception of idea to final working and 'usable' ventilator. For example, looking at open source efforts for Personal Protective Equipment (PPE) we see that the 3D Printing community rapidly converged towards the preferential use of PETG polymers for manufacturing, owing to the factors of mechanical stiffness providing a semi flexible yet robust part, and importantly biocompatibility for limited human contact and being food grade to allow ease of decontamination. Equally, strict protocols were developed to minimise contamination of parts during printing, handling and shipping to both reduce the spread of the virus within the supply chain and to follow some element of best practise, similar to constraints set by medical device companies upon their manufacturing procedures. It would be very useful to the open source community if the author would share their thoughts in more explicit detail to provide a template that could be built upon for future efforts, outlining best practise from initial design ideation to working prototype. Arguably, the imperative of this is much greater than with PPE given potential intrusive nature of ventilation systems providing a clear route to internal infection through the lungs.

One element that appears to be missed in the present review is an overview of current commercial systems and evaluation of their performance characteristics to be both efficacious and
crucially to prevent unintended harm to a patient. The author does cite an existing review of commercial ventilators by Pham et al, and so it would be unnecessary to conduct a repeat of this work. However, what would be relevant in the present manuscript is a discussion focusing on the regulatory and quality assurance aspects and how these would align and differ in the approach of open source ventilation systems. Indeed, commercial devices must adhere to the very strictest regulatory scrutiny to be classified and used as a medical device, particularly given the invasive nature of their operation to either supplement or take complete control of a patient's breathing. As such there are tightly regulated frameworks of 'fail safes' to ensure that every component used falls within acceptable usage limits, that construction of such devices follow strict regimes for assembly and minimisation of contamination, that tests are conducted to evaluate the working performance of each device, that there is a robust training and best practise usage protocol, amongst many other safety measures and supply chain demands. Arguably, one of the biggest reservations by the commercial sector regarding open source hardware for medical device technology are the lack of process control, quality assurances and regulations regarding technological development and best practise. It would be valuable for the readers to have some insight into what safety and evaluation procedures that are conducted and how such measures would be replicated in open source systems in general terms, highlighting key challenges which can be built upon for the future discussion.

The final sections of the review offer an exceptional summary of the state of art in the field of open source ventilators, highlighting several challenges and opportunities in this space. The author rightly highlights the very early stages open source ventilations systems currently are at and that we should not expect these systems to be working within a clinical setting in the immediate future. Despite this, I for one feel optimistic given the current work that is underway, our ability to access a wealth of digital knowledge, alongside the availability of hardware and manufacturing resources at our disposal. Arguably, the open source community has never been better equipped to make positive impact on the world during the COVID-19 pandemic. It is also the opinion of the reviewer that there remains many more opportunities for both growth of the open source community and to leverage the expertise synergistically with other academic groups to more rapidly advance our preparedness for emergency situations.

Reflecting upon the manuscript, I could not help but feel there are differing schools of thought that of the traditionalist and the open source innovator. The traditionalist will operate with robust scientific rigour but will provide limited information scientific manuscripts, be open to patent and potentially restrict free flow of concepts and with respect to medical technology will strictly follow medical regulatory frameworks. The open source innovator is generally driven by an overwhelming sense of openness and transparency in their work, with the belief this will help proliferate and see ideas adopted faster for anyone's benefit, albeit on some occasions operate with a naivety towards regulatory and best practise aspects. Initial discussions by the author drew distinction between efforts of the academic and non-academic communities, which very loosely are comprised of these two types of innovators, with the exception of a few 'maverick' academic groups. However, there was little discussion of strategies as to how these two communities may come together in cooperation and bridge any perceived differences in thinking. Clearly the academic community prides its outcomes based on empirical evidence, the careful scrutiny of data, alongside objective design performance metrics, attributes that the authors clearly outlines as shortfalls of the general open source community. Conversely, the non-academic community provides a wealth of creativity, ingenuity, alongside technical prowess, finding often remarkable and highly efficacious solutions working with limited resources and minimal dependency of
specific supply chains. I would therefore strongly welcome the author to add further discussion towards strategies on how both communities, which for the most part work independently, could align agendas to realise opportunities that transcend the sum of the two parts. Indeed, I think it more critical for the open source community, based upon the reflection of the author, to be more engaged in traditional scientific process and to incorporate this into their thinking to hasten product development for evaluation.

One element that gave me considerable food for thought, was the notion that developing nations have a perceived advantage during global health emergencies due to their more relaxed legal and regulatory frameworks to deploy open source ventilator systems. I am not entirely convinced that this is the case and indeed the author makes several valid arguments to the contrary, mainly that the technology is not significantly mature to function as intended and without consequence. Such suggestions can indeed have several unpleasant connotations from a legal and ethical standpoint and so I would encourage the reviewer to consider an amendment to this comment. Ultimately, I believe it was not the authors intension to imply this given previous discussions, however the context of this point should be framed better.

Summary
Overall, despite the sombre theme of the review, the author has done an admirable job of bringing together all the relevant themes relating to open source ventilation systems. One of biggest take home messages from the review is how much potential exists with the open source community to provide cost effective, robust and timely medical device solutions, which may be far less susceptible to supply chain disruption and leverage a greater capacity for localised fabrication using the distributed manufacturing model. This capacity can only be realised by continued development of existing open source projects, increased dialogue with academic groups to work collaboratively to validate and iteratively improve ventilation system concepts for maximum efficacy. Equally, there is a clear need for regulatory reform which appreciates the evolving circumstances during a global health crisis and could provide an alternative framework to leverage capacity outside of typical medical supply chains to supplement efforts on the ground, as and when appropriate. What this framework should look like is another debate entirely, but this article makes an elegant argument for the debate to be had.

Despite the infancy of open source ventilation systems, much potential exists and it is an exciting time for developers to continue their efforts towards working solution. It is exciting to see what may be a paradigm shift in how we perceive and operate globally in the medical device sector, particularly in light of the recent issues during the COVID-19 pandemic, which have decimated supply chains, while the shear volume of cases has put a drain on medical resources. The pandemic has already seen the use of open source designs, manufactured in a distributed manner, make impact to supplement shortfalls in PPE equipment. Could the same in time be true for ventilator technologies based on the balance of growing demand and available resources? Only time will tell. I reiterate the relevancy of the article by the author and the manuscript has been a very thought provoking document to read and digest. I would whole heartedly recommend this article for publication and encourage researcher and technologist in the field to draw inspiration from the insightful and thought provoking arguments outlined. I do welcome a time when the lessons we are learning during the pandemic lead to a more caring and equitable world for us all, and it feels from a technological standpoint, that open source innovation will be part of that story.
Additional minor points for consideration

○ The hyperlink for Ref 82 needs revising as it links to an error page.

○ One the first paragraph of page 4 ‘breadth’ has mistakenly been used in place of ‘breath’. Additionally there are some minor grammatical errors toward the end of this same paragraph that need amending, specifically the sentence starting with ‘texts area available for the …….’

○ Generally speaking, it would have been very useful to contextualise much of the discussion with annotated diagrams of several key open source innovations, to give the reader a real feel for the types of devices in development. Indeed, this is typical of many academic based reviews of the scientific and gray literature. Unfortunately, this has not been the case in the present manuscript and I would invite the author to consider such an amendment.

Is the topic of the review discussed comprehensively in the context of the current literature?
Yes

Are all factual statements correct and adequately supported by citations?
Yes

Is the review written in accessible language?
Yes

Are the conclusions drawn appropriate in the context of the current research literature?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Medical Devices, Bioengineering, Product Design, Sustainable Technologies and Materials, Open Source Hardware

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 28 Apr 2020

Joshua M. Pearce, Michigan Technological University, Houghton, USA

Thank you for the rigorous and exceptionally detailed review.

Following your recommendation, I have added a much more explicit section detailing the meaning of open source as well as provide examples of it being misused.

I have removed the now incorrect example of the manufacturer suing over reverse engineered valves. Although it may be interesting to note I contacted the individual who
reversed engineered them before the initial article was published and I have yet to receive
the STL files.

I share your frustration with the current lack of appropriate sharing. I have included a more
detailed review of the existing designs – but as you point out it is impractical to do
everything in such a rapidly changing field where websites are being updated hourly or
faster. I have declined to develop diagrams - even if I could find appropriately licensed
images as a summary because of this rapidly changing nature. Particularly in mechanical
designs because those appear to be the most often altered.

I have expanded considerably what the files are for in open hardware to help better
introduce those outside of the open hardware community to the concept as well as make it
clearer why they need to be included.

I have brought in an example from the PPE community.

In addition, following your recommendation I have attempted to provide some best
practices in terms of ‘when to share’ in the discussion. This is meant to bridge the gap
between the traditionalist and maker philosophies you discuss.

After considering your points and those of the other reviewers about the advantages of less
developed regulatory systems I have simply removed this entirely.

I have corrected the minor mistakes you pointed out – thank you for finding them.

Lastly, I don't think that a complete review of all the legal hurdles and regulatory framework
needed in this technical space can be done here - it needs a completely separate review. We
have a major problem here because even some of the standards are not available. When an
artificial lung company contacted their customers on my behalf to get a testing protocol
they referred me to a long list of ASTM standards which I was not able to acquire through
my relatively-well-resourced University library. I purchased the first one and was
disappointed to find that it mostly contained references to other standards and a shocking
dearth of useful technical information. As there is a conflict of interest between the
publishers of such standards and providing all the information in a single location that is
freely available for designers to help in a pandemic, this represents one of the largest
hurdles. Some good-willed publishers have provided open access to some of their
documents during the pandemic. This is a nice start but a better approach is to use public
funding to provide open access documentation as the cost and opacity of these standards
provide a barrier to open source ventilator development and as I pointed out an
unacceptable state during an emergency.

**Competing Interests:** None
Ramon Farre
Unit of Biophysics and Bioengineering, School of Medicine and Health Sciences, University of Barcelona, Barcelona, Spain

This review by Prof. Pierce is an excellent and timely piece of work. I think the author is to be commended for this panoramic perspective on the current state of the art in the open source field of mechanical ventilators for acute respiratory failure. In addition to subscribing to Prof. Bowman's extensive comments, I would particularly like to emphasize how important it is that open access contributions regarding medical devices be made in the context of multidisciplinary cooperation between professionals from different fields. In fact, unlike many other open access development areas, where the proposed devices or products can be used by a wide variety of users, medical devices focus on very specific end users: the doctors responsible for diagnosing and treating the patients. Therefore, any proposal for open access in medical devices must be not only technically sound and cost-effective, but, most importantly, safe and clinically applicable in real-life clinical routine. Consequently, as clearly stated in Professor Pierce's paper, realistically testing new developments is a critical step. Unfortunately, it is not unusual for some new open access contributions for medical applications to be the result of developments by authors drawn exclusively from the technological field, with the result that the proposed devices exhibit applicability limitations that seem obvious to clinicians who are potential users. Therefore, to advance the extension of the open access approach, it is important to make an effort to actively involve health professionals for both design and testing, trying to publish contributions in medical journals, not only in publications focused exclusively on open access. A very recent work, exactly in the field of developing open access mechanical ventilators for resource-poor areas, shows an example of a team of professionals covering the fields of biomedical engineering, respiratory physiology, physiotherapy and respiratory clinical medicine. This multidisciplinary team has conceived, developed and tested a novel non-invasive pressure support ventilator following the conventional method of evaluating ventilators (bench tests in simulated patients and preclinical evaluation in volunteers) and has published the results in a prestigious journal of pneumology. To end my review of Prof. Pierce's excellent work, I would mention an open ethical issue about the requirements for medical devices to be used in resource-poor areas. It is important that these devices have the same high standard level as those approved for use in developed countries (e.g. CE / FDA marks). But it is also important that until the industry can provide such medical devices at affordable prices for LMICs, patients are not deprived of life-saving therapies. Finding a balance on this ethical-legal issue is difficult but fundamental.

References
1. Garmendia O, Rodríguez-Lazaro M, Otero J, Phan P, et al.: Low-cost, easy-to-build non-invasive pressure support ventilator for under-resourced regions: open source hardware description, performance and feasibility testing. *European Respiratory Journal*. 2020. Publisher Full Text

Is the topic of the review discussed comprehensively in the context of the current literature?
Yes

Are all factual statements correct and adequately supported by citations?
Yes

Is the review written in accessible language?
Yes

Are the conclusions drawn appropriate in the context of the current research literature?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Respiratory mechanics, Respiratory medical devices, Biomedical engineering.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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Author Response 28 Apr 2020

Joshua M. Pearce, Michigan Technological University, Houghton, USA

Thank you for the careful and constructive review.

I have included your well-reasoned point about the need for multi-disciplinary collaboration, the need to involve medical personnel, and to aim to publish in the medical literature.

In addition, I have included a deep review of Garmendia et al. design. A sincere thanks for pointing this out as well as for the work itself.

Thank you for the point about ethics – to account for this I have added several points about this in the discussion.

**Competing Interests:** None

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Reviewer Report 02 April 2020

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Richard Bowman
Department of Physics, University of Bath, Bath, UK
This is a really good, thorough review of open source ventilators, including both a good technical background, a review of existing peer reviewed projects, and a summary of current efforts. I would thoroughly recommend it to anyone considering joining or using an open source ventilator project, as it's particularly good at pointing to some relevant literature that describes the requirements and principles of operation. The conclusion of the article is that we're not there yet - most of the published designs are not sufficiently complete to be easily replicated, while most of the current open projects are not rigorously tested.

The review of peer reviewed articles is interesting and does a good job of rating the different solutions in terms of openness; it is disappointing that these articles don't generally give sufficient information to reproduce the ventilator, but also unsurprising. This lends a great deal of weight to the current move towards more openness in science, where protocols, data, and schematics can be shared in data archives along with papers - but of course that's rarely done retrospectively.

The review of "internet and gray literature" seems objective and reasonable to me, and while such a review cannot possibly stay exhaustive given the frequency with which such projects are appearing, it does seem to cover many of the projects I've heard of. More important than an exhaustive list, however, is the discussion of the common issues to most of the DIY projects - the need for careful testing, quality control, and proper authorisation. Most discussions have focused only on technical validation - but as the author rightly points out, this is not the only way medical devices must be assessed. At least as pressing as the technical challenge is the difficulty of getting new suppliers and new devices through a quality assurance process that gives medical professionals the confidence that they can safely use said devices.

Openness is an important, and often surprisingly contentious, issue. Of the projects that are discussed, only relatively few make available complete designs for their solution. This is particularly surprising in the case of some projects from high-profile institutions that have already been widely reported in the media as "open" while not yet having released any designs. The commonly-accepted practice in open software is that complete designs, including source code and documentation, are made available to the public, and that a project is not considered open until this happens. Similar norms are being established for open hardware projects, supported by organisations such as OSHWA and GOSH.

Given the safety-critical nature of a ventilator, it's reasonable to be reluctant to release untested designs out of a desire to be responsible. Given the time-critical situation, sharing documentation and designs may also be considered lower priority than product development. However, the intent to share a design in the future misses the myriad benefits of open hardware - in terms of scrutiny, feedback, and improvements from the community. It also stifles the development of a community around the design, and there are many cases of promised openness never materialising. My own view is that projects ought not to claim openness until their designs are publicly available under an appropriate license, but there are definitely valid ethical and practical concerns here, and I would welcome an open debate on the best way forward.

The one statement in the article that I'm slightly troubled by is the suggestion that developing countries may be at an advantage due to their less robust regulatory systems. Firstly, while it is true that many countries in the Global South do have less formal economies, their regulations are often very tightly aligned with those in richer nations - for example, the Tanzanian medical device regulations closely mirror those used in the EU. If different standards are adhered to, it may be
because the regulations are not implemented fully, rather than because the government has intentionally applied lower standards. Also, the better-resourced regulatory bodies in rich nations are more able to accelerate the process of approval if needed; it is not clear to me that a medical device would clear the bureaucratic hurdles and achieve approval any faster in a developing country, indeed the process can be much slower. It is also a very thorny ethical issue to trial medical interventions in the Global South that would not pass ethical scrutiny in richer nations, particularly as the interventions are often being proposed by people from said richer nations. I don't think the author is suggesting this, but I do feel it's a point worth highlighting. While there is often an argument made that low quality medical supplies may be better than nothing, it is also reasonable to expect that developers of technology shouldn't do anything to citizens of Low and Middle-Income Countries (LMICs) that they wouldn't do to patients in their own nation. Indeed, most ethical review panels in the UK apply exactly this criterion.

The challenge of creating a safety-critical medical device that can be produced in a distributed manner is significant, and I think the article reflects this. I could not agree more with the statement that "technical validation may not be enough" and would probably go further, to say that technical validation alone is not sufficient to ensure patient safety. While many open ventilator projects now exist and have gathered impressive numbers of volunteers, there remains a significant global challenge to enable such projects to be regulated appropriately, either in the current crisis or longer-term. The existing system of medical device regulation is slow, expensive, and conservative; while this conservatism has its roots in the entirely reasonable desire to prevent harm to patients, the way the system is implemented makes it extremely difficult to certify a medical device without the resources of a large company. Reform of these regulatory systems could enable a more agile approach to the design and manufacture of safety-critical components, but a satisfactory supply chain will also require significantly more quality management than is present in a typical "maker space" run by volunteers, hobbyists, or even experienced engineers. Questions around training and liability are also of paramount importance; while litigation against volunteers acting in good faith seems unduly harsh, there must be accountability in the supply chain of medical devices. Otherwise, we push responsibility onto the clinical staff using uncertified equipment, which adds a crippling burden to front-line staff who are already working at the limit of their capacity.

Overall, I think it's right to keep an optimistic tone, while acknowledging the obvious difficulties associated with the current challenge. It's likely that, while there are many 3D printers available around the world, formal structures that do not yet exist will be needed to enable them to be fully employed to solve supply issues in this and future crises. Whether or not it is possible to make use of community designed and built ventilators in the coming months, I look forward to a world where critical supplies can be designed and produced openly for the common good. If we take the opportunity to put LMICs on a more equitable footing with respect to richer nations, the future may be more inclusive, as well as more resilient.

Is the topic of the review discussed comprehensively in the context of the current literature?
Yes

Are all factual statements correct and adequately supported by citations?
Yes
Is the review written in accessible language?
Yes

Are the conclusions drawn appropriate in the context of the current research literature?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Automated microscopy and open source hardware

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Author Response 28 Apr 2020**

**Joshua M. Pearce**, Michigan Technological University, Houghton, USA

Thank you for the thorough review and kind comments. I agree with you that the state of full open source disclosure in the general literature is disappointing and share your hope that the current move towards open science continues to accelerate.

In this revised version I attempted to pull the grey literature review up to date, but as you point out that is a daily battle and I have cited the work by Read et al., trying to do that on Github.

I share your surprise in the wave of what can only be described as “open washing” where high profile organizations announce their development of an open source ventilator, which upon closer inspection is not open and/or not all of the critical files have been shared. To address this, I have greatly expanded the introductory information and definition of openness as well as provided a few examples.

I have also offered additional thoughts on “when to share” in the Future Work Needed Section.

After considering your points and those of the other reviewers about the advantages of less developed regulatory systems I have simply removed this entirely.

**Competing Interests:** None

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**Comments on this article**

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**Version 2**

Author Response 29 May 2020
Joshua M. Pearce, Michigan Technological University, Houghton, USA

It should be noted that the Gamendia et al. article cited above has been amended and now their easy-to-build non-invasive ventilator is completely open source and licensed under CERN OHL v1.2 and GNU GPL v3. It was published along with an op ed in the European Respiratory Journal by Guerin and Levy that argue the open source approach is something not only feasible but necessary.
https://erj.ersjournals.com/content/early/2020/05/07/13993003.01271-2020

**Competing Interests:** None

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Reader Comment 10 Apr 2020

**Manfred Niehus**, isel/it, Lisboa, Portugal

Good review, Thank you. Please re-check the updated Ref 29 and consider to soften the claim in the article accordingly. stay safe!

**Competing Interests:** No competing interests were disclosed.

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Reader Comment 31 Mar 2020

**Santhosh Kumar Rajamani**, Banas medical college and research institute, India

Even the ventilation protocols used in Ventilators is a proprietary and patented. This leads to confusion and patient injuries notably under ventilation, pneumothorax. Even the ventilation strategy has to be Open sourced. Great idea and best of luck!

**Competing Interests:** None
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