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Abstract

Coronavirus disease 2019 (COVID-19) can lead to respiratory failure. Some patients require extracorporeal membrane oxygenation (ECMO) support. During the current pandemic, health care resources in some cities have been overwhelmed and doctors have faced complex decisions about resource allocation. We present a case in which a pediatric hospital caring for both children and adults seeks to establish guidelines for the use of ECMO if there are not enough resources to treat every patient. Experts in critical care, end of life care, bioethics and health policy discuss if age should guide rationing decisions.

Introduction

The COVID-19 pandemic has forced us to examine questions about the fair allocation of medical resources. This novel coronavirus can cause severe cardiopulmonary failure necessitating life-sustaining therapies, including mechanical ventilation and extracorporeal membrane oxygenation (ECMO). At the time of this writing, there have been 1467 confirmed or suspected COVID-19 patients (920 in North America) placed on ECMO.(1)

While the majority of patients admitted for COVID-19 related illnesses have been adults (median age 49 years), children and adolescents in the US with COVID-19 have also required admission to intensive care units (ICUs).(2) Overall, children who require ICU admission for COVID-19 related illnesses do much better than adult patients. Hospital mortality rate is 4.2% for children and adolescents versus over 50% for adults.(3)

Since most COVID-19 related ICU admissions have been for adult patients, many pediatric ICUs have admitted adults when adult ICUs have been overwhelmed.(4) This has resulted in unique medical and ethical challenges for pediatric intensivists. One such ethical challenge is whether our allocation guidelines for ECMO should prioritize younger patients.
In a recent survey of US pediatric ECMO directors, most respondents (68%) that that pediatric patients should get priority. But a substantial minority (32%) disagreed. (5)

The Case

During the current COVID-19 pandemic, a regional hospital system, which includes a free standing Children’s hospital, assembled a working group of administrators, clinicians, and public health officials to establish guidelines for the allocation of extracorporeal membrane oxygenation (ECMO) support.

The hospital only had 8 ECMO devices, and two trained ECMO teams. The experts were concerned that they might have more patients needing ECMO than they had devices or ECMO teams. The working group wanted to establish guidelines for the fair allocation of ECMO in the event of a shortage.

One group member, a pediatric intensivist, argued that pediatric patients should be given priority over adult patients because they have more years to live. Others in the group objected to this proposal arguing that discriminating on the basis of age fails to uphold the equal dignity of all patients. The policy options were sent to the Ethics Committee for their analysis.

Ryan M. Antiel, MD, MSME and Farr A. Curlin, MD Comment

We find ourselves in a situation where the medical demand outweighs our current supply. What criteria should we use to guide the fair allocation of a scarce resource?
The first criterion is always prognosis. Patients should not get a scarce resource if they are likely to die even if they are treated with the resource. Thus, it is important that we start by asking if a particular patient is likely to benefit from the intervention. Even using prognosis, however, we may end up in situations in which two or more patients with an identical prognosis both need the same scarce resource. What then?

One method is to select patients based on social worth. This approach was used in 1962 by a committee (later known as the “god committee”) charged to develop an allocation system for dialysis at the Swedish Hospital in Seattle.(7) The primary criterion for allocating dialysis was social worth. The committee’s attempts to judge who was worthy of saving proved to be highly subjective and discriminatory. People with families, for example, were judged more worthy of dialysis than people living alone. The Seattle approach was roundly criticized because it implicitly claimed that some lives were worth more than others.

We must ration on the basis of what lives can be saved, not whose life is worth saving. This commitment enacts the solidarity of clinicians with those who are sick, which reinforces public trust in the medical profession.

Another approach to rationing is to use a simple lottery. All patients who meet medical criteria have an equal chance of getting the scarce resource. This option avoids all biases, including those based on age, disability, or other non-medical factors. This approach affirms that all patients share a radically equal and intrinsic dignity simply by being human.(6)

Lotteries have their own problems, however. One still must make judgments about who is likely to benefit from an intervention. Is benefit measured simply as an immediate response to therapy, survival to hospital discharge, or some predetermined length of survival? Each would
implicitly incorporate some values or biases and each would result in resources being allocated differently from the others.

Using the patient’s age is controversial. On one hand, taking age into consideration seems to problematically suggest that young people’s lives are worth more than old people’s lives; it seems to recapitulate the problem of judging social worth. On the other hand, taking age into consideration recognizes a common intuition that pediatric patients have more to lose from death than do, say, eighty-year-olds.

Such moral intuitions would support prioritizing a healthy 12 year old over a healthy 92 year old - not because of their worth or utility, but because of the arc of their lives. The 12 year old has simply lived through fewer life cycles - childhood, young adulthood, middle age, and old age – than the nonagenarian.

Age can be seen as a proxy for the magnitude of loss threatened. There is no invidious discrimination involved. Again, we do not prioritize the young because of their worth or utility but rather because of what ethicist Gilbert Meilaender terms *generativity* - “the virtue that makes us ready, even eager, to produce those who will replace us and to sacrifice ourselves on their behalf.”(8) We should not grasp for more life indefinitely but instead should seek to cultivate a gratitude for the gift of life and a willingness to let the next generation take our place.

These considerations lead to a three-part rationing approach. First, we must consider prognosis and whether a patient will benefit from ECMO. We need to consider the severity of their current illness, as well as whether the patient has significant comorbidities for which we would expect the patient to not survive even until hospital discharge. Second, after assessing the likelihood that a particular patient may or may not benefit from ECMO, age ought to be
considered as a tie-breaker. Younger patients have more to lose when the arc of their life is threatened to be cut short and thus they have a greater claim to treatment on the basis of need.

Finally, when two patients, who both stand to benefit from ECMO are roughly the same age, we can resort to a random lottery.

We should keep in mind that, if a patient is ill enough to need ECMO, the prognosis is poor for young and old alike. Perhaps in this situation the ethically important questions lie elsewhere. First, a policy of rationing that considers age as a criterion can only be morally tolerable if we have other means of caring for the elderly besides ECMO and mechanical ventilation. The institutions of hospice and palliative medicine provide a helpful alternative when technology can no longer restore the patient to health.

Furthermore, as Paul Ramsey argues in *The Patient as Person*, before we even get to questions about who will live and who will die, we must first address fundamental questions about why we as a society prioritize exotic medical treatments when the most vulnerable among us do not have access to basic care. The current pandemic has again exposed significant health disparities and mis-allocations of resources that have always been latent in our society.

Bernard Williams wrote that “moral conflicts are neither systemically avoidable, nor all soluble without remainder.” We acknowledge no scheme is without shortcomings, without a moral remainder. Leading a moral life in a pandemic cannot be reduced to a moral calculus, but by its nature is a practice in which dispositions and discernment are cultivated as we once again learn what it means to care even when we cannot cure. Thus, we must seek ways to support the physicians and nurses who can experience significant moral distress while caring for these patients, especially if they must make resource allocation decisions.
Govind Persad, JD, PhD, and Douglas B. White, MD, MAS, Comment

With appropriate precautions, it is legal and ethical to prioritize pediatric patients as one part of an allocation system for scarce, life-saving medical resources.

Ethical justification for prioritizing young patients

The ethical justification for prioritizing pediatric patients is not grounded in value judgments about older individuals’ social worth or quality of life. Rather, it aims to avoid the distinctively bad outcome of dying before one has had the opportunity to live through life’s stages. Living for only 15 years before having one’s life cut short by Covid-19 is an enormously worse outcome than dying of COVID-19 at age 75. On one view, this allocation criterion gives priority to the worst off, in the sense that the young have had the least opportunity to live through life’s stages. On another view, this allocation criterion is egalitarian, in the sense that it seeks to diminish disparities in individuals’ opportunity to live through stages of life.(11) This approach is consistent with recognizing the equal dignity of all patients regardless of age.

It is important to distinguish a priority for pediatric patients, which we believe is ethically sound, from a more general priority for any patient with a longer life-expectancy. The problem with the more general argument is that it runs the risk of disadvantaging people who have already been disadvantaged by their life circumstances. People living in poverty, for example, have a lower life-expectancy than people living in affluence. That doesn’t mean we should prioritize the affluent, even though they are likely to live longer.
In the presented case—and in most cases—pediatric priority will reduce the odds that someone will die before living through life’s stages while also saving more years of life. But these considerations can come apart in ways that reveal the importance of priority to the worst off (i.e., pediatric patients) even when more years of life would not be saved. Consider a 12-year-old with cancer and a 90 year-old with cancer, both of whom are expected to die within 5 years. Although their life expectancies are similar, the pediatric patient is substantially worse off in that she has had the least opportunity to live through life’s stages. When saving more years of life aligns with prioritizing the worst off, this can obviate ethical concerns we might otherwise have about considering long-term life expectancy. Unlike the shorter life expectancy often associated with poverty or disability, the shorter life expectancy associated with older age does not stem from disadvantage or social injustice. Rather, it is a consequence of past advantage—the desirable outcome of already having experienced many years of life.

Giving pediatric patients some priority in access to treatment appeals not only to the value of equality of opportunity to live a complete life, but also to the value of reducing disparities. The life-shortening effect of social inequality means that people from groups who are subject to various forms of disadvantage and social injustice are less likely to live a long life. They are, today, overrepresented in younger age groups and underrepresented in older groups. For example, 38% of all Hispanics and 31% of all Blacks in the United States are under the age of 22 years compared to 23% of all Whites.(12)

Recognizing that it can be ethical to consider age as part of a multi-principle allocation system does not entail accepting that it would be ethical to only consider age. Other values, like a patient’s prognosis for hospital survival and near-term survival, essential worker status, or
disadvantages they have experienced all matter too. These considerations might outweigh pediatric priority.

Numerous US states have adopted as part of their Crisis Standards of Care (CSCs) a multi-principle allocation framework that gives some priority to the young, either as a tie breaker or as one factor that is part of a point system. A reserve system, which gives priority to pediatric patients for some but not all available ventilators, is another way of incorporating pediatric priority into a multi-principle approach.

**Legal permissibility of including age in allocation framework**

Considering age is not only ethically defensible but, contrary to some misunderstandings, legal. In constitutional law, age-based criteria need only to have a “rational basis.” Rational basis scrutiny only requires that a law or policy that considers a given characteristic—such as age or wealth—do so in a way that is rationally connected to the law’s goals, This allows ample room for decision makers to consider age. This contrasts with the legal scrutiny given to policies based on race or sex. They are subject to the more exacting legal standards of “intermediate” or “strict” scrutiny. Those require that the law in question use the characteristic in a way that is precisely tailored to its objectives. For instance, under current Supreme Court precedent, universities with race-based affirmative action programs must show that their use of race is strictly necessary to achieve objectives that are themselves crucial to higher education goals, whereas universities only need a rational basis for employing affirmative action programs based on a family’s geographic location, wealth, or income.
The law regards age differently from other identity characteristics in part because it recognizes that age inevitably changes over time. While the Age Discrimination Act of 1975 goes beyond constitutional law in regulating the use of age, the statute differs sharply from race or sex discrimination statutes. In particular, it—unlike race, sex, disability, or religious discrimination statutes—exempts from review age criteria explicitly enacted in federal, state, or local law or ordinance. If a state or locality passed a law or ordinance explicitly authorizing the use of age in CSCs, that law would be exempt from Age Discrimination Act review. It also permits the use of age as a proxy for other factors that are part of the normal operation or statutory objectives of a program—objectives which might include extending lives or narrowing health disparities—when age is more practical to measure directly. And the Age Discrimination Act explicitly presumes that programs giving children special priority are legal unless such priority excludes others from participation.

Age Discrimination Act claims have to proceed through mediation and other steps before entering the court system. Thus, the Age Discrimination Act has generated little case law. Nonetheless, federal agencies have interpreted the Age Discrimination Act to permit the use of age as one among several factors in multi-principle organ allocation policies, such as kidney allocation policies that allocate older kidneys to older patients. The U.S. Department of Health and Human Services (HHS) currently interprets the Age Discrimination Act to prohibit “a policy that automatically disallows heart transplants to all individuals aged 65 or older”—assuming that the policy was not explicitly adopted in federal, state, or local law and therefore exempt—but recognizes that “it would be permissible for the transplant center to consider age as one factor in assessing the allocation of transplants.” Most recently, HHS reviewed
Pennsylvania’s CSC after a complaint was filed, and permitted the framework’s use of “stage of life” in a multi-principle framework to stand.(25)

Public attitudes about prioritizing the young

In light of the ethical and legal permissibility of incorporating life cycle considerations into triage during a pandemic, it is also relevant to consider public attitudes. In a multi-year structured deliberative democratic engagement with Maryland residents, a substantial majority of participants supported the use life cycle considerations in ventilator allocation frameworks during a pandemic.(26) Similarly, a large study of public opinions allocating scarce organs for transplant revealed support for prioritizing young patients.(27)

Pediatric prioritization aligns with the World Health Organization’s suggestion that allocation “should rely on broad life stages, rather than ranking individuals based on differences of only a few years.” Carl Coleman has similarly suggested that “giving a bump up in priority to broad categories like ‘adolescents’ and ‘young adults’ may pose less of a risk of fostering biases against the elderly than a sliding scale approach in which each additional year of life is treated as a negative.”(28)

Cathy Zhang, BA; Aaron Glickman, MPA; and Ezekiel J. Emanuel, MD, PhD, Comment
Frameworks for allocating scarce medical resources require multiple principles. Any individual principle is insufficient on its own because it ignores some morally relevant factors.

Any individual allocation principle is flawed and therefore insufficient. For example, allocation on the basis of “first-come, first-served” is morally flawed because it usually benefits the wealthy, well-connected, and people who by chance become ill earlier. This principle should be excluded from allocation frameworks.(29, 30)

Similarly, many in medicine believe that allocation of scarce resources should favor the sickest patients.(31) Like first-come-first served, the sickest-first principle is flawed. It ignores prognosis, favors treatment when only minor improvements are possible, and focuses how sick someone is at a current time, at the expense of people who are likely to become just as sick in the future. (32-34)

We propose a multi-principle framework for allocating resources, previously described as the complete lives system.(35) By this approach, the prioritization of youngest patients proposed by the pediatric intensivist is insufficient by itself, but such prioritization could be incorporated into an ethically defensible multi-principle framework.(36)

For scarce life-saving medical interventions, priority should go to patients who could die, but are much more likely to live with the intervention, rather than those who are likely to die even with the intervention or likely to live regardless of receiving it. While there are technical problems in assigning patients to groups, we will assume that it is possible on clinical grounds and reasonably accurate. If there are more patients likely to benefit than there are available ECMO machines to treat them all, the complete lives system should be used for allocation.
The complete lives system aims to allow more people to live complete lives by prioritizing younger patients. However, there are legitimate deficiencies to youngest-first allocation. The complete lives system therefore uses a modified youngest-first approach and incorporates 4 other considerations: prognosis, saving the most lives, lottery (i.e. equity), and prospective instrumental value during a public health crisis.

Priority to the worst off: modified youngest-first

A widely accepted ethical principle is that the “worst off” should receive allocation priority. In medicine, a natural way of thinking about the worst off is as those people who will have a premature death and therefore not be able to live a complete life.(37, 38) The complete lives system thus distinguishes people on the basis of not having something everyone thinks is valuable—a long, complete life. This is a subtle but important distinction from other forms of youngest-first allocation.

Unlike race or sex, age is morally relevant because all people age. Every elderly person has been young. Not prioritizing young people denies them the ability to become old. Therefore, to give life-saving resources to the young does not discriminate against the old on the basis of ageist stereotypes or value of life judgements. It merely recognizes that we all value a complete life.

The complete lives system also considers the value of life lived so far. The death of a 22-year-old, who has developed an identity and attachment to their future through personal and professional projects, built relationships, and received investment from the community, is more tragic than that of a 2-year-old old.(39) This is a widespread moral intuition.(40) Accordingly,
the complete lives system prioritizes adolescents and young adults over infants. This modified youngest-first approach differs from a strict youngest-first approach that would prioritize the 2-year-old over the 22-year-old.

*Maximizing total benefits: prognosis, saving the most lives, and extending life-years*

In addition to saving more life-years, there is value in each individual life saved. Therefore, survival prognosis is a morally relevant consideration. Considering prognosis allows for maximization of benefits by saving the most lives. In cases where patients have similar prognoses with treatment, number of life-years can be a consideration.

Prognosis and life-year considerations raise objections from disability advocates who worry that considerations of life-years will lead to discrimination against people with disabilities whose life-expectancies may be shorter than those of people without disabilities. These objections and concerns are valid. Patients may not be excluded solely on the basis of having a disability. Nor should triage decisions consider long-term life expectancy and quality of life judgments. Because those factors are difficult to predict individually, they are easily subject to bias and abuse and are therefore flawed.

However, these concerns can be addressed without ignoring either short-term prognosis and age. Short-term life expectancy is generally predictable and ignoring it will produce immoral outcomes such as giving equal priority to previously healthy patients and to patients with advanced terminal illnesses. The relevant time range considered for short-term prognosis will need to be determined by a working group; one option is to consider 5-year prognosis, as is done for liver transplant allocation.
Maximizing benefits during public health emergencies: instrumental value

In the context of COVID-19, special considerations apply in the complete lives system. Essential health care workers and participants in clinical research should be prioritized. These patients are not privileged because of inherent moral worth. Rather, prioritizing them is consistent with values of maximizing benefits. Saving additional health care workers will allow for more lives to be saved in the future. Prioritizing research participants will yield useful clinical insights that can in turn save lives and efficiently allocate resources—thereby reducing scarcity.

Treating people equally and respecting dignity

There may be edge cases in which two patients similar in all morally relevant ways may require ECMO machines, but supply is limited. In these circumstances, the equality of persons can be upheld with a random lottery. While allocation by lottery is on its own insufficient because it ignores other relevant principles such as prognosis, it can ethically be used as a tie-breaker. Furthermore, there is no ethical basis for giving COVID-19 patients priority over patients without COVID-19. Allocation decisions should be made by an independent ethics body, not frontline health care workers. This will relieve physicians of the emotional burden of triage. Most importantly, it will promote consistent, equitable decisions.

In summary, the pediatric intensivist’s proposal for a youngest-first allocation framework on the basis of saving more life-years is ethically defensible. The claims that it discriminates on the basis of age and does not respect the dignity of all patients are not persuasive. Life-years must be considered, and it is possible to do so while respecting the dignity of all patients.
Considering life-years maximizes opportunities for all patients to live complete lives. The correct approach is to consider age and incorporate it into an allocation algorithm that also includes the other ethical imperatives noted above.

**John D. Lantos, MD, Comments**

Decisions about allocating scarce life-saving resources are brutal. All the choices are bad. But some are worse than others. And some are illegal. We must not discriminate on the basis of social worth, race, disability, gender, wealth, or fame. That sort of fairness will be difficult to achieve but it is essential. We must scrutinize policies to make certain that they don’t implicitly incorporate considerations that would systematically lead to such discrimination, considerations like a first-come, first-served allocation plan. But, as all three commenters show, broad consideration of age is different, both ethically and legally. We were all young once. If we are lucky, we will all grow old. We all deserve to live a full life and to experience all of life’s stages. Thus, prioritizing younger patients for scarce resources is not just ethically permissible; it is the epitome of fairness.

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