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When to lock, not whom: Managing epidemics using time-based restrictions



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1. Introduction

ABSTRACT

We present novel policy tools to manage epidemics. Rather than using prevalent population restrictions, these policy strategies are based on cyclical time restrictions. Key findings on the outcomes of such policy strategies include: a significant improvement of social welfare, substantially lessening the trade-offs between economic activity and health outcomes; optimally-derived timings of interventions are shown to suppress the disease, while maintaining reasonable economic activity; and outcomes are superior to the actual experience of New York State and Florida over the course of 2020.

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million; the declines in U.S. GDP and consumer expenditures for 2020 have been -3.4% and -3.8%, respectively.

The COVID19 pandemic has created a global health and economic crisis of a magnitude not experienced since the Great Influenza Pandemic of 1918-1919. After 33 months, about 630 million people have become infected, about 6.6 million people worldwide have died, and estimates of excess deaths are more than three times as high¹; IMF (2022) reported a 3.1% drop in world GDP in 2020, and a 4.5% drop in the advanced economies. The death toll in the U.S. is almost 1.1

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URL: https://www.yashiv.sites.tau.ac.il/ (E. Yashiv).

¹ See https://ourworldindata.org/covid-deaths.

We address the issue of policy responses to the pandemic, providing an analysis of new time-based tools to manage epidemics. These policy strategies were proposed in epidemiologically-grounded work by Karin et al. (2020). The contribution of our paper is the economic analysis of these new policy tools. The proposed policy consists of alternating periods of work and lockdown, at pre-defined frequencies, for the entire population. We present both normative and positive analyses. The former applies to future pandemics or epidemics while the latter evaluates policy against real world benchmarks in the U.S. using data.

The proposed policy tools are particularly relevant in light of the difficulties experienced by policymakers in finding a policy strategy that lessens the trade-offs involved. In theory, targeted population lockdowns could constitute "fine tuning" of lockdown measures, which would serve to lessen any economic cost. In practice, however, it turned out to be challenging to identify sub populations to be allowed unrestricted economic activity, while imposing restrictions on other population groups. Political and moral issues, as well as practical implementation issues, have come into play. The time-based public health management policy avoids these difficulties, taking time, rather than population, as the medium of restrictions.

Our model lies within a line of COVID19 research in Economics, which posits a planner problem. The planner formulates an optimal policy of Non-Pharmaceutical Interventions (NPIs), and in particular, lockdowns, subject to a model of disease dynamics, taking into account vaccine arrival. The planner trades off the costs of public health outcomes, such as breach of ICU capacity and death, with the economic costs of suppression policy, including declines in production. Prominent contributions in this line of literature include Acemoglu et al. (2021), Alvarez et al. (2021), Brotherhood et al. (2021), Brotherhood and Jerbashian (2020), Farboodi et al. (2021), and Jones et al. (2021). Our analysis pertains to new policy strategies, which were not considered in the literature.

We highlight five dimensions related to time:

(i) From the normative perspective, we analyze policy that relies on *time restrictions*. Such policy is an alternative to policy based on restrictions of sectors, age groups, regions, or other targeted population groups. The success of the policy hinges on two mechanisms that operate over time. In lockdown days, the disease is suppressed through exogenous government-imposed restrictions. In work days, the disease is suppressed through the endogenous rational behavior of the population. This combination allows the effective reproduction number to be kept at around 1 on average. Notably, a full, prolonged lockdown is not needed to keep death rates low. All that is needed is to have the open periods short enough, backed up by rational precaution of the population, and followed by lockdown periods of reasonable length ahead of the next re-opening.

(ii) The epidemiological rationale for the proposed policy is, inter alia but not exclusively, based on the *timescales of virus transmission*. The idea is that for every 14 day period, there will be *k* days of work and 14 - k days of lockdown. This number, *k*, uses the timescales of the virus against itself, taking into account a latent period after exposure, whereby the infected person does not infect others. In other words, the epidemiological cost of open days is mitigated by the fact that individuals do not become infectious immediately. The epidemiological benefit of lockdown days is enhanced because the schedule locks down the economy when individuals are at their most infectious.

(iii) Using an optimizing social planner model, the control variables for this policy are *the timing of the various measures*-initial lockdown, the cyclical policy phase, and release. Hatchett et al. (2007) highlight the idea that imposing NPIs early in an epidemic can significantly reduce mortality. In the current paper, the exploration of timing issues, both start time and duration, are at the heart of the analysis.

(iv) This policy is compared to a prevalent policy path which sets lockdown and release as *functions of disease prevalence*, *which is time-varying*. Specifically, the latter uses trigger thresholds, such as the number of persons hospitalized in ICU in a given period of time, and gives rise to the pattern of recurrent lockdown and release observed in the U.S. and other countries since the start of COVID19.

(v) The proposed policy is subsequently compared to the actual experience of New York State (NYS) and Florida. The outcomes observed for NYS turn out to depend crucially on *the timing of the policies undertaken;* those for Florida reveal particular policy preferences.

We explicitly model the dynamic path for the reproduction parameter, reflecting both rational individual behavior and the effects of lockdown policy. Thus, we model the endogenous response by individuals, who adjust to the new environment and behave accordingly. Additionally, government interventions in the form of lockdowns induce behavioral change. These elements are included in a model whereby individuals maximize utility and a social planner maximizes a welfare function, which incorporates individual utility. We then take the model to the data and use the resulting estimates to calibrate the relevant parameters, which we subsequently use in simulations.

The social planner in our model uses a PDF in order to form ex-ante expectations of vaccine arrival time. This serves three key roles; first, it sets the horizon for the problem, acting as a rate of leaving the state of the pandemic. Second, it is an expression of the risk and uncertainty embodied in the planner problem. Third, relative to the interest rate, it plays the major quantitative role in discounting future streams. In our simulations vaccine arrival is realized after 540 days. Using these elements, our analysis quantifies outcomes in terms of social welfare. We simulate the optimal cyclical policies and examine their health and economic implications.

The cyclical policy is compared to four non-cyclical benchmarks: two polar cases, of no policy intervention (i.e., no lockdown) or full lockdown till vaccine arrival; a single time span lockdown policy, whereby the starting date and the duration are chosen optimally; and a theoretical path trying to mimic a real-world rationale, whereby the planner chooses thresholds for multiple lockdowns in terms of the critically ill. Subsequently, and additionally, we evaluate the time-based policy tools in relation to the actual experience of NYS and Florida in 2020.

We plot the policy plans in terms of a policy possibilities frontier. These show the outcomes of optimal planner policies, using a two-dimensional graph of the death toll per 1 million people and the value of lost output, in annual GDP terms. Movement along the frontier occurs as the policy instrument in use changes, or as the weight assigned to fatalities in the planner objective function changes, or as the lethality of the disease changes.

Our analysis yields the following key findings.

First, the policy instruments, based on time restrictions, provide for significant improvement, substantially lessening the trade-offs involved relative to the four non-cyclical benchmarks. The latter are situated in points on the graph beyond the frontier.

Second, we quantify social welfare costs. These are given in utility terms and in Present Discounted Value (PDV) consumption terms. Using the consumption loss measure, comparing actual consumption to the pre-COVID steady state (formally defined below), the different cyclical strategies place these losses at 22% to 26%, the no intervention policy results in 36%, full lockdown in 29%, optimal lockdown duration in 27%, and the thresholds strategy in 25%. The underlying rationale for the improvement is that cyclical strategies allow the planner to achieve similar death tolls with fewer lockdowns, or to reduce the death toll dramatically without a significant damage to output. These results are due to the optimally-derived timings of intervention (for example, "front loading" interventions is beneficial in specific cases, which are spelled out) and the ability of the cyclical strategies to suppress the disease while maintaining a reasonable level of economic activity.

Third, using daily data from March 2020 to early 2021, optimal cyclical policies fare much better than actual experience in the states of New York and Florida. When deriving this result, we use estimated state-specific parameters in the simulations.

Importantly, the benefits of the time-based policy tools that we find are likely to be a lower bound of their true advantage over policy strategies that have been implemented. This is so because, for tractability, we are not giving the planner full flexibility when applying the cyclical tools. Similarly, we do not quantify additional benefits, such as predictability of production, gains in non-COVID health matters, transparency, ease of communication, and fairness.

We note that the idea of a cyclical strategy, which is at the focal point of the normative analysis of our analysis, has been brought to the attention of policymakers (see Yashiv (2020), Alon and Yashiv (2020), and Alon et al., 2020) and has been considered or implemented by a host of firms and educational institutions in the U.S., in Europe, and in Latin America. Online Appendix A provides elaboration.

The paper proceeds as follows: Section 2 discusses the model, including the policies we propose, on which we further elaborate in online Appendix A. Section 3 presents the calibration and the solution methodology. Section 4 presents the results. Section 5 explores the underlying mechanism. Section 6 examines the relation between the model planner solution and actual outcomes in two U.S. states – NYS and Florida. Section 7 concludes. Online Appendices B and C provide further elaboration.

2. The model

We model an optimizing social planner who operates within a SEIR model of the epidemic and a model of the macroeconomy with optimizing agents responding to the epidemic. We elaborate on the novel policy strategies based on time restrictions.

2.1. The evolution of the epidemic

We analyze the evolution of the epidemic in two complementary blocks - infection transmission and clinical progression.

2.1.1. The infection transmission block

The infection transmission block is characterized by the SEIR-Erlang model, reflecting the epidemiological properties of COVID19. The model is essentially based on the seminal contribution of Kermack and McKendrick (1927). Its present form is discussed in Champredon et al. (2018). See Bar-On et al. (2021) for a more detailed analysis, where we explain the need for the two complementary model blocks. Before contacting the disease for the first time, a person is Susceptible (S). Once a person gets infected, disease progression is split into distinct compartments – Exposed (E), Infectious (I), and Resolved (*R*). We denote by β_t the infections transmission rate, σ , the transition rate from *E* to *I*, and γ , the transition rate from I to R. An infected individual spends some time in each compartment before moving on to the next one. The person is infectious only when in the I compartment, but not when residing in the preceding E compartment. The time durations spent in the E and I compartments are known as the latent and infectious periods, respectively. Once people move to the Resolved stage, they no longer participate in disease transmission. With Poisson transition rates between compartments, the residence times in each of them are distributed exponentially, and thus have zero mode. Exponential distributions capture the mean but not the mode of the biologically accurate distributions of residence times, because in reality what most people spend in each stage is close to the mean of the distribution, rather than zero. Therefore, we split the E and I compartments into two sub-compartments and double the rate of transition. Now, the latent and infectious periods are the sum of the time spent in the E_1 and E_2 or I_1 and I_2 sub-compartments, respectively. Their distribution is the sum of exponentially distributed random variables, a special case of the Gamma distribution, known as the Erlang distribution. The means of Erlang distributions remain $1/\sigma$ and $1/\gamma$, but the modes are now near the means, as they should be. In the remainder of the paper we shall refer to this model as the SEIR model, without noting the number of sub-compartments.



a. The SEIR-Erlang block



b. The Clinical block

Fig. 1. The Model.

Graphically, this block is presented in panel a of Fig. 1.

The following equations describe this block. Throughout, all stock variables are expressed as a fraction of the population.

$$S_t = -\beta_t (I_{1t} + I_{2t}) S_t \tag{1}$$

$$\dot{E}_{1t} = \beta_t (I_{1t} + I_{2t}) S_t - 2\sigma E_{1t}$$
⁽²⁾

 $\dot{E}_{2t} = 2\sigma E_{1t} - 2\sigma E_{2t}$ (3)

$$\dot{I}_{1t} = 2\sigma E_{2t} - 2\gamma I_{1t} \tag{4}$$

$$\dot{I}_{2t} = 2\gamma I_{1t} - 2\gamma I_{2t} \tag{5}$$

$$\dot{R}_t = 2\gamma I_{2t} \tag{6}$$

An important parameter is the reproduction number \mathcal{R}_t , which is the average number of people infected by a person, and is given by:

$$\mathcal{R}_t = \frac{\beta_t}{\gamma} \tag{7}$$

We use \mathcal{R}_t for the reproduction number at date t and denote the basic reproduction number by \mathcal{R}_0 at the initial stage, when $S_0 = 1$. Beyond the initial t = 0, our formulation will allow for \mathcal{R}_t to be affected by policy and by rational individual behavior, as elaborated below in sub-section 2.2. We shall also be discussing the effective reproduction number, defined as:

$$\mathcal{R}_e = S_t \mathcal{R}_t \tag{8}$$

We model the transmission rate β_t as a function of three arguments, using the following formulation:

$$\beta_t = \beta_W - \beta_N \left(1 - \frac{N_t}{N^{SS}} \right)^{\alpha} + \beta_\Lambda \exp\left(-\Lambda t\right)$$
(9)

The arguments are motivated as follows:

a. β_W is the transmission rate when the economy is open, i.e., production and employment are not restricted. b. β_N parameterizes the scale of the decline in transmission as activity falls (decline in $\frac{N_t}{N^{SS}}$, employment relative to its steady state), using a power function with parameter α .

c. $\beta_{\Lambda} \exp(-\Lambda t)$ expresses the decline in transmission due to learning over time by individuals. At time t = 0, when $\frac{N_t}{N^{SS}} = 1$ we get:

$$\beta_0 = \beta_\Lambda + \beta_W \tag{10}$$

which is the transmission at the initial stage and corresponds to $\mathcal{R}_0 = \frac{\beta_0}{\nu}$.

After a period of time, which depends on the rate of decline Λ , individuals change their behavior, and when exp $(-\Lambda t) \ll \beta_W$ we get that the transmission rate (and hence \mathcal{R}_t) rises with employment:

$$\beta_t = \beta_W - \beta_N \left(1 - \frac{N_t}{N^{SS}} \right)^{\alpha} \tag{11}$$

2.1.2. The clinical block

The clinical block describes the clinical progression of the disease and the progression of new cases through the healthcare system, depending on the development and severity of symptoms. We postulate the following. Once infected, a person enters an incubation period, a *P* state, during which there are no symptoms, lasting for $1/\theta_P$ on average. Following it, a person either remains asymptomatic (*O*) or develops symptoms (*M*). Denote the share of asymptomatic cases by η . The others $(1 - \eta)$ develop symptoms, and with probability ξ are hospitalized (*H*). A given share π of patients become critically ill (denoted *X*), i.e., develop conditions requiring transition to ICU. Following the literature (see, for example, Kaplan et al. (2020) and Brotherhood et al. (2021)), we specify the death probability in this critical state *X* as:

$$\delta(X_t) = \delta_1 + \delta_2 \cdot \frac{\mathbf{I}(X_t > \overline{X}) \cdot (X_t - \overline{X})}{X_t}$$
(12)

where \overline{X} denotes ICU capacity and \mathbf{I} is the indicator function. The reasoning is as follows. When there is no breach of ICU capacity, the death probability is given by δ_1 . Whenever there is an overflow, $\mathbf{I}(X_t > \overline{X}) = 1$, the death probability increases with the risk that a patient will not be provided ICU care when needed. The underlying assumption is that the allocation to ICU is random among all patients in need of it, so the risk to be left out of ICU (for a given patient) is given by $\frac{(X_t - \overline{X})}{X_t}$. At the limit, the death probability $\delta(X_t \gg \overline{X})$ is given by $\delta_1 + \delta_2$.

At any stage, a person may recover (C). The clinical block is represented graphically in panel b of Fig. 1.

The analytical description of the symptomatic branch is:

$$P_{t} = \beta_{t} (I_{1t} + I_{2t}) S_{t} - \theta_{P} P_{t}$$
(13)

$$\dot{M}_t = (1 - \eta)\theta_P P_t - \theta_M M_t \tag{14}$$

$$\dot{H}_t = \xi \cdot \theta_M M_t - \theta_H H_t \tag{15}$$

$$X_t = \pi \theta_H H_t - \theta_X X_t \tag{16}$$

$$\dot{D}_t = \delta(X_t)\theta_X X_t \tag{17}$$

The parameters θ_P , θ_M , θ_H , and θ_X relate to the average time that passes between the stages of infection, symptoms onset, hospitalization, ICU admission, and death, respectively.

Note that a given person moves through the two blocks simultaneously. They relate to two timescales, the infectiousness profile and the clinical progression, which develop in parallel. Each of the blocks characterizes different properties of an infection case: one identifies whether a given person is infectious and the other identifies the severity of the disease (whether one needs hospitalization, for example). Thus, a person might be infectious (stage I of the infection transmission block) and at the same time still show no symptoms, i.e., be in stage P, the incubation period, of the clinical block. Similarly, a person may no longer be infectious (stage R of the infection transmission block) but still be hospitalized (stage H of the clinical block).

2.2. The economy

We first describe optimal individual decisions (2.2.1) and then elaborate on the aggregate outcomes (2.2.2).

2.2.1. Consumer-worker optimization

In each period, there are two groups of individuals in the economy: the majority who are active and a minority who are inactive because of health reasons.

Active workers. Active workers maximize a utility function, which includes standard utility from consumption and disutility from labor; there is no saving/investment. Additionally and importantly, they respond to the epidemic endogenously. The model is thus related to papers which tie macroeconomic dynamics to epidemiological dynamics, which posit that individual rational economic behavior has two-way connections with disease transmission.² Notable COVID-related contributions include Atkeson (2021, 2022), Atkeson et al. (2021), Eichenbaum et al. (2021), Garibaldi et al. (2020), and Krueger et al. (2020). An elaborate analysis, emphasizing heterogeneous agents, is offered by Kaplan et al. (2020).

² This issue was explored long before COVID19; prominent examples include Geoffard and Philipson (1996), Fenichel et al. (2011), and Fenichel (2013), Greenwood et al. (2019), and the surveys by Philipson (2000) and Verelst et al. (2016).

Inactive workers. This group is comprised of the following pools (shares of the population): D_t , deceased; X_t , critically ill; H_t , hospitalized; and ϕM_t , workers with COVID19 symptoms, who do not work because they are isolated. The individuals in these pools, apart from the deceased, get a fixed consumption transfer from the government. This formulation follows a modeling idea of Kaplan et al. (2020, see page 28); it assumes that the government has access to a storage technology for final goods and has accumulated a sufficient quantity of goods to provide for the consumption of the afore-cited pools during the pandemic. As a result, their consumption is not part of the flow resource constraint. This assumption simplifies the simulation exercise and has no bearing on its quantitative results as these pools are negligibly small throughout relative to the active population.

The utility function of the active workers is given by:

$$U(c_t, n_t, \dot{D}_t) = u(c_t) - \frac{1}{\omega^n(\dot{D}_t, t)} v(n_t) - \ln \omega^x(\dot{D}_t, t)$$
(18)

We define and explain each term in turn:

(i) For consumption we postulate a conventional functional form and parameter value.

$$u(c_t) = \frac{c_t^{1-\zeta_c}}{1-\zeta_c} \tag{19}$$

where ζ_c is the constant relative risk aversion parameter, which we set to 1.

(ii) For labor we postulate a conventional functional form and parameter value.

$$v(n_t) = \theta^{1+\zeta_n} \frac{n_t^{1+\zeta_n}}{1+\zeta_n} \tag{20}$$

and ζ_n is the inverse Frisch elasticity of labor supply, set to 4, in line with the synthesis of micro evidence reported by Chetty et al. (2013), pointing to Frisch elasticities around 0.25.

This conventional term is multiplied by $\frac{1}{\omega^n(\dot{D}_{t,t})}$ capturing the individual response to the epidemic, in the spirit of modeling by Kaplan et al. (2020, see their eqs. (21)-(22)). It expresses an increase in the marginal disutility of labor due to the social interactions component of market work in the times of the epidemic. It is a function of the current daily death rate \dot{D}_t , and we assume $1 > \omega^n(\dot{D}_t, t) > 0$, $\omega_{ss}^n = 1$, and $\frac{\partial \omega^n(\dot{D}_t, t)}{\partial \dot{D}_t} < 0$. (iii) The last term in equation (18) offsets the level shift in utility generated by the marginal utility shifter $\frac{1}{\omega^n(\dot{D}_t, t)}$. We

(iii) The last term in equation (18) offsets the level shift in utility generated by the marginal utility shifter $\frac{1}{\omega^n(\dot{D}_{t,t})}$. We assume $1 > \omega^x(\dot{D}_t, t) > 0$, $\omega_{ss}^x = 1$, and $\frac{\partial \omega^x(\dot{D}_{t,t})}{\partial D_t} < 0$ so that this term, $-\ln \omega^x(\dot{D}_t, t)$, is positive and increases in \dot{D} . This offset follows Kaplan et al. (2020) when computing social welfare. The offset term precludes a big fall in one's own individual utility (and consequently in social welfare which aggregates individual utility) when others die, while the individual undertakes precautionary behavior. It can be interpreted as capturing the added value of precautionary behavior, i.e., the value to the individual herself or himself from engaging in precautionary behavior in the face of the death of others.

Under these restrictions, the utility function becomes:

$$U(c_t, n_t, \dot{D}_t) = \ln c_t - \frac{1}{\omega^n(\dot{D}_t, t)} \theta^5 \frac{n_t^5}{5} - \ln \omega^x(\dot{D}_t, t)$$
(21)

The lifetime utility maximization problem of active workers, given the individual discount rate r, is given by:

$$\max_{\{c_t, n_t\}} \int_{t=0}^{\infty} e^{-rt} U(c_t, n_t, \dot{D}_t) dt = \max_{\{c_t, n_t\}} \int_{t=0}^{\infty} e^{-rt} \left[\ln c_t - \frac{1}{\omega^n (\dot{D}_t, t)} \theta^5 \frac{n_t^5}{5} - \ln \omega^x (\dot{D}_t, t) \right] dt$$
(22)

$$s.t. \quad c_t = wn_t \tag{23}$$

Using the budget constraint, the F.O.C implies that individual labor supply of utility-maximizing agents can be written as follows:

$$n_t = \frac{\omega^n (\dot{D}_t, t)}{\theta}^{1/5} \tag{24}$$

We assume the following functional form for the marginal utility shifter $\omega^n(\dot{D}_t, t)$:

$$\omega^n(\dot{D}_t, t) = \left(\omega^x(\dot{D}_t, t)\right)^5 \tag{25}$$

where $\omega^{x}(\dot{D}_{t}, t)$ is defined to be

$$\omega^{\chi}(\dot{D}_t, t) \equiv (1 - g(\dot{D}_t, t)), \quad g(\dot{D}_t, t) < \overline{g} \quad \forall t.$$
(26)

The function $g(\dot{D}_t, t)$ depends on the daily death flow and on time and is bounded from above by \overline{g} . We discuss the restrictions on $g(\dot{D}_t, t)$ and its functional form below. We use estimation to undertake empirically-based parameterization of this epidemic-related feedback function, $\omega^x(\dot{D}_t, t)$.

Hence, optimal labor supply and consumption of individuals drop when death rates rise and are given by:

$$n_t = \frac{1}{\theta} \left(1 - g(\dot{D}_t, t) \right) \tag{27}$$

$$c_t = w \cdot n_t = w \cdot \frac{1}{\theta} \left(1 - g(\dot{D}_t, t) \right)$$
(28)

The parameter $\frac{1}{\theta}$ in equation (27) represents optimal labor supply in steady state (n_{SS}) without the epidemic. The term $(1 - g(\dot{D}_t, t))$ is the endogenous time-varying reduction in the individual labor supply relative to the no-epidemic steady state, due to the rational behavioral response by individuals. In the course of the epidemic, actual labor supply is not necessarily the optimal labor supply given by equation (27), because the government might impose even stricter reductions on activity. Denote the limitation on labor supply mandated by the government by $1 - h(L_t, t)$, where L_t expresses lockdown measures. In that case, actual labor supply will follow the government mandate, and consumption will be set according to the budget constraint $c_t = wn_t$. Therefore, the following inequality should hold true in all periods:

$$n_t \le \frac{1}{\theta} \left(1 - h(L_t, t) \right) \tag{29}$$

Hence, formally:

$$n_{t} = \frac{1}{\theta} \min\{1 - g(\dot{D}_{t}, t), 1 - h(L_{t}, t)\} = \frac{1}{\theta} (1 - \max\{g(\dot{D}_{t}, t), h(L_{t}, t)\})$$
(30)

$$c_t = w \cdot \frac{1}{\theta} \left(1 - \max\{g(\dot{D}_t, t), h(L_t, t)\} \right)$$
(31)

2.2.2. Production and labor

Denote by \mathcal{P}_t the aggregate population of active, representative agents in period *t*. In the pre-epidemic steady-state: $\mathcal{P}_t = \mathcal{P}_{SS}$. During the epidemic, aggregate employment falls due to people being unable to work because of health reasons. The active population is therefore:

$$\mathcal{P}_t = \mathcal{P}_{SS} \cdot (1 - D_t - X_t - H_t - \phi M_t). \tag{32}$$

The hours of labor supplied by the representative agent in period t, n_t , derived in equation (30), is in daily hours. The aggregate hours worked in period t, N_t , are thus:

$$N_t = \mathcal{P}_t \cdot n_t \tag{33}$$

In the pre-epidemic steady-state, $N_{SS} = \mathcal{P}_{SS} \cdot n^{SS}$. During the epidemic, both aggregate employment \mathcal{P}_t and the intensive margin n_t change, so that:

$$N_{t} = \mathcal{P}_{SS} \cdot (1 - D_{t} - X_{t} - H_{t} - \phi M_{t}) \cdot \frac{1}{\theta} \left(1 - \max\{g(\dot{D}_{t}, t), h(L_{t}, t)\}\right)$$
(34)

Therefore, the relative scale of economic activity during the epidemic, aggregate hours worked by the active population relative to the pre-epidemic steady-state, is given by (noting that $n_{SS} = \frac{1}{a}$):

$$\frac{N_t}{N_{SS}} = \frac{\mathcal{P}_{SS} \cdot (1 - D_t - X_t - H_t - \phi M_t) \cdot \frac{1}{\theta} \left(1 - \max\{g(\dot{D}_t, t), h(L_t, t)\} \right)}{\mathcal{P}_{SS} n_{SS}}$$

= $(1 - D_t - X_t - H_t - \phi M_t) \cdot \left(1 - \max\{g(\dot{D}_t, t), h(L_t, t)\} \right)$ (35)

The production function of the firm is linear in aggregate hours:

$$Y_t = A \cdot N_t \tag{36}$$

$$Y_{SS} = A \cdot N_{SS} \tag{37}$$

Where the empirical counterpart of *A* is the average daily output per worker in the pre-epidemic steady-state. The firm maximizes its flow profits w.r.t. employment so:

$$\Pi_t = AN_t - wN_t$$

$$\rightarrow w = A \tag{38}$$

Table	1		
Menu	of the	Cvclical	Strategies

		week 1				week 2												
k	Strategy	Wor	kdays lo	cked	Μ	Т	W	Т	F	S	S	Μ	Т	W	Т	F	S	C
		week 1	week 2	in total														
0	lockdown only	5	5	10														
3	3-11	2	5	7														
4	4-10	1	5	6														
5	week on, week off	0	5	5														
6	3-day week	2	2	4														
7	7-7	1	2	3														
8	long weekend	1	1	2														
14	no intervention	0	0	0														



Thus aggregate consumption of active workers equals the aggregate flow output:

$$C_t = \mathcal{P}_t \cdot c_t = \mathcal{P}_t \cdot w \cdot n_t = \mathcal{P}_t \cdot A \cdot n_t = A \cdot N_t = Y_t \tag{39}$$

We normalize to 1 the pre-epidemic steady-state population of representative agents, i.e., $P_{SS} = 1$.

2.3. Policy based on time restrictions

The new policy, pertaining to the entire population, was introduced in Karin et al. (2020), where its epidemiological implications are analyzed extensively. Following an initial lockdown, move to a regime of k days of work and 14 - k days of lockdown, every 14 days. On work days, people are released from lockdown with strict hygiene and physical distancing measures on the same k weekdays for everyone. On lockdown days, people are kept away from work places as well as from other public spaces. Epidemiological measures need to be used throughout, including rapid testing, contact isolation, and compartmentalization of workplaces. All strategies respect regular weekends, facilitating application. Table 1 offers a visual summary.

Epidemic dynamics using these policies are discussed in detail below, where they are depicted graphically in Fig. 4 (see Section 4).

The rationale for the policy is as follows. Cyclical strategies reduce the average value of the reproduction parameter – which will be shown below to capture the progress of the disease – through two effects: time-restrictions and anti-phasing.

The *time-restrictions effect* is a reduction in the time *T* that an infectious person is in contact with many others, compared to the situation with no lockdown. For example, a 4-day work, 10-day lockdown cycle (k = 4) reduces *T* to $\frac{4}{14}T \approx 0.3T$. The *anti-phasing effect* uses the timescales of the virus against itself. Most infected people are close to peak infectiousness for about 3-5 days, beginning after a latent period of about 3 days (on average) after exposure. A proper work-lockdown cycle, such as a 4-work 10-lockdown schedule, allows most of those infected during work days to reach maximal infectiousness during lockdown, thus avoid infecting many others. Those with significant symptoms can be infectious for longer, but remain hospitalized, isolated, or quarantined along with their household members, preventing secondary infections outside the household.

As Table 1 shows, we only consider $k \le 8$ in our analysis of the cyclical strategies. This is because higher values of k imply shorter periods of lockdowns, for example, locking only on weekends. Though similar lockdown policies have been implemented (for example, in India), they do not line up with the epidemiological rationale of the cyclical policies. Furthermore, we find that, in the U.S. context, such extremely open policy tools are hardly consistent with a policy of efficient epidemic suppression. As noted above, online Appendix A provides further details.

2.4. The planner problem

The Social Welfare Function. The benevolent planner maximizes discounted social welfare, in expected present value terms. In any given period, social flow welfare is the sum of the agents' flow utilities, less the value of lives lost in that period:

$$W(T_0, T_1, T_2) =$$
(40)

$$= \int_{T_V=0}^{\infty} f(T_V) \cdot \int_{0}^{T} e^{-rt} \left[\frac{\mathcal{P}_{ss} \cdot (1 - D_t - X_t - H_t - \phi M_t) U(c_t, n_t, \dot{D}_t)}{-\mathcal{P}_{ss} VSL^U \cdot \dot{D}_t} \right] dt$$

where c_t and n_t are defined in equations (30)–(31), T is the planning horizon, T_V denotes the time till vaccine arrival distributed according to $f(T_V)$, and the term VSL^U denotes the statistical value of life in utility units.

The first term in the square brackets depends on the utility of active workers, from the group $\mathcal{P}_{ss} \cdot (1-D_t-X_t-H_t-\phi M_t)$, and is given by $U(c_t, n_t, \dot{D}_t)$, where c_t, n_t and \dot{D}_t are defined in (30)– (31) and in (17). The planner assumes that the welfare of inactive agents is zero. Hence, relative to their steady state utility, prior to disease outbreak, there is a social welfare loss for the duration of their inactivity.

The second term in the square brackets is a lost lives value term, not part of the individual utility function. It often features in this context; see, for example, Acemoglu et al. (p. 492, 2021), Alvarez et al. (eq. (7) on p. 371, 2021), and Farboodi et al. (eq. (13) on p. 13, 2021). It is made up of the product of the aggregate death flow \dot{D}_t and the value of statistical life in utility terms, VSL^U .

The planner takes into account the possibility of a pharmaceutical resolution of the epidemic. We assume that at time T_V a vaccine and a cure are found simultaneously (similar to the assumptions made by Acemoglu et al. (2021) and Alvarez et al. (2021)). We assume that after T_V no new cases arise and all inactive workers recover immediately. The waiting time till vaccine/cure introduction is uncertain, and it is distributed according to the probability density function $f(T_V)$, known to the planner. We elaborate on the functional form of $f(T_V)$ and the intuition underlying it in sub-section 2.5 below. Immediately after T_V , the economy moves to the no-epidemic steady state.

The Planner Maximization Problem. The planner takes the individual optimization described in sub-section 2.2.1 as given, and maximizes the afore-cited objective function through the choice of three regime-switching dates: T_0 , the start of the initial lockdown, T_1 , the date at which the planner switches from initial lockdown to the cyclical strategy, and T_2 , the date at which the planner abandons the cyclical strategy and removes all restrictions on economic activity. The lockdown regime L_t changes at dates T_0 , T_1 , T_2 , as described below, and the stringency of lockdowns under different regimes is fixed when the planner decides on the optimal timing T_0 , T_1 , T_2 . We describe the calibration of L_t under different regimes in sub-section 3.2.2 below. The full problem of the planner reads:

$$max_{T_0,T_1,T_2}W(T_0,T_1,T_2)$$
(41)

subject to equations (1)–(6), (9), (12), (13)–(17), (21), and (30)–(31), where lockdown policy L_t is defined as follows:

$$L_{t} = \begin{cases} 0 & if \quad t \leq T_{0} \\ L_{L} & if \quad T_{0} < t \leq T_{1} \\ L_{W} = 0 \text{ on open days or } L_{L} \text{ on closed days } if \quad T_{1} < t \leq T_{2} \\ 0 & if \quad t > T_{2} \end{cases}$$

$$(42)$$

where L_L is the lockdown stringency on "closed" days and $L_W = 0$ on "open" days. We further assume that the process of learning (see the last term in equation (9)) is triggered either by the imposition of lockdown of at least 14 days or by the endogenous response that is significant enough (i.e., $g(\dot{D}_t, t) > L_L$).

Social Welfare Loss. One can compute the social welfare loss in consumption or output terms. We define the flow welfare loss as follows:

$$WL(n_t, \dot{D}_t) \equiv \widetilde{V}(n_{ss}, 0) - \widetilde{V}(n_t, \dot{D}_t)$$
(43)

where \tilde{V}_t is the planner instantaneous utility, $U(n_{ss}, 0) = \tilde{V}(n_{ss}, 0)$ is steady state utility with no deaths, and

$$\widetilde{V}\left(n_{t}, \dot{D}_{t}\right) = \ln\left(w \cdot n_{t}\right) - \frac{1}{5} \left(\frac{n_{t}}{n_{ss} \cdot \omega^{x}(\dot{D}_{t}, t)}\right)^{5} - \ln\omega^{x}(\dot{D}_{t}, t) - VSL^{U} \cdot \dot{D}_{t}$$

$$\tag{44}$$

We also define the function WL_0 to be the welfare loss function when no disease is present

$$WL_0(n_t) \equiv WL(n_t, 0) \tag{45}$$

Appendix B shows that we can define the inverse function

$$\widetilde{n}_t = W L_0^{-1} (W L_t) \tag{46}$$

This function maps the instantaneous welfare loss, WL_t , to a corresponding hours value, \tilde{n}_t , that if worked in a world with no disease, would have caused this welfare loss. The decline in consumption (in percentage points) corresponding to WL_t is given by

$$CL(WL_t) = \frac{n_{ss} - WL_0^{-1}(WL_t)}{n_{ss}} = 1 - \frac{WL_0^{-1}(WL_t)}{n_{ss}}$$
(47)

Equation (47) translates the instantaneous welfare loss WL_t to an instantaneous reduction in consumption, CL_t , in percentage terms. The corresponding PDV value of consumption reduction corresponding to the welfare loss due the disease is given by

$$PDV_{CL}(WL_t) = \int_{-\infty}^{\infty} \exp(-rt) CL(WL_t) dt$$

(48)

2.5. Modeling vaccine arrival time

The term $f(T_V)$ is the probability density function of the availability of a vaccine at time T_V . This is an important term as it sets the horizon for the problem, acting as a hazard rate for leaving the state of the pandemic. It is an expression of the essential risk and uncertainty embodied in the planner problem. Note that were we to model an arrival time known with certainty, not only would an important real world aspect be removed, but such modeling might create an artifact in the optimal plan. The planner may enable an outbreak shortly before vaccine arrival, relying on the vaccine to eradicate it. Such a plan is not robust to delays in the arrival time. Relative to the interest rate *r*, expressing time preference, $f(T_V)$ plays the major quantitative role in discounting future streams. The constancy of *r* makes the planner problem time-consistent, as shown by Halevy (2004). We examine two alternative distributions for $f(T_V)$.

The baseline distribution is the Gumbel distribution, which use is justified by the following logic. We assume that the arrival of the vaccine is a result of simultaneous competition among many firms. The time of arrival is the minimum development time across these firms. Note that over the course of 2020–2022 over 110 vaccines were in clinical trials and dozens more in pre-clinical evaluations. The distribution of arrival time is then well approximated by a Gumbel distribution (see Kotz and Nadarajah (2000)), which is a member of the family of extreme value distributions. Specifically, it is used for modelling the minimum of a sample from many distributions, including exponential, logistic, and normal distributions. Under mild regularity conditions, it is suitable to be a model for a sample minimum even when the distributions from which the sample is drawn are unknown. In our setting, we remain agnostic about the distributions of vaccine development time by individual firms.

In terms of the model, T_V refers to the time of sufficient vaccination. With logistics, production times, gradual take-up rates, etc. an ex-ante expected 540 days seems reasonable relative to the March 2020 start date of the epidemic in the U.S.³ This number (540 days) is also the one used by Alvarez et al. (2021) and Farboodi et al. (2021), and is the middle of the range in Acemoglu et al. (2021).

The cumulative distribution function G(x) of a Gumbel distribution is defined over the real numbers and parametrized by a location parameter μ_G and a scale parameter σ_G :

$$G(x; \mu_G, \sigma_G) = 1 - \exp\left(-\exp\left(\frac{x - \mu_G}{\sigma_G}\right)\right)$$
(49)

We anchor the distribution's parameters (μ_G , σ_G), by positing that the mean of the distribution is 540 days, and that the probability of sufficient vaccination before day 360 is only 1%. These assumptions engender two linear equations:

$$E(Gumbel(\mu_G, \sigma_G)) = \mu_G - EulerGamma \cdot \sigma_G = 540$$

$$Q(Gumbel(\mu_G, \sigma_G), q) = \mu_G + \log(-\log(1-q)) \cdot \sigma_G = 360$$

where *E* is the mean and *Q* is the quantile function. Targeting a mean of 540 and Q(q = 0.01) = 360 leads to the solution of $\mu_G = 565.83$, $\sigma_G = 44.74$.

As an alternative we examine the exponential distribution, reflecting a Poisson process for vaccine arrival, at a fixed rate for any given day. This has been used by Acemoglu et al. (2021), Alvarez et al. (2021), Farboodi et al. (2021), and Jones et al. (2021). Its drawback is that it has a mode of zero, which is implausible in the case of vaccine development. We calibrate the parameter of the exponential distribution to 1/540, so that it has the same mean waiting time of 540 days as in Gumbel distribution.

Fig. 2 illustrates the resulting calibrated probability density functions. The Gumbel is shown by the solid line and the exponential by the dashed line.

The Gumbel distribution has an asymmetric bell-shaped form, with mode around 540 days, and a rather long left tail. The density drops quite fast after 540 days. The exponential distribution has the desired mean expected waiting time, 540 days, but the implied probability density is strictly decreasing, implying that the first day of the epidemic is the most likely date of vaccine development. This property of the distribution is highly counter-intuitive. However, assuming the exponential distribution is convenient because it fits in smoothly with the exponential discount factor which is frequently used in dynamic optimization problems in Economics. In Appendix B we compare the use of Gumbel and Exponential distributions as the basis for an optimizing planner, and show that the welfare loss in using the Exponential is higher.

 $^{^{3}}$ We have used 540 days in work on simulating this model since the summer of 2020. Actual developments turned out broadly consistent with this number.



Note: The parameter of the exponential distribution is set to 1/540, and the location and scale of Gumbel distribution are set to 565.83 and 44.74, respectively.

Fig. 2. Alternative Distributions of Waiting Time till Vaccine Arrival.

2.6. Benchmarks

We shall compare the results of simulating the model, calibrated for the U.S. economy, to the following benchmarks:

(i) and (ii) The polar cases of no policy intervention (i.e., no lockdown) or full lockdown till vaccine arrival.

(iii) The case whereby the duration of lockdown is set optimally. This is essentially the case of k = 0, i.e., no open days within a cyclical setting, and we shall denote it Optimal Lockdown.

(iv) A theoretical path trying to mimic real-world rationale by re-interpreting the planner problem as choosing thresholds for lockdown policy in terms of the critically ill, X. The first threshold defines a level X_0 whereby if $X_t > X_0$ an initial lockdown is imposed. Subsequently, a second threshold defines a level X_1 whereby if $X_t < X_1$ lockdown is released. Finally, a third threshold defines a level X_2 whereby if $X_t > X_2$ lockdown is re-imposed. The planner chooses the three thresholds optimally. The logic is that the first eruption of the disease is different from later eruptions, since the initial learning period has not yet occurred, and the endogenous response to death lags the infection. Therefore, X_0 allows the planner the freedom to lock earlier or later in the phase of initial eruption, while X_2 enables the planner to lock at a different threshold for later eruptions. The release threshold X_1 is the same throughout. Note that this strategy leads to recurrent lockdown and release policies, and, that the planner here is not as constrained as in the case of the cyclical strategies, where we have allowed for only three optimal points in time to be chosen.

These four cases are compared to six cyclical strategies, which use k = 3, 4, 5, 6, 7, 8.

In Section 6 we compare the model to the actual experience of the states of New York and Florida.

3. Calibration and solution methodology

We calibrate the model to fit the U.S. economy, which was badly hit by COVID19. Throughout we work in daily terms. In Section 6, we discuss the methodology and calibration values used for the analysis of two specific states in the U.S.

3.1. Calibration of the epidemiological model

In Table 2, we present the relevant parameter values for the two blocks, where we rely on sources in the epidemiological and medical literatures published in Science, Nature, the Lancet, and JAMA, as detailed in the table's notes.

3.1.1. The infection transmission block

For the duration numbers of the latency period $(1/\sigma)$ and the infectiousness period $(1/\gamma)$ we rely on studies that have appeared early on in the COVID19 pandemic and were published in Science; see Tian et al. (2020) and Li et al. (2020). These studies were co-written by university researchers from China (Tsinghua, Hong Kong, and others), from the U.S. (Harvard, Princeton, Columbia, Penn State, UC Davis, and NIH), and from the U.K. (Imperial, Oxford, and Southampton). Their findings are confirmed by studies on infector-infectee pairs; see He et al. (2020) and Ma et al. (2020). A number of papers (Tindale et al. (2020), Kong et al. (2020), and Ren et al. (2021)) highlight the fact that a significant part of disease transmission happens before the onset of symptoms. This implies that the latent period is shorter than the incubation period, which is usually 5-6 days (see the review in Bar-On et al. (2020) and the references therein). The latter appears within the clinical block to which we turn now.

Table 2
Calibration.

Parameter	Interpretation	Range	value
a. The Infection T	ransmission Block (SEIR)		
σ	latent period duration	3—5 days	1/3
γ	infectious period duration	4 – 5 days	1/4
b. The Clinical Blo	ck		
θ_P	incubation period	5 – 6 days	1/5
θ_M	days from symptoms till hospitalization	7 days	1/7
θ_H	days in hospital till ICU	2 days	1/2
θ_X	days in ICU before death	5.5 days	1/5.5
η	Prob. to be asymptomatic	20% - 50%	0.5
ξ	Prob. of hospitalization when symptomatic	$\frac{\#Hospitalized}{\#Infected} = [2\% - 4\%]$	$\frac{0.04}{1-0.5} = 0.08$
π	Prob. of ICU admission	10% - 40%	0.4
IFR	Infection Fatality Rate (implied)		0.008

Sources:

1. Panel a: Bar-On et al. (2020); He et al. (2020); Li et al. (2020); Tian et al. (2020);

2. Panel b - Bar-On et al. (2020); Huang et al. (2020); Richardson et al. (2020); Salje et al. (2020). *Notes*:

a. Range relates to number of days, unless noted otherwise.

b.
$$\frac{\#\text{Hospitalized}}{\#\text{Infected}} = \frac{\#\text{Hospitalized}}{\#\text{Symptomatic}} \cdot \frac{\#\text{Symptomatic}}{\#\text{Infected}} = \xi \cdot (1 - \eta) \implies \xi = \frac{\frac{\#\text{Hospitalized}}{\#\text{Infected}}}{1 - \eta}.$$

3.1.2. The clinical block

The value for the duration till death is given by $\frac{1}{\theta_P} + \frac{1}{\theta_M} + \frac{1}{\theta_H} + \frac{1}{\theta_X}$ and is set at 19.5 days based on CDC estimates. We use $\overline{X} = \frac{58,094}{329,529*10^6} = 1.8 \times 10^{-4}$ based on an estimate of 58, 094 ICU beds by the Harvard Global Health Institute. The implied Infection Fatality Rate (IFR) is given by $IFR = \xi \cdot \pi \cdot (1 - \eta) \cdot \delta_1$. Estimates of the Imperial College COVID-19

The implied Infection Fatality Rate (IFR) is given by $IFR = \xi \cdot \pi \cdot (1 - \eta) \cdot \delta_1$. Estimates of the Imperial College COVID-19 Response Team (2020) and the meta-analysis findings of Meyerowitz-Katz and Merone (2020) put *IFR* at 0.8%. Meyerowitz-Katz and Merone (2020) state: "Based on a systematic review and meta-analysis of published evidence on COVID-19 until July 2020, the IFR of the disease across populations is 0.68% (0.53% – 0.82%). However, due to very high heterogeneity in the meta-analysis, it is difficult to know if this represents a completely unbiased point estimate. It is likely that, due to age and perhaps underlying comorbidities in the population, different places will experience different IFRs due to the disease. Given issues with mortality recording, it is also likely that this represents an underestimate of the true IFR figure." Using this number and the numbers for ξ , π , and η in Table 2, we derive $\delta_1 = 0.5$. We then calibrate $\delta_2 = 0.5$ to capture the fact that, with extreme loads on the public health system, the probability to die increases to 1 for each patient in need of an ICU bed (see the reasoning in sub-section 2.1 above).

3.2. Calibration based on estimation

We employ daily U.S. data to estimate key relations and use the point estimates to calibrate the model.

3.2.1. Data

The data series used are daily deaths, daily employment, lockdown measures, and the derived transmission rate. Appendix C elaborates on data definitions and sources. Fig. 3 plots these four variables.

The transmission rate falls rapidly till early April 2020; it then rises somewhat till June 2020 and subsequently fluctuates around a fairly stable level. The disease is represented by the daily death series, where a big surge starts the first wave in March 2020 and where the decline starts in late April 2020; a second wave peaks in January 2021. Employment falls rapidly to a low of 80% of its pre-pandemic level in mid April 2020 and recovers, first rapidly and then slowly, though not all the way to its pre-pandemic level. Policy is shown by the weighted lockdown stringency index, which rises to a high level by the last third of March 2020. It subsequently drops in June 2020 and stays constant except for a rise in part of November 2020.

3.2.2. Dynamics of transmission and the reproduction parameter

We estimate equation (9), as reported in panel a of Table 3

The estimates imply the following. When $\frac{N_t}{N^{SS}} = 1$ i.e., the economy is not locked and there are no sick or dead, at time t = 0:

$$\beta_0 = \beta_\Lambda + \beta_W = 0.339 + 0.376 = 0.715$$



Notes: Data sources and definitions are elaborated in Appendix C.

Fig. 3. Key Variables.

Table 3 Estimation Results.

a. Infection Transmission

 $\beta_t = \beta_W - \beta_N \left(1 - \frac{N_t}{N^{5S}}\right)^{\alpha} + \beta_\Lambda \exp\left(-\Lambda t\right)$

March 15, 2020-February 28, 2021

β_{Λ}	Λ	β_W	β_N	α
0.339***	0.12***	0.376***	0.53***	0.69
(0.05)	(0.01)	(0.05)	(0.12)	(0.31)

 R²
 0.74

 RMSE
 0.0307

 n
 351

b. The Endogenous Response

 $\frac{N_t}{Nss} = 1 - \max(\varkappa_t L_t, \kappa_t \dot{D}_t) + \varepsilon_t$

March 1, 2020-February 28, 2021

κ	φ_{κ}	х	φ_{\varkappa}	μ_f	$\sigma_{_f}$
0.095*** (0.0001)	0.18*** (0.01)	0.42*** (0.007)	0.59*** (0.026)	245*** (2.36)	27.5*** (2.68)
R ² RMSE n	0.99 0.0139 365				

Notes:

a. Tables report point estimates with s.errors in parentheses.

b. p < 0.10, p < 0.05, p < 0.01.

and so $\mathcal{R}_0 = \frac{\beta_0}{\gamma} = 2.86$ Given the estimated rate of decline, $\Lambda = 0.12$, in a little less than a month individuals adjust their behavior to the presence of the disease; subsequently, when $\exp(-\Lambda t) \ll \beta_W$ we get:

$$\beta_t = \beta_W - \beta_N \left(1 - \frac{N_t}{N^{SS}}\right)^{\alpha}$$

This implies β_t drops endogenously below β_W as a function of employment, yielding reproduction parameter \mathcal{R}_t variation between 1.5 and 0.8.

3.3. Calibration of the economic model

Discounting. We posit a 4% annual discount rate (r = 0.04), converted to daily terms (used by individuals and consequently by the social planner).

The value of ϕ . As in Glover et al. (2020), we assume that anyone who has any symptoms self-isolates and does not work $(\phi = 1).$

The endogenous response, lockdowns, and employment.

In order to calibrate other values appearing in sub-section 2.2 we proceed as follows.

Without restrictions, equation (30) would allow employment to drop very significantly. This can even reduce employment to levels below what is usually regarded as essential employment plus work from home. Empirical estimates for the U.S. indicate that the minimum employment level under the most stringent lockdown measures was around 0.65 - 0.70 of full employment (see Blau et al. (2021) and Gregory et al. (2020)). We therefore calibrate this level of employment to be 0.68 and set $\overline{g} = 0.32$.

We postulate that the functions $g(\dot{D}_t, t), h(L_t, t)$ are given by:

$$g(\dot{D}_t, t) = \kappa_t \dot{D}_t \tag{50}$$

$$h(L_t, t) = \varkappa_t L_t \tag{51}$$

As discussed above, there is an overlap of compliance with lockdown and the endogenous response, so we use the maximal response. The variables x_t and κ_t express the idea that both endogenous individual response and compliance with lockdown are modeled with time decay f_t . We use the functional form proposed by Atkeson (2021, 2022) as follows:

$$f_t = \operatorname{normal_cdf}\left(\frac{t - \mu_f}{\sigma_f}\right) \tag{52}$$

$$\varkappa_t = \varkappa \left(1 - (1 - \varphi_{\varkappa}) \cdot f_t \right) \tag{53}$$

$$\kappa_t = \kappa \left(1 - (1 - \varphi_\kappa) \cdot f_t \right) \tag{54}$$

where normal_cdf is the CDF of the normal distribution, x_t and κ_t are the time-varying parameters of the effects of compliance lockdown and individual response on employment, respectively, and the parameters μ_f , σ_f , \varkappa , φ_{\varkappa} , κ and φ_{κ} are to be estimated.

The resulting estimating equation is given by:

$$\frac{N_t}{Nss} = 1 - \max(\varkappa_t L_t, \kappa_t \dot{D}_t) + \varepsilon_t$$
(55)

We non-linearly estimate equation (55) using data on $\frac{N_t}{Nss}$, *L*, and \dot{D}_t in the period from March 1, 2020 to February 28, 2021. The results are reported in Table 3b. We note that L_L is calibrated to be the highest share of workers who did not work during the course of the disease in the U.S. i.e., 0.20.

Employment, Wages, and the Utility Function.

For the planner problem and the simulations we further need to calibrate θ and A. To do so we use two U.S. data points, as do Eichenbaum et al. (2021, pp. 5162-3): the representative person works 28 hours per week reflecting the average number of hours worked from the Bureau of Labor Statistics 2018 ATUS and earns an annual income of \$58,000, using the 2019 estimate from the U.S. Bureau of Economic Analysis.

Thus, pre-epidemic, which we call steady-state (SS), when $g(D_t, t) = 0$ we get, converting weekly hours to daily terms:

$$n^{SS} = \frac{1}{\theta}$$

$$\theta = 1/4 \tag{56}$$

In the aggregate, no-epidemic steady-state: $\mathcal{P}^{SS} = 1$, and $N^{SS} = \mathcal{P}^{SS} \cdot n^{SS} = 4$.

Using daily income W and daily hours, the hourly wage is:

$$w = \frac{W}{n^{SS}} = \frac{\frac{58000}{365}}{4} = 39.73$$
(57)

We use equation (38) and set A = w. Value of Statistical Life.

The planner objective includes the term $\mathcal{P}^{SS} \cdot \dot{D}_t \cdot VSL^U$, where VSL^U is the value of statistical life. We determine its value and show how it fits in the social welfare function.

The central estimate in Hall et al. (2020) for the monetary value of statistical life lost to COVID19, *VSL^{USD}*, is 3.81 million USD, based on the EPA estimate of 270,000 USD per year and the authors' estimate of 14.1 years of remaining life on average. For robustness, we look at two sets of alternative values estimates in sub-section 5.3 below.⁴

To include these values in the social welfare function, we apply an oft-used methodology (see, for example, Farboodi et el. (2021, p. 16)), as follows: denote the Value of Statistical Life in utility terms by VSL^U , so that the event of death in the model is associated with utility loss of VSL^U . Individuals are indifferent between paying $SHARE_C$ of their flow consumption and avoiding the risk ε of losing VSL^U , and not paying $SHARE_C$ of their flow consumption and carrying the ε risk of losing VSL^U . Given the no-epidemic steady-state utility, this logic means that VSL^U should satisfy the following indifference condition:

$$\frac{\ln c_t - \theta^5 \frac{n_t^2}{5}}{r} - \varepsilon VSL^U = \frac{\ln \left((1 - SHARE_C)c \right) - \theta^5 \frac{n_t^2}{5}}{r}$$
(58)

where

$$SHARE_{C} = \frac{\varepsilon VSL^{USD} \times r}{\frac{C^{USD}}{365}}$$
(59)

The representative agent would be willing to pay $SHARE_C$ so as not to lose VSL^U with an ε risk of death; the payment, $SHARE_C$, is given by equation (59), paying $\varepsilon VSL^{USD} \times r$ each day, where r is the daily discounting rate is r.

Assuming $SHARE_C \ll 1$ we get $-\ln(1 - SHARE_C) \simeq SHARE_C$ and using our modeling of C = Y (which we have taken to be 58,000 USD), we get:

$$\varepsilon VSL^{U} = \frac{SHARE_{C}}{r} = \frac{\frac{\varepsilon VSL^{USD} \times r}{\frac{YUSD}{365}}}{r}$$
$$VSL^{U} = \frac{VSL^{USD} \times 365}{Y^{USD}}$$
(60)

Thus the VSL^U value we get is 23,977 for the baseline VSL^{USD} value of 3.81 million USD.

4. Results

We present the results for the baseline calibration and explore some variations in the next section. Section 2 of Online Appendix C delineates the methodology of the simulations.

The optimizing planner chooses how long to wait till first lockdown (T_0), when to start implementing a cyclical strategy (T_1), and when to release completely (T_2), for each instrument, namely for each given k. Optimal timing is based on probability-weighted scenarios for vaccine arrival over the horizon (T) of two years. According to the vaccine arrival time distribution that we assume, the probability that it will take more than two years to introduce the vaccine is practically 0. In the simulations, vaccine arrival is actually realized on day 540, at its arrival time in expectation, using the afore-discussed Gumbel distribution.

We present the results for the benchmarks discussed in sub-section 2.6 above – no intervention, full lockdown, optimal duration lockdown, and a thresholds strategy on X – and the cyclical strategies. Table 4 reports the planner's optimal timing T_0 , T_1 , T_2 , the resulting values of social welfare loss in terms of utility and in terms of consumption (both delineated in the table's notes), loss of annual GDP (in PDV, per annum terms, evaluated over two years), and the cumulative number of deaths, per 1 million people.

Fig. 4 shows the time evolution of key series for three benchmarks (to avoid clutter we omit the optimal lockdown case) and three of the cyclical strategies (k = 4, 5, 8). The series plotted are as follows: flow deaths (D_t), the endogenous response $\kappa_t D_t$, the effective reproduction parameter \mathcal{R}_e , the employment level relative to its pre-epidemic steady state $\frac{N_t}{N_{SS}}$, and the

⁴ In Section 2.2 of Appendix B we delineate an elaborate procedure to get such VSL estimates using a number of data sources, including Greenstone and Nigam (2020), and taking into account age groups, finding very similar numbers.

Table 4

Outcomes of the Different Policy Strategies - Baseline.

		T_0	T_1	T_2	WL (utility)	WL (cons)	GDP loss	D (per 10 ⁶)
Non-cyclic	al							
(<i>i</i>)	No Intervention	540	540	540	68.70	0.361	0.098	2,482
(<i>ii</i>)	Full lockdown	0	540	540	46.75	0.291	0.287	21
(iii)	Thresholds	n.a.	n.a.	n.a.	38.35	0.248	0.201	425
(iv)	Optimal L ($k = 0$)	8	487	487	42.24	0.268	0.256	84
Cyclical								
(v)	k = 3	0	14	499	34.65	0.226	0.211	43
(vi)	k = 4	0	14	511	32.95	0.221	0.197	76
(vii)	k = 5	0	98	534	33.30	0.218	0.201	42
(viii)	k = 6	0	120	540	36.14	0.240	0.194	218
(<i>ix</i>)	k = 7	0	296	540	38.68	0.252	0.224	113
(<i>x</i>)	<i>k</i> = 8	0	366	540	39.75	0.257	0.236	75

Notes: a. We use

$$\overline{g} = 0.32$$
, $VSL^{USD} = 3.81M$ \$, $IFR = 0.8\%$

b. T_0 is the start day of lockdown, T_1 start day of the cyclical strategy, and T_2 the release day; the numbers in these columns indicate day number since the start of the epidemic. The assumption is that vaccine arrival happens on day 540.

c. Welfare loss (utility) – the realized welfare loss when the vaccine actually arrives on day 540, in PDV terms, i.e., **the loss** of *W*, which is defined as follows:

$$WL(n_t, \dot{D}_t) \equiv \widetilde{V}(n_{ss}, 0) - \widetilde{V}_t(n_t, \dot{D}_t)$$

where \tilde{V}_t is the planner instantaneous utility and $U(n_{ss}, 0) = \tilde{V}(n_{ss}, 0)$ is steady state utility with no deaths, and

$$\widetilde{V}\left(n_{t}, \dot{D}_{t}\right) = \ln\left(w \cdot n_{t}\right) - \frac{1}{5}\left(\frac{n_{t}}{n_{ss} \cdot \omega^{x}(\dot{D}_{t}, t)}\right)^{5} - \ln\omega^{x}(\dot{D}_{t}, t) - VSL^{U} \cdot \dot{D}_{t}$$

d. Welfare loss (consumption terms) – the realized welfare loss when the vaccine actually arrives on day 540, in PDV terms, in consumption terms as given by:

$$PDV_CL(WL_t) = \int_{t=0}^{\infty} \exp(-rt) CL(WL_t) dt$$

where

$$CL(WL_t) = 1 - \frac{WL_0^{-1}(WL_t)}{n_{ss}}$$

See sub-section 2.4 for details. e. GDP loss – loss in GDP terms computed as:

$$V_{\rm Y} = \int_{0}^{730} e^{-rt} \left(\frac{Y_{\rm SS}}{N_{\rm SS}} (N_{\rm SS} - N_t) \right) dt$$

f. Cumulative deaths D (per 10^6) – stock of death per million.

flow (first difference) welfare loss in consumption terms $PDV_CL(W_t)$, as computed from equations (47)-(48). For clarity of presentation, the series (except for the endogenous response) are smoothed and what is presented is a 14 days moving average.

Table 4 and Fig. 4 show that the results can be characterized as follows.

The *cyclical strategies with low k* (k = 3, 4) lock the economy immediately for 14 days and then implement the cyclical strategy for an extended period of time. Because open days are relatively few, these strategies keep the epidemic under control. Flow deaths (D_t) and the stock of cumulative deaths (D) are very low, and \mathcal{R}_e is kept at 1 for most of the time – see the k = 4 case in Panel A of Fig. 4. As a consequence, the endogenous response $\kappa_t \dot{D}_t$ is low as well for most of the time, peaking at 1% of the work force during the initial breakout of the epidemic until it is contained.

During the prolonged cyclical phase, with relatively few open days, employment averages around 85% of its steady state level; GDP loss is around 20% - 21% in PDV per annum terms evaluated over two years. Welfare loss, in terms of the PDV over the two year period, is around 33 - 35 in utility terms (to be compared to other strategies below) and is around 22% - 23% in annual consumption terms. In flow terms, the loss is for most of the time around 15% of steady state consumption.



Cyclical Strategies k=4,5,8: the Dynamics of Key Variables

a. *k* = 4, 5, 8

Fig. 4. Six Policy Strategies. (For interpretation of the colors in the figure(s), the reader is referred to the web version of this article.)

The cyclical strategies with higher k (= 5, 6, 7, 8) lock the economy immediately for an extended period of time; the higher the k, the longer the initial lockdown period. They then implement the cyclical strategy up to vaccine arrival. During the initial lockdown period, starting at T_0 , \mathcal{R}_e is below 1, but when the cyclical regime is in effect, \mathcal{R}_e is above 1, and the higher the k, the higher is \mathcal{R}_e . Flow deaths (D_t) rise; see, for example, the k = 8 case in Panel a of Fig. 4. Consequently, the endogenous response $\kappa_t D_t$ rises as well enhancing the efficacy of cyclical tools and driving the effective reproduction parameter \mathcal{R}_e towards 1.

In the case of k = 5, \mathcal{R}_e converges to just above 1 almost immediately with the start of the cyclical phase, while for higher k values vaccine arrival occurs before this convergence has been completed.

Looking at the k = 8 case in Panel a of Fig. 4 we see the following: employment is low, at 80% during the initial lockdown, and then jumps to around 90% at the beginning of the cyclical phase. It gradually declines thereafter, due to the endogenous response. The loss of GDP and welfare is higher relative to the low k ones: GDP loss is 24% as compared to 20% - 21% in PDV per annum; welfare loss is 40 as compared to 33 - 35 in PDV terms (over two years), and 26% as compared to 22% - 23% in annual consumption terms. In flow terms, welfare loss is quite stable at around 20% of steady-state consumption during the year-long initial lockdown, it then drops with the transition to the cyclical regime, and subsequently gradually rises to around 16% due to the increase in the death toll, until vaccine arrival.





Cyclical Strategy k=5 and Non-Cyclical Benchmarks: the Dynamics of Key Variables

Notes:

a. The series plotted are as follows: the effective reproduction parameter \mathcal{R}_{e} ,

flow deaths (D_t), the employment level relative to its pre-epidemic steady state $\frac{N_t}{N_{SS}}$, the endogenous response $\kappa_t D_t$, and the flow welfare loss in consumption terms $PDV_CL(WL_t)$, as computed from equations (47)-(48).

b. For all series, except for the endogenous response, we use a smoothed 14 day average.

Fig. 4. (continued)

The results for three benchmarks, discussed in sub-section 2.6 above, to which we compare the afore-going cyclical policies, are as follows.⁵ Note, that the first two are non-optimizing, extremal benchmarks.

The no policy intervention case has the disease erupt: flow deaths (D_t) and the stock of cumulative deaths (D_t) are the highest across all the policies examined here; total deaths are higher by a factor of almost 60 relative to the k = 3 or k = 5

⁵ While here we compare to theoretical benchmarks, note that in Section 6 below we directly compare the model results to two actual U.S. cases – New York State and Florida.

cases. There is a strong endogenous response and employment falls to its lower limit, 68% of the steady state level. This endogenous response and the fast erosion of the Susceptibles pool, S_t , bring the effective reproduction number \mathcal{R}_e to unity in less than three months. GDP loss is relatively low, at around 10% in PDV per annum terms evaluated over two years, and results solely from the endogenous precautionary behavior of individuals. In PDV terms, the cumulative welfare loss is extremely high, at around 69 (compared to 33-40 in the cyclical strategies), which amounts to 36% of annual consumption (compared to 22% – 26% for the cyclical strategies). In flow terms, the welfare loss surpasses 50% of steady state flow consumption during the initial eruption and fluctuates between 10% and 30% of steady state consumption thereafter.

The full lockdown case has the disease under control as it entails an immediate lockdown, remaining in place until vaccine arrival. It leads to a very low number of deceased, 21 people per million. As it leads to a decline of employment of 80% of its steady state value, it has a substantial cost, at almost 29% of annual GDP in PDV terms over two years. This output loss translates to a welfare loss: in PDV terms the V loss is almost 47, as compared to 33 - 40 in the cyclical strategies, and is 29% in annual consumption terms. In flow terms, it is around 19% of steady state consumption for most of the time.

The thresholds policy case follows a prevalent policy rationale, though this is not the policy actually implemented anywhere. The planner optimally chooses thresholds for lockdown policy in terms of ICU hospitalizations, *X*, and keeps the epidemic

under control. Flow deaths (D_t) and the stock of cumulative deaths (D) are relatively low, and \mathcal{R}_e fluctuates between 0.76

and 1.40 most of the time. The endogenous response $\kappa_t D_t$ is low for much of the time, except for an initial spike. With the recurring lockdowns, employment fluctuates between 80% and 100% of its steady state level; GDP loss is around 25% in PDV per annum terms evaluated over two years. In terms of PDV welfare loss, the *V* decline is around 38 in utility terms, compared with 33 – 34 in the low *k* cyclical strategies, and is 25% in annual consumption terms, compared with 22% – 23% in the low *k* cyclical strategies. In flow terms, the social welfare loss fluctuates between 4% and 20% of steady state flow consumption.

The optimal lockdown case (k = 0) features a long lockdown beginning on day 12 and results in outcomes that lie between the two extremal benchmarks and the thresholds policy case in terms of social welfare loss (see row (iv) in Table 4). It has a lower death toll relative to the thresholds policy with a higher GDP loss. It is clearly inferior to the cyclical strategies.

Overall, the emerging picture is that the extremal benchmarks yield high welfare losses. This is so because policy is either non-existent or non-optimal. All other strategies have lower welfare losses because they optimally balance the cost of deaths and economic costs. A policy using thresholds on *X* achieves substantial improvement in welfare terms but is still inferior to the low *k* strategies (with $k \le 6$). The cyclical policies obtain better welfare results within a narrow range. Hence the different cyclical policies perform the best, within the set of policies examined, but between them there is relatively little difference. In Section 6 below, we compare the cyclical strategies with real world results for two states, New York and Florida.

5. Exploring planner policies

While the preceding section presented the baseline results, in this section we analyze the mechanism, its implications, and robustness. First, we study the underlying mechanism, exploring the rationale for the planner optimal decisions (subsection 5.1). Subsequently, we evaluate the cyclical policies by comparing them to the alternative benchmark policies using a plot of the policy possibilities frontier (5.2). Finally, we study variations in key parameters (5.3).

5.1. The mechanism

The underlying mechanism is as follows. The progression of the epidemic can be classified by the *effective* reproduction number \mathcal{R}_e , which depends both on the current reproduction number \mathcal{R}_t and on the current fraction of susceptibles S_t , i.e., $\mathcal{R}_e = \mathcal{R}_t \cdot S_t$, where \mathcal{R}_t reflects the current policy intervention regime and/or the endogenous behavioral response of individuals to the epidemic. A low \mathcal{R}_e is achieved (i) following an extensive exposure of the population to the virus (engendering a low S_t), (ii) by the endogenous response of individuals who decrease their activity, thereby reducing \mathcal{R}_t , (iii) by imposing stringent policy restrictions to curb virus spread, thereby lowering \mathcal{R}_t . The following classification is useful:

(i) When $\mathcal{R}_e < 1$, the epidemic is suppressed and the number of newly infected people declines with time.

(ii) When $\mathcal{R}_e = 1$, the disease is stable and an outbreak (i.e., a spurt of disease growth) is no longer possible, though susceptibles still do get infected at some slowly declining rate.⁶

(iii) When $\mathcal{R}_e > 1$, the amount of people who are infected daily grows, some of whom become ill, hospitalized, and may eventually die. However, this case is not sustainable over time: the rising daily death toll elicits an endogenous response from individuals, causing them to be cautious and to restrict their economic activity, which in turn reduces the infection transmission rate and thus reduces \mathcal{R}_e . Thus the endogenous response provides for a negative feedback mechanism, which prevents the disease from spiraling out of control.

We can now formulate the outcomes of the optimal planning problem in these terms. The outcomes presented in the preceding section basically follow one of two basic paths.

⁶ There is a decline in the rate of infection since the number of susceptibles S_t is declining with each person infected.

Exogenous containment. This path implies that \mathcal{R}_e is kept below or around 1 throughout most of the planning period using stringent restrictions, while preserving the pool of the susceptibles intact to a large extent. This path requires strong suppression measures to be imposed for long periods of time. These measures either reduce the epidemic or keep it growing at a very slow pace. The costs in terms of foregone output are relatively high, but the death toll is low.

Endogenous containment. This path implies that there is a reduction in \mathcal{R}_e to around 1 through the endogenous response of individuals, while policy interventions are loose or short-lived. Since this path involves less prolonged and delayed interventions, it is cheaper in terms of the loss of output, while the ensuing death toll is inevitably higher (relative to the *exogenous* containment path).

These two policy paths reflect planner choices in relation to the fundamental trade-off between economic costs and death tolls in managing the epidemic. The resolution of the trade-off, the optimal policy choice, depends on a number of factors: the extent to which economic activity can be maintained in lockdowns, the fatality rate of the virus, the scope of the endogenous response of individuals to death, and the value of statistical life. Importantly, it depends on the type of policy instruments available to the planner. It turns out that when cyclical strategies are in the policy toolkit, the fundamental trade-offs can be softened in a way that allows achieving lower economic costs and/or lower death tolls, all while waiting for vaccine arrival. The underlying rationale behind the social welfare improvement engendered by the cyclical strategies is that they average out *k* working days and (14 - k) lockdown days and therefore reduce \mathcal{R}_t . The approximation of the average \mathcal{R}_t , to be denoted \mathcal{R}_a , is given by⁷:

$$\mathcal{R}_a(k) \simeq \frac{(14-k) \cdot \mathcal{R}_L + k \cdot \mathcal{R}_{W_t}}{14} \tag{61}$$

where \mathcal{R}_L is the reproduction number on days of lockdown, given by:

$$\mathcal{R}_L = \frac{\beta_W}{\gamma} - \frac{\beta_N}{\gamma} \left(L_L \right)^{\alpha} \tag{62}$$

This is so under the assumptions that $L_L > \kappa_t \dot{D}_t$ and that the economy has passed the initial stages of the disease so $\exp(-\Lambda t) \ll 1$.

 \mathcal{R}_W is the reproduction number on open days (no lockdown) and is given by:

$$\mathcal{R}_{W_t} = \frac{\beta_W}{\gamma} - \frac{\beta_N}{\gamma} \left(\kappa_t \dot{D}_t \right)^{\alpha} \tag{63}$$

This is so under the assumption that the economy has passed the initial stages of the disease so $\exp(-\Lambda t) \ll 1$.

An effective lockdown policy measure L_L , should be set such that $\mathcal{R}_L < 1$ i.e. $L_L > \left(\frac{\beta_W - \gamma}{\beta_N}\right)^{1/\alpha}$. Combining equations (61),(62), and (63) yields:

$$\mathcal{R}_{a}(k,\dot{D}_{t}) \simeq \frac{\beta_{W}}{\gamma} - \frac{\beta_{N}}{\gamma} \left(\frac{k}{14} \left(\kappa_{t} \dot{D}_{t}\right)^{\alpha}\right) - \frac{\beta_{N}}{\gamma} \left(1 - \frac{k}{14}\right) \left(L_{L}\right)^{\alpha}$$

$$\tag{64}$$

The last equation shows that the reduction in the reproduction number (the terms with a minus sign) depends on a weighted average of lockdown measures and the endogenous response to death flow numbers.

We now make the connection between the afore-cited concepts: the effective reproduction parameter, \mathcal{R}_e , types of containment (*exogenous* and *endogenous*), and the average reproduction parameter, \mathcal{R}_a .

The basic classification into containment types is made under the assumption that the death toll is low $(\dot{D}_t \simeq 0)$.

If $\mathcal{R}_a(k, 0) \leq 1$, the cyclical strategy is based on *exogenous* containment, and the disease is contained as long as the cyclical policy is in effect. This is so because when $\dot{D}_t \simeq 0$ and $R_a(k, 0) \leq 1$, then $R_e \leq 1$ holds true throughout, the disease does not grow, and death rates remain suppressed so that the endogenous response plays no role.

If $\mathcal{R}_a(k, 0) > 1$, *endogenous* containment takes place. The disease grows, triggering an endogenous response that drives $R_a(k, \dot{D}_t)$ down until $R_e = 1$ and the disease is subsequently contained at some fixed level.

Note that all optimal cyclical strategies are based on restricting the size of the epidemic, so by the time of vaccine arrival, only 1% - 3% of the population is exposed to the virus. In other words, the policy-maker never relies on the erosion of the Susceptibles pool, S_t , as a means to drive the effective reproduction number \mathcal{R}_e down.

From the parameters estimated for the U.S., one can see that the low-*k* strategies (k = 3, 4) are used to achieve *exogenous* containment, using a very early lockdown phase followed by an extremely prolonged cyclical stage. Exogenous containment is possible since the average reproduction parameter \mathcal{R}_a implied by these strategies, even without an endogenous response, is low, and so they achieve an effective \mathcal{R}_e below unity. Crucially, they can bring down the infection while not closing down the economy completely.

⁷ The true average is not exactly a linear function of \mathcal{R}_W and \mathcal{R}_L (due to the anti-phasing effect and non-linearities of disease dynamics) and so this is a first order approximation.

⁸ The estimates for the relevant parameters satisfy this condition.

The k = 5 strategy is based on *endogenous* containment, following an initial lockdown period of just above 3 months. However, the endogenous response is not quantitatively large: since \mathcal{R}_a is only a little over 1, as the disease starts to grow, the endogenous response immediately brings \mathcal{R}_e to 1, keeping the disease, and death rates, at low levels ever since.

The less stringent strategies (k = 6, 7, 8) are used to achieve *endogenous* containment as well. These strategies use much longer initial lockdown periods (3 – 5 months), before moving to the cyclical phase. After initial suppression, using the cyclical strategies, the disease grows slowly, and with its rise, the endogenous response brings \mathcal{R}_e back to 1. These dynamics can be seen in Fig. 4 above. In Section 3 of online Appendix C we report the R_a and S_t values associated with these different policies.

5.2. The policy possibilities frontier

As discussed above, there are two major ways to deal with the epidemic, *exogenous* and *endogenous* containment. The trade-offs involved are most easily seen in a two-dimensional graph, that maps the outcome obtained under each instrument on two axes, the death toll per million people and the value of lost output in annual GDP terms. One can trace out a policy possibilities frontier using this graphical representation. Fig. 5 presents several variations of this frontier.

Panel a of the figure shows the frontier plot using the baseline values, discussed in Section 3 and presented in Table 4 and Fig. 4. Thus it shows the cyclical strategies as well as the benchmark strategies, discussed in sub-section 2.6. The analysis is conditional on the baseline $VSL^{USD} = 3.81M$ \$. The two extreme benchmarks of full lockdown (labeled 'lock till vaccine' arrival) and no intervention (labeled 'do nothing') are represented by two extreme points on the graph, with huge output or death toll, respectively, well beyond the frontier. The benchmark optimal duration lockdown policy (labeled 'lockdown only') lies above the frontier too, as does the thresholds strategy benchmark. All cyclical strategies are located at the left part of the graph. The figure clearly shows that the use of cyclical strategies brings about a very substantial improvement in outcomes relative to the two extremes of no intervention or full lockdown (as also seen in the values reported in Table 4). Compared to the optimal thresholds strategy, the cyclical strategies provide significant improvement too.

In the zoomed-in graph, the cyclical strategies trace out two linear frontier curves, of the two types of containment. The strategies marked in green follow *exogenous* containment policies. These are low-k strategies with just a few days open each fortnight. They trade off a higher cost of lost output for a lower death toll. They are on the frontier marked in green, nearer the origin. The other cyclical strategies, marked in red (k = 6, 7, 8), are not as powerful in suppressing the epidemic, as more open days are allowed every fortnight and therefore rely on *endogenous* containment. They are represented by the frontier schedule in red, further away for the origin.

Notably, the outcome of the k = 5 cyclical strategy, which belongs to the *endogenous* containment set, is located in the vicinity of the *exogenous* containment strategies outcomes. This is so as it achieves a low R_a (1.05) and is superior to the other *endogenous* containments strategies. With this exception, under the baseline calibration, the outcomes achieved by *exogenous* containment strategies are superior to the outcomes of *endogenous* containment strategies. The former limit the economic activity of agents, using the cyclical strategy so that the disease does not erupt. The latter strategies rely on the reaction of people to the death toll in order to limit economic activity. In order to limit the death toll, these strategies rely on a longer initial lockdown period, which is more costly in terms of social welfare loss.

5.3. Variations in key parameters

We turn to examine robustness of our results and look at variations in key parameter values. We continue using frontier terminology and plots.

First, we look at variations in the Value of Statistical Life. Varying the planner's preferences does not alter the big picture: the planner opts for a low death toll, and *exogenous* containment is preferred to *endogenous* containment. In the case of the *exogenous* containment strategies (k = 3, 4) as well as in the case of the k = 5 strategy, the planner is near the limit of the possibility to minimize death, so the optimal planning choice does not change much with the change in value of *V SL*. In the case of the *endogenous* containment cyclical strategies (k > 5) the change of *V SL* translates into a movement along the frontier with an almost constant slope. A rise of *V SL* causes the planner to lengthen the initial lockdown period (an increase in $T_1 - T_0$), raising the loss of GDP and reducing death. A lower *V SL* causes the planner to shorten the initial lockdown period, which translates into a lower GDP loss and a rise in the death toll. We present the results in full in sub-section 2.2.3 of Appendix B.

Second, the cited trade-offs also depend on the infection fatality rate. Panel b in Fig. 5 and Table 5 present the results with the alternative *IFR* values of 0.6%, and 1.2%.

A lower *IFR* means a less lethal disease so better outcomes can be achieved by the planner and thus welfare loss, compared to the baseline, is always lower. This enables the planner to shorten the initial lockdown period, which, combined with the lower *IFR*, leads to a lower death toll, lower GDP loss, or both. A higher *IFR* means that the disease is more lethal, and thus welfare loss compared to the baseline is higher. This leads the planner to lengthen the initial lockdown period, and with the higher *IFR*, there is either a higher death toll, a higher GDP loss, or both.

The most conspicuous feature of Fig. 5d is that low-k strategies fare much better, whatever the *IFR*. For all cyclical strategies, a higher *IFR*, usually translates into a upward-rightward movement (moving from black to blue points), with an



a. Cyclical and Benchmark Strategies

See notes d and e to Table 4.



Note:

Second graph is a "zoom-in" on the first graph.

Fig. 5. The Policy Frontier.

increase of both death and output loss. A lower *IFR* usually translates into a movement downward-leftward (moving from black to green points), with a decrease in death and output loss.⁹

We conclude that the finding of low-*k*, exogenous containment strategies being superior is robust to the parameter value variations examined.

⁹ The k = 7 strategy with low *IFR* is an exception. Under the baseline *VSL*^{USD} = 3.81*M*\$ and with lower *IFR* the planner narrowly prefers the 712 deaths per million and 0.17 output loss, to the 100 deaths per million and 0.22 output loss, which would have made it closer to the other 7 – 7 points.



b. Alternative Values of IFR

Notes: a. Legend is color IFR blue high, 1.2% black baseline, 0.8% green low, 0.6%

b. For each IFR value we plot strategies k = 3...8. These are labelled only for the black points.

Fig. 5. (continued)

Table 5

Parameter Variations - IFR.

value of life: $IFR = 0.6\%$						
k	<i>k</i> = 3	k = 4	<i>k</i> = 5	k = 6	<i>k</i> = 7	<i>k</i> = 8
T ₀	0	0	0	0	18	0
T_1	14	14	98	113	35	352
<i>T</i> ₂	498	507	530	540	540	540
WL (utility)	34.44	32.49	32.99	35.26	39.02	39.31
WL (cons)	0.224	0.217	0.215	0.235	0.262	0.254
Output loss	0.210	0.195	0.200	0.192	0.166	0.233
D (per 10 ⁶)	33	60	32	185	712	73
value of life: $IFR = 1.2\%$						
$\frac{\text{value of life: } IFR = 1.2\%}{k}$	k = 3	k = 4	<i>k</i> = 5	<i>k</i> = 6	k = 7	k = 8
value of life: $IFR = 1.2\%$ k T_0	k = 3	k = 4 0	k = 5 0	k = 6	k = 7 0	k = 8
value of life: $IFR = 1.2\%$ k T_0 T_1	k = 3 0 14	k = 4 0 37	k = 5 0 99	k = 6 0 162	k = 7 0 317	k = 8 0 380
value of life: $IFR = 1.2\%$ k T_0 T_1 T_2	k = 3 0 14 501	k = 4 0 37 509	k = 5 0 99 540	k = 6 0 162 540	k = 7 0 317 540	k = 8 0 380 540
value of life: $IFR = 1.2\%$ k T_0 T_1 T_2 WL (utility)	k = 3 0 14 501 35.05	k = 4 0 37 509 33.37	k = 5 0 99 540 33.85	k = 6 0 162 540 37.42	k = 7 0 317 540 39.56	k = 8 0 380 540 40.47
value of life: $IFR = 1.2\%$ k T_0 T_1 T_2 WL (utility) WL (cons)	k = 3 0 14 501 35.05 0.230	k = 4 0 37 509 33.37 0.221	k = 5 0 99 540 33.85 0.223	k = 6 0 162 540 37.42 0.247	k = 7 0 317 540 39.56 0.257	k = 8 0 380 540 40.47 0.261
value of life: $IFR = 1.2\%$ k T_0 T_1 T_2 WL (utility) WL (cons) output loss	k = 3 0 14 501 35.05 0.230 0.212	k = 4 0 37 509 33.37 0.221 0.201	k = 5 0 99 540 33.85 0.223 0.204	k = 6 0 162 540 37.42 0.247 0.204	k = 7 0 317 540 39.56 0.257 0.230	k = 8 0 380 540 40.47 0.261 0.240

Notes: See Table 4.

5.4. Discussion

What is the broader picture emerging from the analysis? Even in the absence of any restrictions, two mechanisms lead to output loss. The first is the direct effect generated by infected people who cannot work. The second, and more significant one, is that individuals self restrict their economic activity in order to avoid infection. In our analysis output loss without policy intervention was shown to be significant. This demonstrates the critical role of the endogenous individual response to the disease and holds true under other active planner strategies too. To be more specific, under laissez-faire, i.e., no intervention, the negative feedback mechanism engendered by the endogenous response alone curbs the spread of the disease; as shown in Fig. 4d, \mathcal{R}_e drops to below 1 in a month and subsequently converges to 1. After 18 months, only

30% of the population are infected and the ICU limit is almost never breached. Without such endogenous response, disease growth would be so rapid that even after a considerable amount of the population is infected and the susceptible pool is greatly depleted (such that $\mathcal{R}_e = 1$), it takes a while for the disease to slow down, with a considerable death toll.

At the other extreme, under full lockdown, it is not possible to eliminate the disease completely. A seed of the disease is always present in the population, or, alternatively, is imported from the outside.

These findings give perspective on the role of policy. The main aim of policy is not to "flatten the curve" or prevent an overshoot. These aims are achieved by the endogenous individual response. Policy cannot eliminate the disease or avoid its economic costs. Using lockdowns, policy is truly needed in order to "buy time." This is the time needed for individuals to adapt to the disease and for governments and for the pharmaceutical industry to develop vaccines and treatments. The question, then, is how best to do so. We show that cyclical lockdown strategies are useful in this context, as they enable the planner to smooth out the costs over a longer period of time. Doing so they reduce the total social welfare loss in PDV terms. We have analyzed the dynamics of these policies in detail, including its two modes of containment and the fact that these strategies keep \mathcal{R}_e close to 1. Note that our simulations incorporate vaccine arrival time as discussed in sub-section 2.5 above and in Appendix B. Iverson et al. (2021) analyze and emphasize the importance of expected arrival time in this context.

6. The cyclical strategies vs actual experience

The cyclical strategies can be compared to actual real world experience. We do so by simulating optimal plans under the cyclical strategies and comparing them to a policy path based on the experience of the states of New York and Florida. The choice of these two states is motivated by the fact that both experienced high levels of the epidemic but very different dynamics.

6.1. Data and methodology

We use state-level data on daily deaths, daily employment, lockdown measures, and the derived transmission rate, from the same sources spelled out in Appendix C for the national-level data. N_{SS} is taken to be state-level employment on March 1, 2020.

We employ the same methodology for the simulation as described above, after re-estimating the transmission equation and the endogenous response equation for each state. The equations specifications were adapted to cater for the statespecific data. For the transmission equation we used $\alpha = 1$ for both states and a different specification for β_{Λ} in the Florida equation. The endogenous response equation had no time decay term, as none is apparent in the data. See the specifications delineated in the table. The estimates are reported in Table 6.

The fit of the equations is good for the most part; the estimated κ indicates a lower endogenous response relative to the U.S. as a whole but close to the post-fatigue value in the U.S. The implication is that in both Florida and New York, employment was controlled more by the policy response and less by the endogenous response.

In New York the estimated β_t captures well the rapid drop of the reproduction parameter \mathcal{R}_0 from its initial value due to strict lockdown, as well as the subsequent gradual rise to values well above $\mathcal{R}_t = 1$. In Florida, the range of values for β_t , in estimation as in the data, is much narrower; for most of the time the reproduction parameter is above 1.

6.2. Results

We report the results as we did in Table 4 and Fig. 5 above for the whole U.S. economy. We compare the results to actual data.¹⁰

Both Table 7 and Fig. 6 refer to the data sample period used for the estimation reported in Table 6. The first panel of the figure looks at the data together with the cyclical strategies, while the second looks only at the latter set.

There is a substantial difference between the two states and we analyze each in turn.

New York State. As seen in Fig. 6, the actual data point for NYS is very far from all the cyclical strategy points. In comparison to the cyclical strategies, the actual data exhibit a far higher death toll and just a slightly lower GDP loss. In the data the death toll is 2, 346 dead per million and the GDP loss is 0.15. The cyclical strategies entail a far lower death toll, between 7 and 29 per million, and a GDP loss of between 0.16 and 0.21, the former number being just a little above the data number.

Looking within the cyclical set, both the second panel of Fig. 6 and Table 7 show that there is variation, with the strategies using k = 3 and k = 4 being different from those using k = 5, 6, 7, 8. This panel depicts linear policy possibilities

$$V_{Y} = \int_{0}^{730} e^{-rt} \left(\frac{Y^{SS}}{N^{SS}} (N^{SS} - N(t)) \right) dt$$

¹⁰ As explained in Table 4 above, GDP loss is given by

Table 6

Estimation Results for New York State and Florida.

a. New York State, March 15, 2020 to February 28, 2021.

a. Infection Transmission

 $\beta_t = \beta_W - \beta_N \left(1 - \frac{N_t}{N^{SS}} \right) + \beta_\Lambda \exp\left(-\Lambda t\right)$

β_{Λ}	Λ	β_W	β_N
0.202	0.967***	0.517***	1.519***
(0.437)	(1.777)	(0.0112)	(0.0797)

 R²
 0.559

 RMSE
 0.0631

 n
 351

b. The Endogenous Response

 $\frac{N_t}{Nss} = 1 - \max(\varkappa L_t, \kappa \dot{D}_t) + \varepsilon_t$

κ	х
0.00506***	0.224***
(0.000218)	(0.00274)

 R²
 0.998

 RMSE
 0.0339

 n
 352

Notes

a. The tables report point estimates with s.errors in parentheses.

b. p < 0.10, p < 0.05, p < 0.01.

b. Florida, March 27, 2020 to January 31, 2021

a. Infection Transmission

 $\beta_t = \beta_W - \beta_N \left(1 - \frac{N_t}{N^{SS}} \right) + \left(\beta_{Max} - \beta_W \right) \exp\left(-\Lambda t \right)$

Λ	β_W	β_N
0.0912	0.329***	0.496***
(0.174)	(0.0121)	(0.135)

 $\beta_{Max} = 0.364$

0.971
0.0495
311

b. The Endogenous Response

 $\frac{N_t}{Nss} = 1 - \max(\varkappa L_t, \kappa \dot{D}_t) + \varepsilon_t$

к	х
0.0118***	0.174***
(0.000754)	(0.00334)

 R²
 0.999

 RMSE
 0.0326

 n
 324

Notes:

a. Tables report point estimates with s.errors in parentheses.

b. *p < 0.10, **p < 0.05, ***p < 0.01.

frontier schedules. Table 7 shows that the values of welfare losses vary across these strategies within a narrow range. The policy differences across the cyclical strategies are manifested in T_1 , which values are much higher in the k = 5, 6, 7, 8 group, so GDP losses and welfare losses are higher. We can call the former group *exogenous* containment strategies, and the latter group *endogenous* containment strategies, using the terms delineated above. Only the *exogenous* containment strategies bring the average effective reproduction parameter, \mathcal{R}_a , to values close to unity. The *endogenous* containment strategies per se are unable to do so and do not contain the epidemic, hence the planner resorts to a very lengthy initial lockdown period









Fig. 6. Policy Possibilities Frontier Plots for New York State and Florida.

(see Table 7). The reasons for this inability to contain the epidemic are two: the implied \mathcal{R}_W , derived from the estimated β_W , is relatively high and the estimated endogenous response parameter, κ , is relatively low.

It is clear that actual policy underperformed and achieved only weak containment. In particular, note that the use of moderately stringent cyclical tools implies a dramatic reduction in the death toll and only a slight increase in output costs relative to actual experience. Timing is key here – the optimal use of the cyclical strategies here involves an immediate initial lockdown, which proves to be critical in curbing the disease during its exponential growth phase. Compared to the optimal policy, waiting with the initial lockdown resulted in death tolls that are higher by a factor of at least 81. It thus appears that lockdown in New York State started very late and that a significant outbreak was facilitated.

Florida. This state had a death toll of 1,230 dead per million and a GDP decline of 0.08; the cyclical strategies entail once more a far lower death toll, between 11 and 18 per million, but a bigger loss of GDP, around 0.14 – 0.17. The cyclical strategies are able to bring the average effective reproduction parameter \mathcal{R}_a to values close to unity and so to contain the epidemic. Only the k = 3 strategy is based on *exogenous* containment, with $\mathcal{R}_a < 1$, with all other cyclical strategies

b. Florida, March 27, 2020 to January 31, 2021.



2. Zoomed in



Notes: See notes d and e of Table 4.

Fig. 6. (continued)

k = 4, 5, 6, 7, 8 based on *endogenous* containment. Initial lockdown duration, T_1 , is a month and half with k = 3, 3-4 months for k = 4, 5, and longer durations for higher k values, with higher GDP losses and welfare losses.

Actual policy in Florida seems to have preferred a lower GDP decline and a much higher death toll. Using the current set-up this can be viewed from two perspectives:

(i) Absent cyclical tools, actual policy outcomes are compatible with a strategy based mostly on *endogenous* containment, such as to lock and then release until a vaccine is found, without a cyclical phase. For example, a simulation of locking for 56 days (8 weeks) and then releasing yields a GDP loss of 0.076 and a death toll of 1,306 dead per million, very close to the data.

(ii) When cyclical tools are available, and when VSL is at 2.15*M*\$, then an optimal cyclical strategy can lead to the data outcome. In panel c of Table 7 we show that a planner using k = 10 (which locks only on weekends) will optimally choose $T_0 = 49$, $T_1 = 63$, $T_2 = 421$, i.e., delay the initial lockdown for 7 weeks thereby letting the disease erupt, lock for two weeks, and then lock only on weekends (k = 10). Such a strategy yields a GDP loss of 0.079 and a death toll of 1,294 dead per million, similar to the actual data. Note that this value of statistical life, 2.15*M*\$, is 56% of our baseline value (3.81*M*\$) and

0.080

0 1 5 4

1294

Table 7

State Level Outcomes (at the end of the sample period). a. New York State - March 15, 2020 and February 28, 2021.

		T_0	T_1	<i>T</i> ₂	WL (utility)	WL (cons)	GDP loss	D (per 10 ⁶)		
$L_L = 0.22$; $VSL^{USD} = 3.81M$ \$; $IFR = 0.8\%$										
(<i>i</i>)	k = 3	0	14	540	29.45	0.170	0.168	20		
(<i>ii</i>)	k = 4	3	58	540	28.41	0.165	0.160	33		
(iii)	k = 5	0	345	540	36.43	0.206	0.210	7		
(iv)	k = 6	0	449	540	36.74	0.208	0.212	7		
(v)	k = 7	0	479	540	36.74	0.208	0.212	7		
(vi)	k = 8	0	500	540	36.74	0.208	0.212	7		
b. Florida, March 27, 2020 to January 31, 2021										
		T_0	T_1	T_2	WL (utility)	WL (cons)	GDP loss	D (per 10 ⁶)		
$L_L = 0.21; VSL^{USD} = 3.81 M$ \$, $IFR = 0.8\%$										
(<i>i</i>)	<i>k</i> = 3	0	44	449	24.55	0.148	0.146	17		
(<i>ii</i>)	k = 4	0	105	473	24.30	0.146	0.144	14		
(iii)	k = 5	0	115	506	23.38	0.141	0.139	15		
(iv)	k = 6	0	127	540	22.52	0.137	0.133	18		
(v)	k = 7	0	205	540	24.90	0.149	0.148	13		
(vi)	k = 8	0	282	540	28.35	0.168	0.169	11		
c. Florida, March 27, 2020 to January 31, 2021, VSL= 2.15 <i>M</i> \$										
		T_0	T_1	T_2	WL (utility)	WL (cons)	GDP loss	D (per 10 ⁶)		
$L_L = 0.21; VSL^{USD} = 2.15M$ \$, <i>IFR</i> = 0.8%										
(<i>i</i>)	k = 4	49	63	444	23.18	0.142	0.115	469		
(<i>ii</i>)	k = 6	49	63	454	23.37	0.144	0.100	741		
(iii)	k = 8	49	63	442	24.63	0.149	0.089	1039		

(iv) Notes:

1. See Section 7 of the paper.

k - 10

2. See the notes to Table 4.

is \$152,000 in annual terms. Such a low value, though still within the range cited by Hall et al. (2020) of \$100,000 to \$400,000 per year, may have been underlying the policy actually used in Florida.

25 99

63

49

421

7. Conclusions

Looking forward, the COVID19 pandemic is unlikely to be the last one. There are dozens of coronaviruses (according to CDC data, seven of which can infect humans); new influenza strains can be as deadly as the ones that have killed millions in the pandemics of the twentieth century; and Zoonotic infections (like Ebola, SARS, or zoonotic influenza) have increased significantly over time (see Jones et al. (2008) and Salyer et al. (2017)). These diseases pose an increasing and critical threat to global health security, with ever growing connectivity and mobility of people, animals, and goods. The June 2021 report of a high level G20 panel¹¹ states that "We are in an age of pandemics.... There is every likelihood that the next pandemic will come within a decade – arising from a new influenza strain, another coronavirus, or one of several other dangerous pathogens. Its impact on human health and the global economy could be even more profound than that of COVID-19." Meganck and Baric (2021), May (2021), and references therein, provide detailed elaboration of the key forecasted threats.

Our paper has studied the optimal timing patterns of a new, cyclical policy strategy, consisting of alternating days of opening and lockdown. Given significant trade-offs between health outcomes (deaths, breaches of ICU capacity) and economic outcomes (loss of output and employment), the analysis has shown that epidemic management policy based on such time restrictions may lead to significant improvement in terms of social welfare. The comparison was made relative to four hypothetical benchmark policies, as well as to the experience of New York State and Florida.

The analysis, which is relevant for any future epidemic, laid down the principles for time restrictions policy, as well as a framework for comparative policy analysis. Exploring this policy seems a promising avenue for future research in the context of managing epidemics. The analysis clearly shows that such time-based strategies allow for a nuanced response to

¹¹ See the report of the High Level Independent Panel on Financing the Global Commons for Pandemic Preparedness and Response at https://pandemic-financing.org/report/foreword/.

observed epidemic dynamics, without resorting to singling out a particular population group. It is important to note that the afore-mentioned advantages of the cyclical strategies over prevalent policies are likely to be a lower bound of their full benefits. First, in our model the planner is deliberately constrained in the way cyclical tools are applied; for example, the planner is not permitted to mix within the set of strategies, or apply them in a staggered way. Giving the planner additional degrees of freedom, as we do for the benchmark thresholds strategies, should increase the advantages of the novel instruments over prevalent policies. Second, and not less significant, our model does not allow us to quantify the additional benefits of cyclical tools, such as the predictability of production that they entail, as well as their potential to alleviate part of the negative impact of prolonged isolation on mental well-being. These issues require a more elaborate analysis and are left for future research.

It should be noted that we have abstracted from heterogeneity across individuals and from modeling the coordination of work and leisure decisions. We have also abstracted from policy such as testing or contact tracing. Such issues have been addressed in the COVID19 context, albeit in different modeling frameworks, by a number of recent papers. Thus, Berger et al. (2022) examine how testing and targeted quarantine will work in an efficient reopening, whereby output increases while deaths are reduced. They use virological and serological testing, and their model features permanently asymptomatic individuals, incomplete information, and a reduced-form behavioral response. Chari et al. (2021) examine how targeting of lockdown policies in a world with heterogeneity achieves better outcomes relative to indiscriminate lockdowns. Glover et al. (2020) analyze old-age issues and redistribution policy. Fuchs-Schündeln et al. (2021) look at the heterogeneous impact of school closures on children during COVID19. Extensions of the analysis along these lines are also left for future research.

Data availability

All data are available online on journal web site.

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Appendix A. Supplementary material

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.red.2023.01.004.

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