

Adaptive Targeted Infectious Disease Testing

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Abstract

We show how to efficiently use costly testing resources in an epidemic, when testing outcomes can be used to make quarantine decisions. If the cost of false quarantine and false release exceed the cost of testing, the optimal myopic testing policy targets individuals with an intermediate likelihood of being infected. If individuals arrive over time, the policymaker faces a dynamic tradeoff: using tests for individuals for whom testing yields the maximum immediate benefit vs. spreading out testing capacity across the population to learn prevalence rates thereby benefiting later individuals. We describe a simple policy that is nearly optimal from a dynamic perspective. We discuss how to implement an adaptive targeted testing policy in practice.

1 Introduction

We have a simple message for all countries: test, test, test. Test every suspected case. If they test positive, isolate them and find out who they have been in close contact with up to 2 days before they developed symptoms, and test those people too...Once again, our key message is: test, test, test. (Tedros Adhanom Ghebreyesus, WHO Director-General’s opening remarks at the media briefing on COVID-19, 16 March 2020).

Testing is a critical part of a response to an epidemic. At an individual level, testing allows authorities to identify and quarantine sick people, thereby stopping the spread of the disease. At a country level, testing helps authorities keep track of the disease spread, make decisions about social distancing rules, and plan for provision of supplies. However, during a sudden epidemic, such as COVID-19, testing resources can be limited (Gupta, 2020). Evidence from the COVID-19 pandemic suggests that countries had very different testing capacities (Hasell et al., 2020). One way to think about the cost of a test is in terms of the value of testing the best possible alternative person. In other words, testing has a (shadow) cost because capacity might be difficult or impossible to ramp up quickly. Even if testing kits themselves are cheap, large-scale laboratory testing capacity might be infeasible (Hope, 2020) or it might be difficult to quickly reach all those who need testing (Weaver and Ballhaus, 2020). In this paper, we offer a simple framework that formalises the key tradeoffs

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that policymakers might face under limited testing capacity, and propose an adaptive policy that can help them allocate their testing capacity as effectively as possible.

Throughout the paper, we work under the assumption that the policymaker’s objective is to minimize the total cost of disease spread. These costs are of many kinds: cost of human lives, cost of lost labour income, cost of testing kits, and reputation cost of unnecessary quarantine.

In our model, potentially sick individuals arrive over time. People might show up at the hospital because they think they have the necessary symptoms, or doctors might go out to survey and actively test people. The policymaker can take one of three actions, for each individual:

- Test the individual.
 - If the individual tests positive, they are quarantined.
 - If the individual tests negative, they are not quarantined.
- Not test the individual, but quarantine them.
- Not test the individual and release (i.e., not quarantine) them.

In our model, the policymaker observes characteristics of individuals. These characteristics can be health-related: whether the individual has the relevant symptoms or whether the individual has been in contact with others who have symptoms or have tested positive. The characteristics can also be observables relevant to the social and economic cost of the disease: whether the individual is in a critical occupation, whether the individual has child-care responsibilities etc.

In our model, the policymaker has access to a statistical model that can estimate the probability of the individual’s having the disease conditional on the observables. The policymaker can therefore assess the overall expected costs associated with quarantining or releasing the individual. The statistical model is imperfect in the sense that it cannot perfectly predict whether a given individual is infected based on their characteristics.

We assume that the test is perfect, but costly, so it is not possible to test everyone.¹

If the policymaker decides to test an individual, they incur a testing cost, but they will subsequently take an optimal quarantining decision because the test is perfect. If the policymaker decide not to test the individual, they can make one of two costly errors:

- False Quarantine: Quarantining an individual who is not infected.
- False Release: Not quarantining an individual who is infected.

In summary, the policymaker faces three types of costs:

- Cost of testing (marginal cost or the cost of relaxing the capacity constraint).
- Cost of false quarantine.
- Cost of false release.

During the COVID-19 pandemic, countries appear to have used their testing capacity in different ways. For example, there is a substantial variation in the number of confirmed cases per test even after controlling for prevalence and testing capacity (Hasell et al., 2020). The question we answer in this paper is: What is the testing policy that minimises the overall costs?

The following example elucidates the key tradeoffs. Suppose that the policymaker only has 10,000 testing kits, but there are 20,000 individuals who have arrived at the hospital.

¹This assumption does not affect the key messages of this paper as we show in Section 5. For a discussion of these issues, see Galeotti et al. (2020).

Whom should the policymaker test? Consider two policies have been used repeatedly in the current pandemic.

Priority Testing: Rank all individuals according to how likely they are to have the disease. Then test 10,000 people who are most likely to have the disease.

Several countries, such as the United States and United Kingdom, implicitly used the Priority Testing policy during the initial stages of COVID-19 pandemic by restricting testing to patients with strong symptoms, to those who travelled to infected area or to those who have been in contact with infected people (Padula, 2020).

Priority Testing might be the optimal policy only if the cost of falsely quarantining individuals who are not infected is extremely high. But by testing individuals that are likely to have the disease, the policymaker could potentially be “wasting” tests: if the cost of a false quarantine errors is not too high, the people with the highest estimated likelihood of the disease could be quarantined without testing. During the COVID-19 pandemic, many countries eventually followed this logic and advised that anyone who has symptoms or who lives with someone who has symptoms of COVID-19 must self-isolate without testing for an extended period.

Random Testing: Test 10,000 individuals at random.

During the COVID-19 pandemic, a few countries and cities used random testing and there have been several calls to expand Random Testing (Oster, 2020; Padula, 2020). Random Testing is a sensible policy if tests are very cheap. The policymaker can learn the prevalence of the disease (thereby being able to make better decisions about testing of individuals in the future), but most people tested will not be infected. Therefore, many tests will, once again, be “wasted”.

We proceed as follows. In Section 3, we point out that to make optimal decisions about testing the policymaker needs to trade off the costs of false quarantine and false release relative to the cost of testing. Under fairly mild conditions, the optimal myopic testing policy is to test individuals with an *intermediate* likelihood of the disease. Priority Testing is therefore not myopically optimal in general because the policymaker would prefer to quarantine individuals with a high likelihood of infection without testing them and would not test or quarantine individuals who are very unlikely to be infected.

In Section 4, we look at how the policymaker’s problem changes when she cares about the future. In this case, we show that the policymaker will not initially want to follow the myopic policy. Rather the policymaker would want to “explore” by initially spreading out some of her testing capacity and sacrificing some immediate benefit. The reason is that such exploratory testing gives the policymaker valuable information about the prevalence of the disease which she can use to make better decisions about the testing of future individuals. A simple dynamic testing policy due to Thompson (1933) tells the policymaker how much exploration is (nearly) optimal. The Thompson policy starts by initial exploratory testing. If the prevalence rate is stable, the payoff to exploration disappears as the number of tested individuals grows because disease prevalence becomes precisely estimated. Over time, the Thompson policy converges to the optimal myopic testing policy.

In Section 5, we discuss some practical implementation issues, including imperfect testing. We also emphasise that our main discussion assumes that true prevalence rates across groups do not change over time. However, in epidemics, prevalence rates can change considerably. We sketch how such “non-stationarity” can be taken into account in the context of our dynamic policies. Section 6 is a conclusion.

Figure 1: Policymaker’s decisions and costs.

2 Model

2.1 Policymaker’s information `policy.pdf`

Let Y_i be a binary random variable denoting whether individual i (he) is infected. Let X_i be a vector of discrete characteristics that are observable and potentially predictive of Y_i . For example, X_i could include whether or not the individual has symptoms related to the disease, whether they have travelled to an infected area, whether they have been in contact with another person who has been infected, or whether they might have already had the disease and therefore built up immunity. We say individuals with characteristics $X_i = x$ are in “group” x .

We denote by Θ_x the true prevalence of the disease among individuals in group x . The true prevalence is unknown and the policymaker (she) has a prior over Θ_x .

After observing the test results of individuals who were previously tested, the policymaker can update her prior of the prevalence of the disease in each group using Bayes’ Theorem. We denote the posterior probability that an individual from group x is infected by \hat{y}_x . Appendix A.1 shows how the posterior probability \hat{y}_x can be calculated from a simple prior used for illustration.

The policymaker makes her decisions having potentially observed a sequence of n individuals, their characteristics, and their outcomes if they have been tested. We denote by n_x the number of individuals from group x who have been tested and by \bar{y}_x the average disease prevalence among these n_x tested individuals. We assume that Θ_x does not change over time. As a result, the policymaker cannot obtain any further information about Θ_x once \bar{y}_x and n_x are known. Constant disease prevalence over time might not be a realistic assumption, but the basic tradeoffs in the model will not be affected by it. We discuss the practical consequences of changing disease prevalence over time in Section 5.

2.2 Policymaker’s choices and costs

The policymaker’s choices when she observes individual i are summarised in Figure 1.

First, the policymaker has two choices: to test the individual ($D_i = 1$) or not to test the individual ($D_i = 0$). Testing someone for the disease comes with a cost of $C \geq 0$. Cost C can either represent the marginal cost of a testing kit or the cost of marginally relaxing the testing capacity constraint (i.e., shadow cost). The test reveals with certainty the value of Y_i and the policymaker observes whether the individual is infected or not.²

Second, the policymaker can quarantine ($Q_i = 1$) or release (not quarantine, $Q_i = 0$) the individual with or without testing. If a test has been conducted, the policymaker can condition her quarantining decision on the observed value of Y_i .

Making wrong decisions (i.e., $Q_i \neq Y_i$) is costly. Falsely quarantining someone who is not infected comes with a cost of $F_Q > 0$.³ Falsely releasing someone who is infected incurs a cost of $F_R > 0$. We normalize the cost of a correct decision (i.e., $Q_i = Y_i$) to 0.⁴ These costs can differ by group x , but we ignore that in our notation for the sake of exposition.

²Imperfect testing does not qualitatively affect the main result (see Section 5).

³We assume that all costs are commensurate and can be measured in a single currency.

⁴The main result is not qualitatively affected by costly correct quarantine decisions (see Section 5).

3 Optimal Myopic Targeted Testing Policy

We now turn to the policymaker's optimal myopic decision. The policymaker takes prior beliefs as given and takes an optimal decision (D_i, Q_i) for individual i in group x having observed a sequence of (the characteristics of all) individuals, testing decisions, and the outcomes for tested individuals, i.e., $(X_j, D_j, D_j Y_j)_{j=1}^n$. This is a two-stage decision problem that can be solved by backward induction.

First, consider the case where a test is conducted ($D_i = 1$) so Y_i is observed. Recall that in this case the policymaker can make the quarantining decision conditional on Y_i . Since $F_Q > 0$ and $F_R > 0$, the optimal decision is to set $Q_i = Y_i$. Total cost incurred in this case is the cost C of testing.

Second, consider the case where no test is conducted ($D_i = 0$) so Y_i is not observed. Recall that having observed prevalence \bar{y}_x in group x , the policymaker's posterior expectation of the prevalence in individual i 's group x is \hat{y}_x . Therefore, the *expected* cost of releasing an untested individual is $\hat{y}_x \cdot F_R$ while the *expected* cost of quarantining an untested individual is $(1 - \hat{y}_x) \cdot F_Q$. Hence, the optimal quarantining decision is to set $Q_i = 1$ if and only if the expected cost of a false release exceeds the expected cost of a false quarantine:

$$\hat{y}_x \cdot F_R \geq (1 - \hat{y}_x) \cdot F_Q,$$

that is, if and only if,

$$\hat{y}_x \geq \frac{F_Q}{F_Q + F_R}.$$

In particular, absent a test, if $F_Q = 0$ the policymaker would quarantine everyone and if $F_R = 0$ the policymaker would not quarantine anyone. Comparing the ex-ante expected costs of testing or not testing, we get that it is optimal to test ($D_i = 1$), if and only if

$$C \leq \min(\hat{y}_x \cdot F_R, (1 - \hat{y}_x) \cdot F_Q),$$

that is, if and only if

$$\hat{y}_x \in \left[\frac{C}{F_R}, 1 - \frac{C}{F_Q} \right].$$

The following proposition summarises the policymaker's optimal myopic policy.⁵

Proposition 1 *The optimal myopic policy for individual i in group x is:*

1. If $\hat{y}_x < \frac{C}{F_R}$, then do not test; release.
2. If $\hat{y}_x \in \left[\frac{C}{F_R}, 1 - \frac{C}{F_Q} \right]$ then test; quarantine if the test is positive; release if the test is negative.
3. If $\hat{y}_x > 1 - \frac{C}{F_Q}$, then do not test; quarantine.

The intuition for this result is as follows. Other things equal, if the cost F_R of a false release in a group increases, the policymaker should start testing individuals who were previously untested (and released) because their likelihood of disease was too low.

On the other hand, if the cost of false quarantine F_Q in a group increases, the policymaker should start testing individuals who were previously untested (and quarantined) because their likelihood of disease was too high.

Lower testing costs symmetrically expand the range of disease likelihoods in which individuals are tested. If $C = 0$, all individuals are tested.

⁵Ties are broken in favour of testing.

Figure 2: Optimal myopic testing and quarantining policy for $C = 1$, $F_R = 5$ and $F_Q = 10$.

Figure 2 provides a concrete illustration of an [optimal myopic.pdf](#) testing policy. In this example, $C = 1$, $F_R = 5$ and $F_Q = 10$. If the policymaker could not test the individual, then she would quarantine the individual if and only if the estimate of the group's disease prevalence is greater than $\frac{10}{10+5} = \frac{2}{3}$. The policymaker tests an individual if her estimate of the group's disease prevalence is between $\frac{1}{5}$ and $\frac{9}{10}$.

4 Adaptive Targeted Testing Policy

The optimal myopic targeted testing policy for individual i ignores any potential value of acquired information for making future decisions. In this section, we consider what happens when the policymaker takes this informational value into account.

Suppose that one individual arrives in every period; thus individual i arrives in period i . The policymaker needs to make a decision (D_i, Q_i) about testing and quarantining this individual immediately, i.e., before the next individual arrives. When making a decision for individual i , the policymaker has access to information $(X_j, D_j, D_j Y_j)_{j=1}^{j < i}$ for all prior individuals $j < i$.

As before when observing an individual i from group x , policymaker needs to make one of two decisions: (i) to test, or (ii) not to test. Denote by R_i the costs associated with the binary testing decision: (i) cost $R_i(1) = C$ of the test, and (ii) the expected cost $R_i(0)$ of the error associated with either quarantining or not quarantining the individual without testing. While the cost of the test is certain, the expected cost of not testing will depend on the prevalence Θ_x of the disease in group x .

Suppose that Θ were known. Then the policymaker will test an individual if the cost of testing is lower than the expected cost of not testing, i.e, if

$$\mathbf{E}[R_i(0) - R_i(1)|\Theta] \geq 0.$$

Recall that if $\hat{y}_x < \frac{F_Q}{F_R + F_Q}$, then the policymaker will release an untested individual; in this case, the policymaker will test if

$$\mathbf{E}[R_i(0) - R_i(1)|\Theta] = C - F_R \cdot \Theta_x \geq 0.$$

Otherwise, the policymaker wants to release an untested individual; in this case, the policymaker will test if

$$\mathbf{E}[R_i(0) - R_i(1)|\Theta] = C - F_Q \cdot (1 - \Theta_x) \geq 0,$$

If the policymaker knew Θ_x , she would be able to take an optimal (in expectation) decision for each individual i in group x . However, the policymaker can only form a posterior over Θ_x given then information she has access to. Armed with this posterior, the policymaker can calculate the probability that her action is, in fact, myopically optimal (see Appendix A.2).

The policymaker's objective is to maximise societal outcomes, i.e., to minimise cumulative expected costs over time. However, the optimal myopic testing policy that "exploits" the full benefit of testing to the current individual (described in Section 3) is not dynamically optimal. The reason is that testing an individual has "exploration" value, i.e., it allows the

polymaker to obtain a more precise estimate of the true prevalence, thereby making better decisions for individuals arriving later. As a result, the polymaker faces an exploration-exploitation tradeoff. In order to explore, the polymaker initially spreads out some of her testing capacity and sacrifices some immediate benefit.

The optimal extent of exploration comes from a solution to a complex dynamic stochastic optimisation problem. But solving for the optimal dynamic testing policy is computationally infeasible. Remarkably, however, a simple policy due to Thompson (1933) turns out to be almost optimal:

**Test individual i with probability equal to
the probability that $D_i = 1$ is myopically optimal.**

The Thompson policy neatly captures the exploration-exploitation tradeoff faced by the polymaker. Initially, the polymaker has little data and exploration is valuable so the probability of a testing decision for any given individual will rely heavily on the polymaker’s prior. As a result, the Thompson policy recommends to spread out testing capacity in the vicinity of the cutoffs for testing under the optimal myopic policy. As the sample size within a group x becomes large, the polymaker learns the true prevalence Θ_x within the group and exploration becomes unnecessary. As a result, Thompson policy—as well as the dynamically optimal policy—coincides with the optimal myopic policy described in Section 3 in the limit when n_x is large.

A spate of recent work has shown that, surprisingly, the expected total cost achieved by Thompson policy asymptotically matches the expected total costs under the fully optimal policy (Agrawal and Goyal, 2012; Kaufmann et al., 2012). Therefore, in large samples the polymaker loses almost nothing by following the Thompson algorithm (see Appendix A.3). The Thompson algorithm is used ubiquitously online in product assortment planning, revenue management, and recommendation systems by companies such as Microsoft, Google, and LinkedIn (see, e.g., (Russo et al., 2018) and is gradually making inroads into economic policy evaluation (Kasy and Sautmann, 2019; Kasy and Teytelboym, 2020; Caria et al., 2020).

4.1 Illustration of myopic vs. dynamic testing policies

We now illustrate how the optimal myopic testing is affected by changes in testing costs and in the costs of false quarantine/release. We also show how the Thompson policy spreads out testing compared to the optimal myopic policy.

We fix $C = 1$ and $n = 50$. Let us consider four types of individuals—nurses, truck drivers, academics, party clowns—whose costs of false release and false quarantine differ as in the table below. Here we assume that the individuals’ types is just one of many characteristics: the types determine the individuals’ relative costs, but not their probability of infection.

	$F_R = 20$	$F_R = 5$
$F_Q = 10$	Nurse	Truck Driver
$F_Q = 3$	Party Clown	Academic

These numbers are purely illustrative; we use them for exposition and to showcase comparative statics of testing policies. Here we imagine that there are many groups of observable characteristics for every type: for example, there are nurses with and without symptoms, academics who have and who have not travelled to a conference in a disease hot spot, etc. We have already considered the optimal myopic testing policy for truck drivers in Section 3.

The optimal myopic testing policies for all four types are illustrated in gray lines in Figures 3a–3d. For example, in an optimal myopic testing policy, the range for testing of nurses (Figure 3c) is greater than the range of testing for academics (Figure 3b) because in our illustration the costs of false quarantine and false release for nurses is greater than that of academics.

Let us first consider the Thompson testing policy for nurses and truck drivers. Figures 3c and 3d show that after 50 observations the Thompson testing policy still smoothly spreads out testing around the optimal myopic testing cutoffs. In order to learn the true prevalence rate which will help make better future decisions, the Thompson policy suggests testing some nurses and truck drivers who would not have been tested under the optimal myopic testing policy (because their infection likelihood is either too high or too low). These tests come at the expense of lowering the probability of testing for some nurses/truck drivers who would have definitely been tested under the optimal myopic policy.

The Thompson sampling policy for party clowns and academics has two interesting further features (see Figures 3a–3b). First, the Thompson policy for academics does not recommend testing any academic with probability 1 (Figure 3b). Second, the probability of testing for academics and party clowns jumps at precisely at the quarantine cutoff in the absence of a test. Intuitively, in the absence of a test the decision changes from not quarantining to quarantining around the cutoff. As a result the probability that testing is optimal also jumps.

In Appendix A.4, we illustrate the Thompson policy when the number of observations is 10 and 500. As more tests have been conducted, the policymaker’s estimate of the prevalence becomes more precise, and the rewards from exploration become smaller. As a result, the Thompson policy becomes closer and closer to the optimal myopic testing policy.

5 Extensions and implementation

Imperfect testing Suppose that the test is imperfect with a false positive rate F_P and a false negative rate F_N . Given the estimated prevalence \hat{y}_x , the expected cost of testing becomes

$$C + \hat{y}_x F_N F_R + (1 - \hat{y}_x) F_P F_Q.$$

The costs of not testing remain the same. Therefore, the policymakers test the individual if

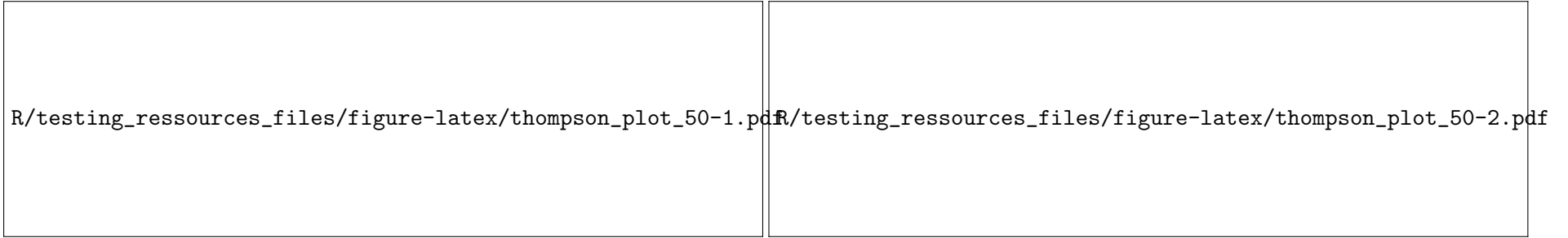
$$C + \hat{y}_x F_N F_R + (1 - \hat{y}_x) F_P F_Q \leq \min(\hat{y}_x \cdot F_R, (1 - \hat{y}_x) \cdot F_Q),$$

that is if

$$\hat{y}_x \in \left[\frac{C + F_P F_Q}{F_R - F_N F_R + F_P F_Q}, \frac{F_Q - C - F_P F_Q}{F_Q + F_N F_R - F_P F_Q} \right],$$

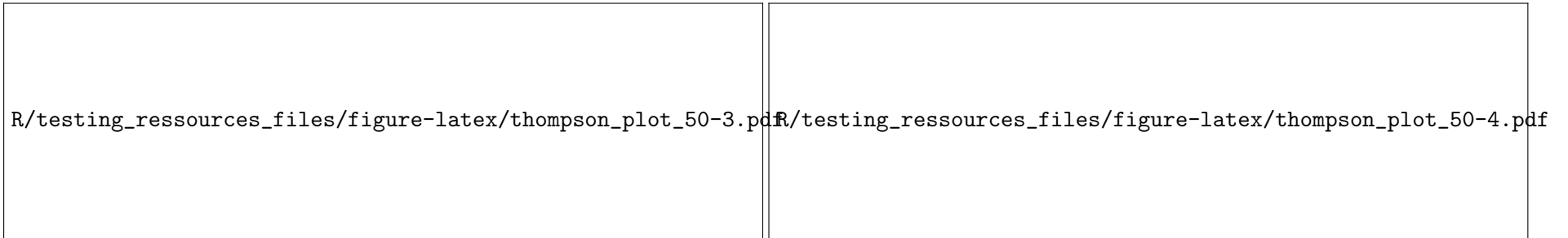
as long as the cutoffs are well-defined given the parameters. In general, the presence of false positive and false negative rates changes both the lower and the upper cutoffs for testing resulting in more or less testing.

Correct quarantine decision is costly Suppose that a correct quarantine decision comes at a cost K , but the correct release decision still has a cost of zero. Then the optimal quarantining decision is to set $Q_i = 1$ if the expected cost of a false release exceeds the



(a) Party clowns. Costs: $C = 1$, $F_Q = 3$, $F_R = 20$.

(b) Academics. Costs: $C = 1$, $F_Q = 3$, $F_R = 5$.



(c) Nurses. Costs: $C = 1$, $F_Q = 10$, $F_R = 20$.

(d) Truck drivers. Costs: $C = 1$, $F_Q = 10$, $F_R = 5$.

Figure 3: Optimal myopic and Thompson testing policies. Solid gray line: optimal myopic testing policy. Dashed gray line: quarantine cutoff in the absence of testing. Solid black line: Thompson policy. Number of observations (n) is 50.

expected cost of a quarantine:

$$\hat{y}_x \cdot F_R \geq (1 - \hat{y}_x) \cdot F_Q + \hat{y}_x \cdot K,$$

that is, if

$$\hat{y}_x \geq \frac{F_Q}{F_Q + F_R - K}.$$

Now it is optimal to test ($D_i = 1$) if

$$C + \hat{y}_x K \leq \min(\hat{y}_x \cdot F_R, (1 - \hat{y}_x) \cdot F_Q + \hat{y}_x \cdot K),$$

that is, if

$$\hat{y}_x \in \left[\frac{C}{F_R - K}, 1 - \frac{C}{F_Q} \right].$$

Therefore, introducing a cost of a correct quarantine decision does not affect the upper bound for testing (as all of these individuals would have been quarantined absent a test), but it increases the lower bound for testing (as testing has become more costly).

Choice of prior In our simulations, we have assumed that the prevalence rates across groups are independent (see Appendix A.1. In practice, the performance of the Thompson policy will, however, depend on the choice of the prior. In particular, we could allow for very complex relationships in the prevalence rates across different groups. If the relationships are incorporated into a prior using existing epidemiological models, the policymaker could learn about prevalence in one group by observing outcomes of another group thereby requiring less exploration. Therefore, a well-specified prior can help the policymaker make better decisions.

Non-stationarity Disease prevalence rates change over time and the process is not stationary. If parameters of the model that tracks the disease spread (e.g., a SIR model) could be accurately estimated, then our methods could be adapted to learning the parameters of such a model. However, there can be a lot of disagreement among experts about the trajectory and extent of disease spread.⁶ Therefore, an alternative is to adapt our methods to an environment with non-stationary prevalence rates, which might for instance follow a random walk. The Thompson policy in such an environment would involve the model “forgetting” older observations which might not be informative of the current state of the world (Russo et al., 2018, Section 6.3). The extent of “forgetting” would depend on the rate at which information about prevalence rates becomes obsolete (Raj and Kalyani, 2017; Besbes et al., 2019)

Ethics of targeting Targeted testing policies target. Disease prevalence as well as costs of false quarantine and false release might well vary across income, race, gender, etc. Policymakers need to make sure their choices of parameters and covariates do not discriminate, especially among the most vulnerable groups. These concerns are not novel to public health experts, but they can go unnoticed when decisions about individuals’ lives are being made using statistical models. Our paper implies that resource allocation can be improved by carefully considering costs and benefits of testing and quarantine within any non-discrimination constraints adopted by the policymaker.

⁶Consider, for example, the difference between two highly influential models for the UK during COVID-19 due to Ferguson et al. (2020) and Lourenço et al. (2020).

Estimating local prevalence How should a policymaker maximise the precision of an estimate of the prevalence rate with a given number of tests? If the policymaker is not concerned about the welfare of the individuals in the experimental sample, then she should sample different groups x in proportion to $\sqrt{\hat{y}_x(1 - \hat{y}_x)}$, i.e., standard deviation of prevalence in the group given its prevalence rate (Neyman, 1934). Therefore, groups with prevalence closer to $\frac{1}{2}$ should be sampled proportionally more. Such stratified testing strategies are particularly useful if the policymaker subsequently uses the estimate to make a decision about a local area lockdown.

6 Conclusion

Testing policies that use testing resources efficiently need to take into account the costs of testing, false quarantine, and false release. Our simple framework illuminated various tradeoffs faced by the policymaker when testing resources are limited. Our testing policies balance the information value of wide-ranging testing with the immediate benefit of testing and quarantining those who are likely to be infected. Practical implementation of our policies does not require any additional statistical sophistication beyond what is typically deployed to fight epidemics, but any application will require careful parameter and model calibration.

References

- Agrawal, S. and N. Goyal (2012). Analysis of thompson sampling for the multi-armed bandit problem. In *Conference on Learning theory*, pp. 39–1.
- Besbes, O., Y. Gur, and A. Zeevi (2019). Optimal exploration–exploitation in a multi-armed bandit problem with non-stationary rewards. *Stochastic Systems* 9(4), 319–337.
- Bubeck, S. and N. Cesa-Bianchi (2012). Regret Analysis of Stochastic and Nonstochastic Multi-armed Bandit Problems. *Foundations and Trends® in Machine Learning* 5(1), 1–122. <http://dx.doi.org/10.1561/22000000024>.
- Caria, S., G. Gordon, M. Kasy, S. Osman, S. Quinn, and A. Teytelboym (2020). Adaptive treatment assignment in experiments for policy choice. Mimeo.
- Ferguson, N., D. Laydon, G. Nedjati Gilani, N. Imai, K. Ainslie, M. Baguelin, S. Bhatia, A. Boonyasiri, Z. Cucunuba Perez, G. Cuomo-Dannenburg, et al. (2020). Report 9: Impact of non-pharmaceutical interventions (NPIs) to reduce COVID19 mortality and healthcare demand.
- Galeotti, A., J. Steiner, and P. Surico (2020, April). An economic approach to testing. Mimeo. <http://home.cerge-ei.cz/steiner/informationvalue.pdf>.
- Gupta, P. (2020, March). Global shortage of coronavirus testing kits critical: Experts. *Al-Jazeera*. <https://www.aljazeera.com/news/2020/03/global-shortage-coronavirus-testing-kits-critical-experts-200323085504252.html>.
- Hasell, J., E. Ortiz-Ospina, E. Mathieu, H. Ritchie, D. Beltekian, B. Macdonald, and M. Roser (2020, May). Our World In Data: Coronavirus (COVID-19) Testing. Technical report. <https://ourworldindata.org/coronavirus-testing>.

- Hope, C. (2020, May). 50,000 coronavirus tests secretly flown to the US after UK lab issues . *The Daily Telegraph*.
- Kasy, M. and A. Sautmann (2019). Adaptive treatment assignment in experiments for policy choice. Technical report. <https://maxkasy.github.io/home/files/papers/adaptiveexperimentpolicy.pdf>.
- Kasy, M. and A. Teytelboym (2020). Adaptive combinatorial allocation policy. Mimeo.
- Kaufmann, E., N. Korda, and R. Munos (2012). Thompson sampling: An asymptotically optimal finite-time analysis. In *International Conference on Algorithmic Learning Theory*, pp. 199–213. Springer.
- Lourenço, J., R. Paton, M. Ghafari, M. Kraemer, C. Thompson, P. Simmonds, P. Klennerman, and S. Gupta (2020). Fundamental principles of epidemic spread highlight the immediate need for large-scale serological surveys to assess the stage of the SARS-CoV-2 epidemic. *medRxiv*.
- Neyman, J. (1934). On the two different aspects of the representative method: The method of stratified sampling and the method of purposive selection. *Journal of the Royal Statistical Society* 97(4), 558–625.
- Oster, E. (2020, April). We Cant Get a Handle on the Coronavirus Pandemic Without Random Testing. *Slate*. <https://slate.com/technology/2020/04/coronavirus-research-random-testing.html>.
- Padula, W. V. (2020, April). Why Only Test Symptomatic Patients? Consider Random Screening for COVID-19. *Applied Health Economics and Health Policy*.
- Raj, V. and S. Kalyani (2017). Taming non-stationary bandits: A Bayesian approach. *arXiv preprint arXiv:1707.09727*.
- Russo, D. and B. Van Roy (2016). An information-theoretic analysis of thompson sampling. *The Journal of Machine Learning Research* 17(1), 2442–2471.
- Russo, D. J., B. V. Roy, A. Kazerouni, I. Osband, and Z. Wen (2018). A Tutorial on Thompson Sampling. *Foundations and Trends® in Machine Learning* 11(1), 1–96. <http://dx.doi.org/10.1561/22000000070>.
- Thompson, W. R. (1933). On the likelihood that one unknown probability exceeds another in view of the evidence of two samples. *Biometrika* 25(3/4), 285–294.
- Weaver, C. and R. Ballhaus (2020, April). Coronavirus Testing Hampered by Disarray, Shortages, Backlogs. *Wall Street Journal*. <https://www.wsj.com/articles/coronavirus-testing-hampered-by-disarray-shortages-backlogs-11587328441>.

A Details of the model

A.1 Policymaker's information

To avoid technical subtleties, we assume that all characteristics are discrete and the distribution of characteristics has finite support.

Let $\Theta_x \in [0, 1]$ be the prevalence of the disease among persons with characteristics (“group”) $X_i = x$. The policymaker does not precisely know the prevalence of the disease among the group. To keep things simple, let us assume that the policymaker's prior over the prevalence is uniformly and independently distributed across groups, i.e.,

$$\Theta \sim \text{Uni}([0, 1]^k)$$

That is, we assume that the policymaker essentially has no prior knowledge of the prevalence of the disease among any group and thinks that the prevalences of the disease are independent across groups. Conditional on the policymaker's prior, the outcomes of group x follow a Bernoulli distribution with parameter Θ_x , i.e.,

$$Y_i | X_i = x, \Theta \sim \text{Ber}(\Theta_x).$$

We assume that the parameters (p_1, \dots, p_k) of the distribution of characteristics are independent of the prevalence vector. Therefore, the policymaker cannot learn about prevalence of disease simply from observing the distribution of characteristics across individuals.

The policymaker observes a sequence $(X_j, D_j, D_j Y_j)_{j=1}^n$ of n individuals, their characteristics, and their outcomes if they have been tested. Formally, we define

$$n_x = \sum_i \mathbf{1}(X_i = x, D_i = 1),$$

and

$$\bar{y}_x = \frac{1}{n_x} \sum_i \mathbf{1}(X_i = x, D_i = 1) Y_i.$$

After observing n individuals, the policymaker can update her prior of the prevalence of the disease in each group. Because the policymaker's prior is uniform, the posterior has the following closed-form:

$$\Theta_x | (X_j, D_j, D_j Y_j)_{j=1}^n \sim \text{Beta}(1 + n_x \bar{y}_x, 1 + n_x(1 - \bar{y}_x)).$$

Since the mean of a random variable distributed as $\text{Beta}(\alpha, \beta)$ is $\frac{\alpha}{\alpha + \beta}$, the expected prevalence \hat{y}_x in group x conditional on observing a sequence of individuals $(Y_i, X_i)_{i=1}^n$ is

$$\hat{y}_x = \mathbf{E} [\Theta_x | (X_j, D_j, D_j Y_j)_{j=1}^n] = \frac{1 + n_x \bar{y}_x}{2 + n_x}.$$

Average prevalence in the observed outcomes is sufficient to pin down the posterior prevalence, so

$$\hat{y}_x = \mathbf{E}[Y_i | X_i = x, \bar{y}_x].$$

A.2 Testing probabilities under the Thompson policy

Recall that the expected difference in rewards between not testing and testing is given by

$$\mathbf{E}[R_i(0) - R_i(1) | \Theta] = C - F_R \cdot \Theta_x$$

if $\hat{y}_x < \frac{F_Q}{F_R + F_Q}$, and by

$$\mathbf{E}[R_i(0) - R_i(1)|\Theta] = C - F_Q \cdot (1 - \Theta_x)$$

otherwise. The posterior distribution of Θ_x is $\text{Beta}(1 + n_x \bar{y}_x, 1 + n_x(1 - \bar{y}_x))$.

Thompson sampling tests with probability equal to the posterior probability that testing is optimal. This posterior probability is given by

$$\mathbf{P}(\mathbf{E}[R_i(0) - R_i(1)|\Theta] < 0).$$

Let $F(y|\alpha, \beta)$ denote the cumulative distribution function of a Beta distribution with parameters α and β , evaluated at y . Then the posterior probability that testing is optimal, if $\hat{y}_x < \frac{F_Q}{F_R + F_Q}$, is given by

$$\mathbf{P}(C - F_R \cdot \Theta_x < 0) = 1 - F\left(\frac{C}{F_R}; 1 + n_x \bar{y}_x, 1 + n_x(1 - \bar{y}_x)\right).$$

The posterior probability that testing is optimal, if $\hat{y}_x \geq \frac{F_Q}{F_R + F_Q}$, is given by

$$\mathbf{P}(C - F_Q \cdot (1 - \Theta_x) < 0) = 1 - F\left(\frac{C}{F_Q}; 1 + n_x(1 - \bar{y}_x), 1 + n_x \bar{y}_x\right).$$

A.3 Regret bound of the Thompson algorithm

The dynamic setup we discuss in this paper can be thought of as a so-called two-armed *contextual bandit* problem. Units arrive sequentially, we observe their group membership X_i (the “context”), and then decide between the two “arms” of testing ($D_i = 1$) or not ($D_i = 0$). Then rewards are realised. The rewards are $-C$ in the case of testing, and either $-F_R Y_i$ (if $\hat{y}_x < \frac{F_Q}{F_R + F_Q}$) or $-F_Q(1 - Y_i)$ (otherwise) in the case of not testing.

Conditional on the quarantining decision absent testing, this is exactly the bandit framework. We can “concentrate out” the quarantining decision when analyzing our setting. With consistency of posteriors the sign of $\hat{y}_x - \frac{F_Q}{F_R + F_Q}$ is the same as the sign of $\Theta_x - \frac{F_Q}{F_R + F_Q}$ in large samples, and theoretical results for bandit problems carry over. The learning problem for the decision maker in this limit then becomes to figure out whether our not Θ_x lies on one or the other side of the myopic cutoffs $\frac{C}{F_R}$ or $1 - \frac{C}{F_Q}$.

A large literature discusses guarantees for the performance of bandit algorithms (e.g., Bubeck and Cesa-Bianchi (2012) and Russo et al. (2018)) and more specifically of Thompson sampling (see, in particular, Agrawal and Goyal (2012) and Russo and Van Roy (2016)). These analyses bound the *regret*—the cumulative difference between realized costs and the costs that could have been achieved by taking optimal decisions given the knowledge of Θ . The guarantees in the literature are distinguished by whether they consider large sample limits for fixed Θ , or worst case bounds over all possible Θ .

For the large sample limit case, Agrawal and Goyal (2012) show that normalized regret $\frac{T}{\log(T)} \bar{R}_T$ converges to a constant equal to the efficiency bound. This constant is larger the smaller the difference between treatment arms since it then takes disproportionately longer to learn which arm is optimal. In our case, the constant is large when Θ is close to either cutoff for the myopic decision rule. We can interpret this bound to mean that only about $\log T/T$ decisions are sub-optimal, relative to those that we would have made given knowledge of Θ . For the worst-case scenario, Russo and Van Roy (2016) show that regret satisfies a finite sample prior-independent bound that grows as \sqrt{T} . This worst-case bound is driven by parameter values that are in a $1/\sqrt{T}$ neighborhood of the cutoffs for the myopic rule.

A.4 Thompson policy for $n = 500$ and $n = 10$

Figures 4a–4d illustrate the Thompson policy after 500 observations.

As we showed in Section A.1,

$$\hat{y}_x = \frac{1 + n_x \bar{y}_x}{2 + n_x},$$

so in small samples the support of estimated prevalence \hat{y}_x can deviate significantly from the support of \bar{y}_x (i.e., $[0, 1]$). Figures 5a–5d illustrate the Thompson policy after 10 observations.

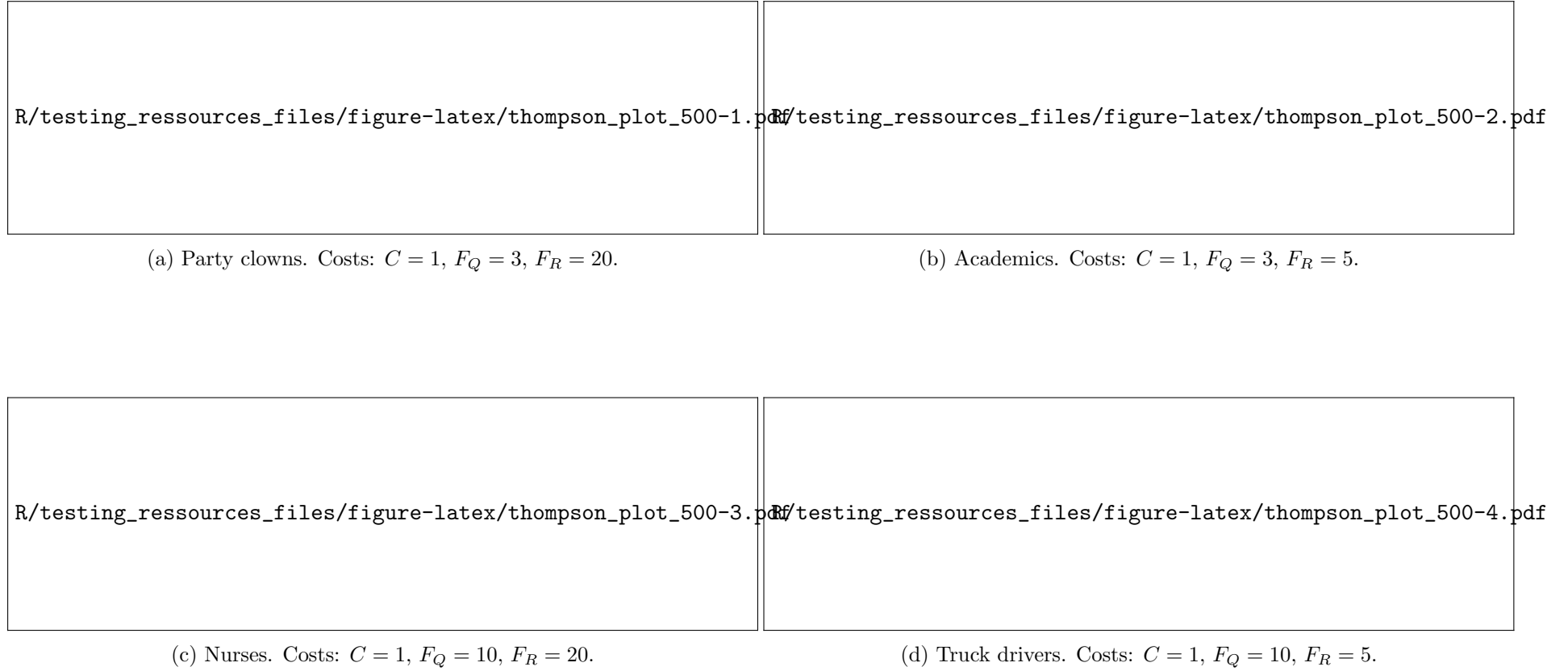


Figure 4: Optimal myopic and Thompson testing policies. Solid gray line: optimal myopic testing policy. Dashed gray line: quarantine cutoff in the absence of testing. Solid black line: Thompson policy. Number of observations (n) is 500.

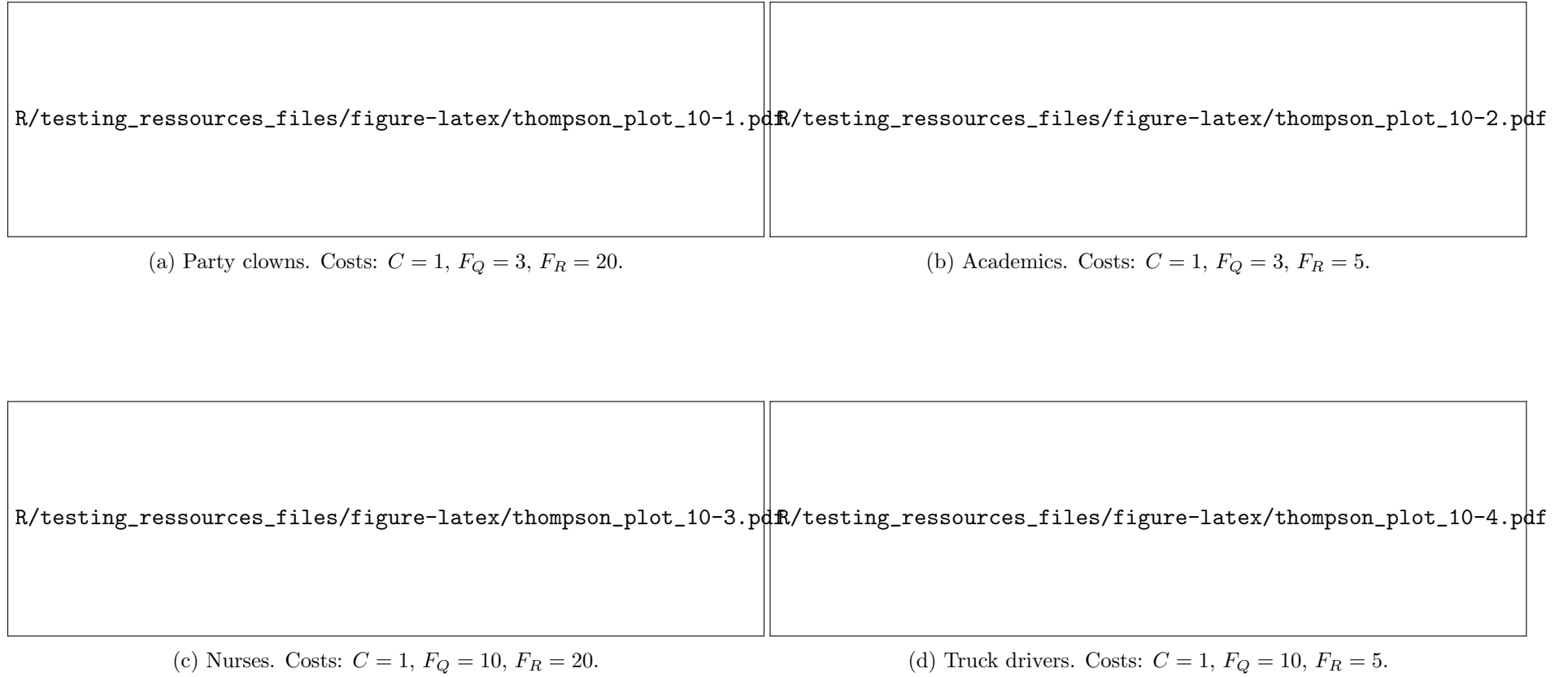


Figure 5: Optimal myopic and Thompson testing policies. Solid gray line: optimal myopic testing policy. Dashed gray line: quarantine cutoff in the absence of testing. Solid black line: Thompson policy. Number of observations (n) is 10.