ESTIMATING COVID-19'S R_t IN REAL-TIME

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Original author: Kevin Systrom—April 17.

Source:

- (1) https://rt.live/(live page)
- (2) https://github.com/k-sys/covid-19 (notebook);

SHA1: cfba8cf5979da87adb7f57d6ff9ff3903d7a4f01

1. Introduction

In any epidemic, R_t is the measure known as the *effective reproduction number*. It's the number of people who become infected per infectious person at time t. The most well-known version of this number is the basic reproduction number: R_0 when t = 0. However, R_0 is a single measure that does not adapt with changes in behavior and restrictions.

As a pandemic evolves, increasing restrictions (or potential releasing of restrictions) changes R_t . Knowing the current R_t is essential. When $R \gg 1$, the pandemic will spread through a large part of the population. If $R_t < 1$, the pandemic will slow quickly before it has a chance to infect many people. The lower the R_t : the more manageable the situation. In general, any $R_t < 1$ means things are under control.

The value of R_t helps us in two ways.

- (1) It helps us understand how effective our measures have been controlling an outbreak and
- (2) It gives us vital information about whether we should increase or reduce restrictions based on our competing goals of economic prosperity and human safety. Well-respected epidemiologists argue that tracking R_t is the only way to manage through this crisis.

Yet, today, we don't yet use R_t in this way. In fact, the only real-time measure I've seen has been for Hong Kong. More importantly, it is not useful to understand R_t at a national level. Instead, to manage this crisis effectively, we need a local (state, county and/or city) granularity of R_t .

What follows is a solution to this problem at the US State level. It's a modified version of a solution created by [Bettencourt and Ribeiro, 2008] to estimate real-time R_t using a Bayesian approach. While this paper estimates a static R value, here we introduce a process model with Gaussian noise to estimate a time-varying R_t .

If you have questions, comments, or improvments feel free to get in touch: hello@systrom.com. And if it's not entirely clear, I'm not an epidemiologist. At the same time, data is data, and statistics are statistics and this is based on work by well-known epidemiologists so you can calibrate your beliefs as you wish. In the meantime, I hope you can learn something new as I did by reading through this example. Feel free to take this work and apply it elsewhere – internationally or to counties in the United States.

Additionally, a huge thanks to Frank Dellaert who suggested the addition of the process and to Adam Lerer who implemented the changes. Not only did I learn something new, it made the model much more responsive.

2. Bettencourt & Ribeiro's Approach

Every day, we learn how many more people have COVID-19. This new case count gives us a clue about the current value of R_t . We also, figure that the value of R_t today is related to the value of R_{t-1} (yesterday's value) and every previous value of R_{t-m} for that matter.

With these insights, the authors use Bayes' rule to update their beliefs about the true value of R_t based on how many new cases have been reported each day.

This is Bayes' Theorem as we'll use it:

$$Pr(R_t \mid k) = \frac{Pr(k \mid R_t) \cdot Pr(R_t)}{Pr(k)}$$

This says that, having seen k new cases, we believe the distribution of R_t is equal to:

- The **likelihood** of seeing k new cases given R_t times ...
- The **prior** beliefs of the value of $Pr(R_t)$ without the data ...
- divided by the probability of seeing this many cases in general.

This is for a single day. To make it iterative: every day that passes, we use yesterday's prior $Pr(R_{t-1})$ to estimate today's prior $Pr(R_t)$. We will assume the distribution of R_t to be a Gaussian centered around R_{t-1} , so

$$Pr(R_t \mid R_{t-1}) = \mathcal{N}(R_{t-1}, \sigma),$$

where σ is a hyperparameter (see below on how we estimate σ). So on day one:

$$\Pr(R_1|k_1) \propto \Pr(R_1) \cdot \mathcal{L}(R_1 \mid k_1)$$

On day two:

$$\begin{aligned} \Pr(R_2|k_1, k_2) &\propto \Pr(R_2) \cdot \mathcal{L}(R_2|k_2) \\ &= \sum_{R_1} \Pr(R_1|k_1) \cdot \Pr(R_2|R_1) \cdot \mathcal{L}(R_2|k_2) \end{aligned}$$

etc.

3. Choosing a Likelihood Function $Pr(k_t | R_t)$

A likelihood function function says how likely we are to see k new cases, given a value of R_t .

Any time you need to model "arrivals" over some time period of time, statisticians like to use the Poisson Distribution. Given an average arrival rate of λ new cases per day, the probability of seeing k new cases is distributed according to the Poisson distribution:

$$\Pr(k \mid \lambda) = \frac{\lambda^k e^{-\lambda}}{k!}$$

```
# Column vector of k
k = np.arange(0, 70)[:, None]

# Different values of Lambda
lambdas = [10, 20, 30, 40]

# Evaluated the Probability Mass Function (remember: poisson is discrete)
y = sps.poisson.pmf(k, lambdas)

# Show the resulting shape
print(y.shape)
```

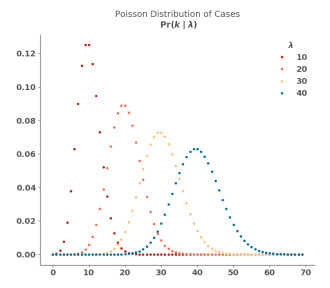
```
(70, 4)
```

Note: this was a terse expression which makes it tricky. All I did was to make k a column. By giving it a column for k and a "row" for lambda it will evaluate the pmf over both and produce an array that has k rows and lambda columns. This is an efficient way of producing many distributions all at once, and **you will see it used again below!**

```
def plot_it(filename):
           fig, ax = plt.subplots(figsize=(8,6.5))
           ax.set(title='Poisson Distribution of Cases\n $\Pr(k\mid\lambda)$')
          plt.plot(k, y,
5
           marker='o'
6
               markersize=3.
               1w=0
8
          plt.legend(title="$\lambda$", labels=lambdas)
9
           fig.savefig(filename)
10
11
           return filename
12
      plot_it(filename='./img/systrom-1.png')
13
```

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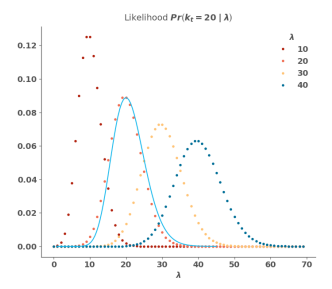


The Poisson distribution says that if you think you're going to have λ cases per day, you'll probably get that many, plus or minus some variation based on chance.

But in our case, we know there have been k cases and we need to know what value of λ is most likely. In order to do this, we fix k in place while varying λ . **This is called the likelihood function.**

For example, imagine we observe k = 20 new cases, and we want to know how likely each λ is:

```
k = 20
       lam = np.linspace(1, 45, 90)
3
       likelihood = pd.Series(
           data=sps.poisson.pmf(k, lam),
           index=pd.Index(lam, name='$\lambda$'),
name='lambda'
5
6
7
8
       def plot_it(filename):
9
            likelihood.plot(
10
                title=r'Likelihood $Pr\left(k_t=20\mid\lambda\right)$',
figsize=(8,6.5)
11
12
13
           plt.legend(title="$\lambda$", labels=lambdas)
14
15
           plt.savefig(filename)
            return filename
16
17
       plot_it(filename='./img/systrom-2.png')
18
```



This says that if we see 20 cases, the most likely value of λ is (not surprisingly) 20. But we're not certain: it's possible λ was 21 or 17 and saw 20 new cases by chance alone. It also says that it's unlikely λ was 40 and we saw 20.

Great. We have $Pr(\lambda_t \mid k_t)$ which is parameterized by λ but we were looking for $Pr(k_t \mid R_t)$ which is parameterized by R_t . We need to know the relationship between λ and R_t

4. Connecting λ and R_t

The key insight to making this work is to realize there's a connection between R_t and λ . The derivation is beyond the scope of this notebook, but here it is:

$$\lambda = k_{t-1} e^{\gamma(R_t - 1)}$$

where γ is the reciprocal of the serial interval (about 7 days for COVID19). Since we know every new case count on the previous day, we can now reformulate the likelihood function as a Poisson parameterized by fixing k and varying R_t .

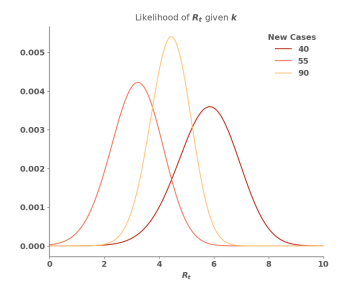
$$\Pr\left(k \mid R_t\right) = \frac{\lambda^k e^{-\lambda}}{k!}$$

5. Evaluating the Likelihood Function

To continue our example, let's imagine a sample of new case counts k. What is the likelihood of different values of R_t on each of those days?

```
k = np.array([20, 40, 55, 90])
2
       # We create an array for every possible value of Rt
4
      R\_T\_MAX \ = \ {\color{red} 12}
      r_t_n = np.linspace(0, R_T_MAX, R_T_MAX*100+1)
6
       # Gamma is 1/serial interval
       # https://wwwnc.cdc.gov/eid/article/26/7/20-0282_article
       # https://www.nejm.org/doi/full/10.1056/NEJMoa2001316
10
11
       # Map Rt into lambda so we can substitute it into the equation below
12
13
      \# Note that we have N-1 lambdas because on the first day of an outbreak
14
       # you do not know what to expect.
      lam = k[:-1] * np.exp(GAMMA * (r_t_range[:, None] - 1))
```

```
16
17
       # Evaluate the likelihood on each day and normalize sum of each day to 1.0
       likelihood_r_t = sps.poisson.pmf(k[1:], lam)
18
19
       likelihood_r_t /= np.sum(likelihood_r_t, axis=0)
20
       def plot_it(filename):
21
22
23
           ax = pd.DataFrame(
24
               data = likelihood_r_t,
25
               index = r_t_range
26
           ).plot(
27
               title='Likelihood of $R_t$ given $k$',
28
               xlim=(0,10),
               figsize=(8,6.5)
29
30
31
           ax.legend(labels=k[1:], title='New Cases')
           ax.set_xlabel('$R_t$');
32
33
           plt.savefig(filename)
           return filename
34
35
       plot_it(filename='./img/systrom-3.png')
```



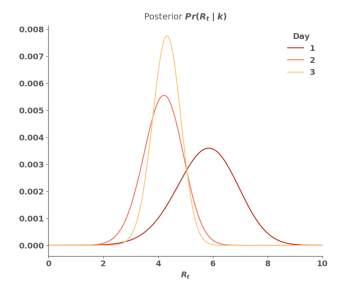
You can see that each day we have a independent guesses for R_t . The goal is to combine the information we have about previous days with the current day. To do this, we use Bayes' theorem.

6. Performing the Bayesian Update

To perform the Bayesian update, we need to multiply the likelihood by the prior (which is just the previous day's likelihood without our Gaussian update) to get the posteriors. Let's do that using the cumulative product of each successive day:

```
posteriors = likelihood_r_t.cumprod(axis=1)
      posteriors = posteriors / np.sum(posteriors, axis=0)
2
3
      columns = pd.Index(range(1, posteriors.shape[1]+1), name='Day')
4
5
      posteriors = pd.DataFrame(
                  = posteriors,
6
          data
          index = r_t_range,
7
          columns = columns
8
9
10
11
      def plot_it(filename):
12
13
          ax = posteriors.plot(
```

```
14
               title='Posterior $Pr(R_t \mid k)$',
15
               xlim=(0,10),
               figsize=(8,6.5)
16
17
           ax.legend(title='Day')
18
           ax.set_xlabel('$R_t$');
19
20
           plt.savefig(filename)
21
           return filename
22
      plot_it(filename='./img/systrom-4.png')
```



Notice how on Day 1, our posterior matches Day 1's likelihood from above? That's because we have no information other than that day. However, when we update the prior using Day 2's information, you can see the curve has moved left, but not nearly as left as the likelihood for Day 2 from above. This is because Bayesian updating uses information from both days and effectively averages the two. Since Day 3's likelihood is in between the other two, you see a small shift to the right, but more importantly: a narrower distribution. We're becoming **more** confident in our believes of the true value of R_t .

From these posteriors, we can answer important questions such as:

"What is the most likely value of R_t each day?"

```
most_likely_values = posteriors.idxmax(axis=0)
print(most_likely_values)
```

```
Day
1 5.85
2 4.22
3 4.33
dtype: float64
```

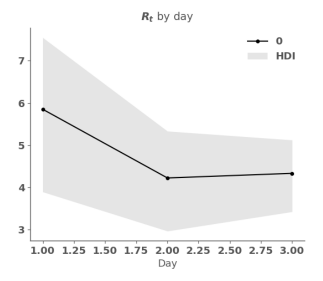
We can also obtain the highest density intervals for R_t :

```
hdi = highest_density_interval(posteriors, debug=True)
print( f'.\n{hdi.tail()}' )
```

```
Low_90 High_90
Day
1 3.89 7.55
2 2.96 5.33
3 3.42 5.12
```

Finally, we can plot both the most likely values for R_t and the HDIs over time. This is the most useful representation as it shows how our beliefs change with every day.

```
def plot_it(filename):
2
              need a pd.DataFrame here
            most_likely = pd.DataFrame(
                most_likely_values
4
5
            fig = plt.figure(figsize=(8, 6.5))
6
           ax = most_likely.plot(
                marker='o',
8
                label='Most Likely',
9
                title=f'$R_t$ by day',
10
                c='k',
11
                markersize=4,
12
13
           ax.fill_between(
14
                hdi.index,
hdi['Low_90'],
hdi['High_90'],
15
16
17
18
                color='k',
19
                alpha=.1,
20
                1w=0,
                label='HDI'
21
22
23
            ax.legend();
            plt.savefig(filename)
24
25
            return filename
26
27
       plot_it(filename='./img/systrom-5.png')
```



We can see that the most likely value of R_t changes with time and the highest-density interval narrows as we become more sure of the true value of R_t over time. Note that since we only had four days of history, I did not apply the process to this sample. Next, however, we'll turn to a real-world application where this process is necessary.

7. REAL-WORLD APPLICATION TO US DATA

7.1. **Setup.** Load US state case data from CovidTracking.com

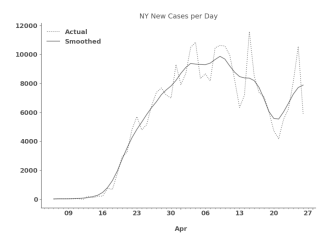
```
url = 'https://covidtracking.com/api/v1/states/daily.csv'
states = pd.read_csv(
    url,
    usecols=['date', 'state', 'positive'],
    parse_dates=['date'],
    index_col=['state', 'date'],
    squeeze=True
).sort_index()
```

Taking a look at the state, we need to start the analysis when there are a consistent number of cases each day. Find the last zero new case day and start on the day after that.

Also, case reporting is very erratic based on testing backlogs, etc. To get the best view of the 'true' data we can, I've applied a gaussian filter to the time series. This is obviously an arbitrary choice, but you'd imagine the real world process is not nearly as stochastic as the actual reporting.

```
state_name = 'NY'
cases = states.xs(state_name).rename(f'{state_name} cases')
original, smoothed = prepare_cases(cases)
```

```
def make_plottable_df(series, name):
            date = pd.to_datetime(
                series.index, infer_datetime_format=True,
3
                errors='coerce'
5
            df = pd.DataFrame(
6
                {name: series.values}
8
9
            df.index = date
10
            return df
11
       def plot_it(filename):
12
            df_o = make_plottable_df(original, 'Actual')
df_s = make_plottable_df(smoothed, 'Smoothed')
13
14
            df = pd.concat([df_o, df_s], axis=1)
15
            fig, ax = plt.subplots()
for cn in df.keys(): df[cn].plot(
16
17
18
                ax=ax.
                c='k'
19
                linestyle = ':' if cn == 'Actual' else '-',
20
21
                alpha=.5.
                title=f"{state_name} New Cases per Day",
22
23
                legend=True,
24
                figsize=(10, 6.5)
25
            ax.get_figure().set_facecolor('w')
26
27
            plt.savefig(filename)
28
            return filename
29
30
       plot_it(filename='./img/systrom-6-a.png')
```



7.2. Running the Algorithm.

7.2.1. Choosing the Gaussian σ for $Pr(R_t \mid R_{t-1})$. Note: you can safely skip this section if you trust that we chose the right value of σ for the process below. Otherwise, read on.

The original approach simply selects yesterday's posterior as today's prior. While intuitive, doing so doesn't allow for our belief that the value of R_t has likely changed from yesterday. To allow for that change, we apply Gaussian noise to the prior distribution with some standard deviation σ . The higher σ the more noise and the more we will expect the value of R_t to drift each day. Interestingly, applying noise on noise iteratively means that there will be a natural decay of distant posteriors. This approach has a similar effect of windowing, but is more robust and doesn't arbitrarily forget posteriors after a certain time like my previous approach. Specifically, windowing computed a fixed R_t at each time t that explained the surrounding t0 days of cases, while the new approach computes a series of t1 values that explains all the cases, assuming that t2 fluctuates by about t3 each day.

However, there's still an arbitrary choice: what should σ be? Adam Lerer pointed out that we can use the process of maximum likelihood to inform our choice. Here's how it works:

Maximum likelihood says that we'd like to choose a σ that maximizes the likelihood of seeing our data k: $P(k|\sigma)$. Since σ is a fixed value, let's leave it out of the notation, so we're trying to maximize P(k) over all choices of σ .

Since $P(k) = P(k_0, k_1, ..., k_t) = P(k_0)P(k_1)...P(k_t)$ we need to define $P(k_t)$. It turns out this is the denominator of Bayes rule:

$$Pr(R_t \mid k_t) = \frac{Pr(k_t \mid R_t) Pr(R_t)}{Pr(k_t)}$$

To calculate it, we notice that the numerator is actually just the joint distribution of k and R:

$$Pr(k_t, R_t) = Pr(k_t \mid R_t) Pr(R_t)$$

We can marginalize the distribution over R_t to get $Pr(k_t)$:

$$Pr(k_t) = \sum_{R_t} Pr(k_t \mid R_t) Pr(R_t)$$

So, if we sum the distribution of the numerator over all values of R_t , we get $Pr(k_t)$. And since we're calculating that anyway as we're calculating the posterior, we'll just keep track of it separately.

Since we're looking for the value of σ that maximizes Pr(k) overall, we actually want to maximize:

$$\prod_{t,i} p(k_{ti})$$

where t are all times and i is each state.

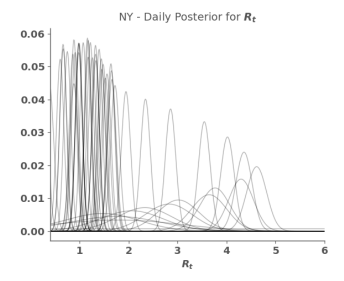
Since we're multiplying lots of tiny probabilities together, it can be easier (and less error-prone) to take the log of the values and add them together. Remember that $\log ab = \log a + \log b$. And since logarithms are monotonically increasing, maximizing the sum of the log of the probabilities is the same as maximizing the product of the non-logarithmic probabilities for any choice of σ .

- 7.3. **Function for Calculating the Posteriors.** To calculate the posteriors we follow these steps:
 - (1) Calculate λ , the expected arrival rate for every day's poisson process;
 - (2) Calculate each day's likelihood distribution over all possible values of R_t ;
 - (3) Calculate the process matrix based on the value of σ we discussed above;
 - (4) Calculate our initial prior because our first day does not have a previous day from which to take the posterior
 - Based on info from the cdc we will choose a Gamma with mean 7.
 - (5) Loop from day 1 to the end, doing the following:
 - Calculate the prior by applying the Gaussian to yesterday's prior.
 - Apply Bayes' rule by multiplying this prior and the likelihood we calculated in step 2.
 - Divide by the probability of the data (also Bayes' rule)

```
# Note that we're fixing sigma to a value just for the example
posteriors, log_likelihood = get_posteriors(smoothed, sigma=.25, gamma=GAMMA)
```

7.4. **The Result.** Below you can see every day (row) of the posterior distribution plotted simultaneously. The posteriors start without much confidence (wide) and become progressively more confident (narrower) about the true value of R_t .

```
def plot_it(filename):
           ax = posteriors.plot(
               title=f'{state_name} - Daily Posterior for $R_t$',
               legend=False,
               lw=1,
c='k',
6
               alpha=.3,
8
               xlim=(0.4,6)
10
           ax.set_xlabel('$R_t$')
11
           plt.savefig(filename)
           return filename
12
13
      plot_it(filename='./img/systrom-7.png')
```



7.5. Plotting in the Time Domain with Credible Intervals. Since our results include uncertainty, we'd like to be able to view the most likely value of R_t along with its highest-density interval.

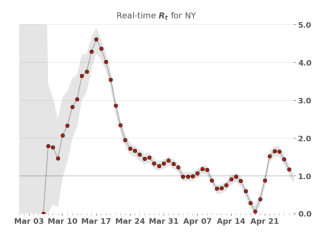
```
hdis = highest_density_interval(posteriors, p=.9)
most_likely = posteriors.idxmax().rename('ML')

# Look into why you shift -1
result = pd.concat([most_likely, hdis], axis=1)
print(f'.\n{result.tail()}')
```

```
ML Low_90 High_90
date
2020-04-22
          1.52
                   1.36
                            1.65
2020-04-23 1.65
                   1.51
                            1.78
2020-04-24 1.64
                   1.50
                            1.76
2020-04-25 1.44
                   1.29
                            1.55
2020-04-26 1.17
                   1.04
                            1.29
```

```
def plot_it(filename):
    fig, ax = plt.subplots(figsize=(600/72,400/72))
    plot_rt(result, ax, state_name, fig)
    ax.set_title(f'Real-time $R.t$ for {state_name}')
    ax.xaxis.set_major_locator(mdates.WeekdayLocator())
    ax.xaxis.set_major_formatter(mdates.DateFormatter('%b %d'))
    plt.savefig(filename)
    return filename

plot_it(filename='./img/systrom-8.png')
```



7.6. Choosing the optimal σ . In the previous section we described choosing an optimal σ , but we just assumed a value. But now that we can evaluate each state with any sigma, we have the tools for choosing the optimal σ .

Above we said we'd choose the value of σ that maximizes the likelihood of the data P(k). Since we don't want to overfit on any one state, we choose the sigma that maximizes P(k) over every state. To do this, we add up all the log likelihoods per state for each value of sigma then choose the maximum.

Note: this takes a while!

```
states.loc[('MA', pd.Timestamp('2020-04-20'))] = 39643
```

```
print( f".\n{states.xs('MA').diff().tail()}" )
```

```
.
date
2020-04-22 1745.0
2020-04-23 3079.0
2020-04-24 0.0
2020-04-25 7325.0
2020-04-26 1590.0
Name: positive, dtype: float64
```

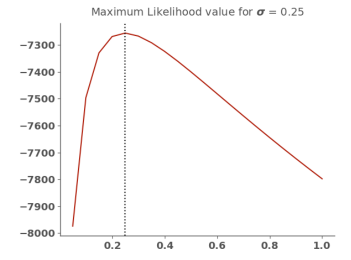
```
sigmas = np.linspace(1/20, 1, 20)
       targets = ~states.index.get_level_values('state').isin(FILTERED_REGION_CODES)
       states_to_process = states.loc[targets]
      if not 'results' in locals() or not len(results):
           results = {}
           for state_name, cases in states_to_process.groupby(level='state'):
              print(state_name)
10
               new, smoothed = prepare_cases(cases, cutoff=25)
11
               if len(smoothed) == 0:
12
                  new, smoothed = prepare_cases(cases, cutoff=10)
13
               # Holds all posteriors with every given value of sigma
               result['posteriors'] = []
               # Holds the log likelihood across all k for each value of sigma
16
               result['log_likelihoods'] = []
17
18
               for sigma in sigmas:
19
                   posteriors, log_likelihood = get_posteriors(smoothed, sigma=sigma)
                   result['posteriors'].append(posteriors)
20
                   result['log_likelihoods'].append(log_likelihood)
```

```
# Store all results keyed off of state name
results[state_name] = result
print('Done.')
```

```
Done.
```

Now that we have all the log likelihoods, we can sum for each value of sigma across states, graph it, then choose the maximum.

```
# Each index of this array holds the total of the log likelihoods for
       # the corresponding index of the sigmas array
      total_log_likelihoods = np.zeros_like(sigmas)
       # Loop through each state's results and add the log likelihoods to the running total.
      for state_name, result in results.items():
           total_log_likelihoods += result['log_likelihoods']
       # Select the index with the largest log likelihood total
      max_likelihood_index = total_log_likelihoods.argmax()
10
11
       # Select the value that has the highest log likelihood
12
13
      sigma = sigmas[max_likelihood_index]
14
      def plot_it(filename):
           fig, ax = plt.subplots()
15
           ax.set_title(f"Maximum Likelihood value for $\sigma$ = {sigma:.2f}");
16
17
           ax.plot(sigmas, total_log_likelihoods)
           ax.axvline(sigma, color='k', linestyle=":")
18
19
           plt.savefig(filename)
          return filename
20
21
22
      plot_it(filename='./img/systrom-9.png')
```



7.7. Compile Final Results. Given that we've selected the optimal σ , let's grab the precalculated posterior corresponding to that value of σ for each state. Let's also calculate the 90% and 50% highest density intervals (this takes a little while) and also the most likely value.

```
if not 'final_results' in locals() or final_results is None:
    final_results = None

for state_name, result in results.items():
    print(f'{state_name} ')
    posteriors = result['posteriors'][max_likelihood_index]
```

```
hdis_90 = highest_density_interval(posteriors, p=.9)
hdis_50 = highest_density_interval(posteriors, p=.5)
most_likely = posteriors.idxmax().rename('ML')
result = pd.concat([most_likely, hdis_90, hdis_50], axis=1)
if final_results is None:
final_results = result
else:
final_results = pd.concat([final_results, result])

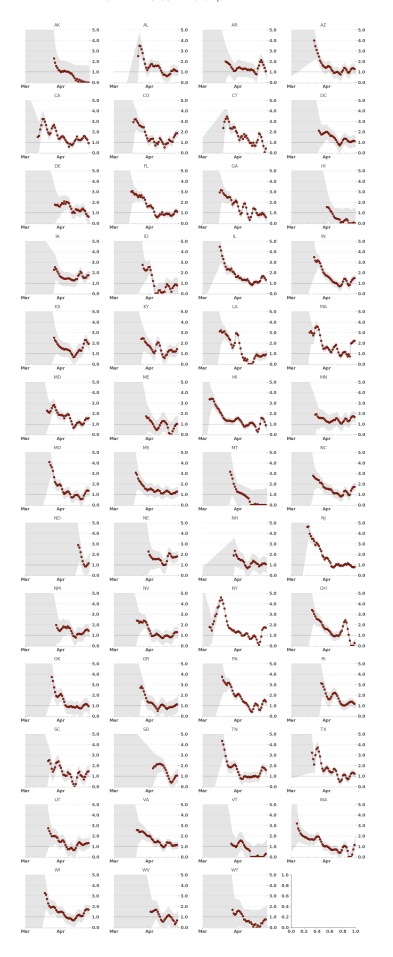
print('Done.')
```

```
Done.
```

```
ncols = 4
nrows = int(np.ceil(len(results) / ncols))

def plot_it(filename):
    fig, axes = plt.subplots(nrows=nrows, ncols=ncols, figsize=(15, nrows*3))
    for i, (state_name, result) in enumerate(final_results.groupby('state')): plot_rt(result.iloc[1:], axes.flat[i], state_name, fig)
    fig.tight_layout()
    fig.set_facecolor('w')
    plt.savefig(filename)
    return filename

plot_it(filename='./img/systrom-10.png')
```



```
# Since we now use a uniform prior, the first datapoint is pretty bogus, so just truncating it here final_results = final_results.groupby('state').apply(lambda x: x.iloc[1:].droplevel(0))
```

7.9. **Export Data to CSV.** Create a data folder:

```
mkdir -p ./data
```

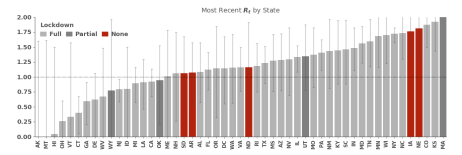
Export the data the data folder:

```
filtered = final_results.index.get_level_values(0).isin(FILTERED_REGION_CODES)
mr = final_results.loc[~filtered].groupby(level=0)[['ML', 'High_90', 'Low_90']].last()

def plot_it(filename):
    mr.sort_values('ML', inplace=True)
    plot_standings(mr, figsize=(16, 4.5));
    plt.savefig(filename)
    return filename

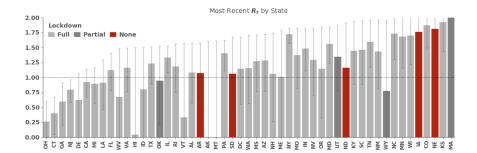
plot_it(filename='./img/systrom-11.png')
```

7.10. Standings.



```
def plot_it(filename):
    mr.sort_values('High_90', inplace=True)
    plot_standings(mr, figsize=(16, 4.5));
    plt.savefig(filename)
    return filename

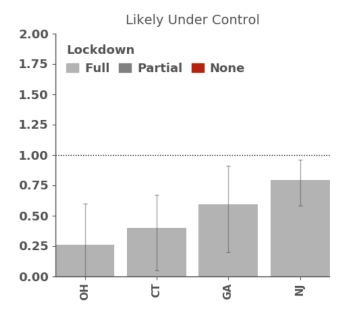
plot_it(filename='./img/systrom-12.png')
```



```
def plot_it(filename):
    show = mr[mr.High_90.le(1)].sort_values('ML')
    fig, ax = plot_standings(show, figsize=(5.25,4.5), title='Likely Under Control');
    plt.savefig(filename)
```

```
return filename

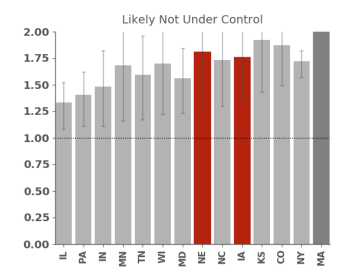
plot_it(filename='./img/systrom-13.png')
```



```
show = mr[mr.High_90.le(1)].sort_values('ML')
fig, ax = plot_standings(show, title='Likely Under Control');

def plot_it(filename):
    show = mr[mr.Low_90.ge(1.0)].sort_values('Low_90')
    fig, ax = plot_standings(show, figsize=(6, 4.5), title='Likely Not Under Control');
    ax.get_legend().remove()
    plt.savefig(filename)
    return filename

plot_it(filename='./img/systrom-14.png')
```



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REFERENCES

[Bettencourt and Ribeiro, 2008] Bettencourt, L. M. A. and Ribeiro, R. M. (2008). Real Time Bayesian Estimation of the Epidemic Potential of Emerging Infectious Diseases. PLOS ONE 5.

APPENDIX A. BUILD: PYTHON PACKAGES

pd.show_versions()

```
INSTALLED VERSIONS
commit: None
python: 3.7.1.final.0
python-bits: 64
OS: Linux
OS-release: 5.3.0-46-generic
machine: x86_64
processor: x86_64
byteorder: little
LC_ALL: None
LANG: en_US.UTF-8
LOCALE: en_US.UTF-8
pandas: 0.23.4
pytest: 4.0.2
pip: 18.1
setuptools: 40.6.3
Cython: 0.29.2
numpy: 1.15.4
scipy: 1.1.0
pyarrow: None
xarray: None
IPython: 7.2.0
sphinx: 1.8.2
patsy: 0.5.1
dateutil: 2.7.5
pytz: 2018.7
blosc: None
bottleneck: 1.2.1
tables: 3.4.4
numexpr: 2.6.8
feather: None
matplotlib: 3.0.2 openpyxl: 2.5.12
xlrd: 1.2.0
xlwt: 1.3.0
xlsxwriter: 1.1.2
lxml: 4.2.5
bs4: 4.6.3
html5lib: 1.0.1
sqlalchemy: 1.0.1
sqlalchemy: 1.2.15
pymysql: None
psycopg2: None
jinja2: 2.10
s3fs: None
fastparquet: None
pandas_gbq: None
pandas_datareader: None
```