

# Lecture 8: Advanced Topics: Advanced Forecasting Methods, Ensembling, and Calibration

Introduction to Time Series, Fall 2023

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## 1 Advanced forecasters

- Thus far, we've learned in-depth about ARIMA and ETS as our two major forecasting frameworks. Undoubtedly, these are battle-tested frameworks that have been used for decades and will take you far, albeit with proper scrutiny
- Of course, there are actually many other forecasting methods out there, and this continues to be an active topic of research. Below, we briefly summarize three other forecasting methods, chosen based on their popularity (they also seem to constitute a common cast of characters, together with ARIMA and ETS, in R and Python forecasting packages)

### 1.1 Theta model

- First on our list is the *Theta model*, proposed by [Assimakopoulos and Nikolopoulos \(2000\)](#). This is on our list not as an advanced forecaster per se (as you will see, it's actually quite simple)
- Instead, it is a simple and popular method that began gaining a lot of attention in parts of the forecasting community, and is closely connected to something you already know: exponential smoothing
- Our presentation here follows [Hyndman and Billah \(2003\)](#), who give a nice, clear perspective on the Theta method and its connection to exponential smoothing
- Given data  $x_t$ ,  $t = 1, 2, 3, \dots$ , the Theta method starts by defining a smoothed sequence  $y_{\theta,t}$ ,  $t = 1, 2, 3, \dots$  by the second-order difference equation:

$$\Delta^2 y_{\theta,t} = \theta \Delta^2 x_t$$

Here  $\Delta^2 = (1 - B)^2$  is the second-order difference operator, just like in our ARIMA lecture, and  $\theta \geq 0$  is a parameter. The solution to the above is given by

$$y_{\theta,t} = a_{\theta} + b_{\theta}t + \theta x_t$$

for some (constant in time) intercept and slope parameters  $a_{\theta}, b_{\theta}$

- For fixed  $\theta$ , the intercept and slope parameters are fit by minimizing the sum of squared errors to the original sequence

$$\min_{a_{\theta}, b_{\theta}} \sum_{t=1}^n (x_t - y_{\theta,t})^2 \iff \min_{a_{\theta}, b_{\theta}} \sum_{t=1}^n \left( (1 - \theta)x_t - a_{\theta} - b_{\theta}t \right)^2$$

which is simply a linear regression of  $(1 - \theta)x_t$  on time  $t$

- Once these estimates  $\hat{a}_{\theta}, \hat{b}_{\theta}$  are found, forecasts are made by running simple exponential smoothing (SES) on the sequence

$$\hat{y}_{\theta,t} = \hat{a}_{\theta} + \hat{b}_{\theta}t + \theta x_t$$

if  $\theta > 0$ , or by extrapolating the line  $\hat{a}_{\theta} + \hat{b}_{\theta}t$  forward in time if  $\theta = 0$

- [Assimakopoulos and Nikolopoulos \(2000\)](#) make the following general recommendation:
  - produce forecasts  $\hat{y}_{0,t+h|t}$  with  $\theta = 0$  (recall this is just extending the line  $\hat{a}_0 + \hat{b}_0 t$ );
  - produce forecasts  $\hat{y}_{2,t+h|t}$  with  $\theta = 2$  (recall this is given by just running SES on  $\hat{y}_{2,t}$ )
  - return their average:  $\hat{y}_{t+h|t} = (\hat{y}_{0,t+h|t} + \hat{y}_{2,t+h|t})/2$ .

Seasonality, if present, is estimated and removed before running this procedure, and added back in at the end. Figure 1 gives a visualization of the canonical “Theta lines”:  $\hat{y}_{0,t}$  and  $\hat{y}_{2,t}$ , for  $\theta = 0, 2$

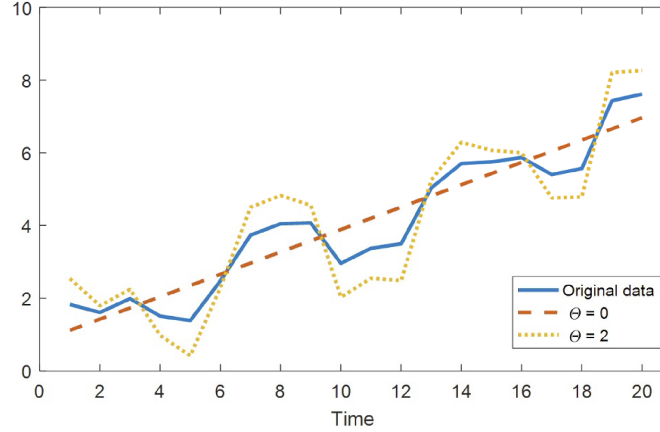


Figure 1: The canonical “Theta lines”, from [Dudek \(2019\)](#).

- [Hyndman and Billah \(2003\)](#) show that this procedure is quite similar to running SES with an added linear trend, whose slope is  $\hat{b}_0/2$ . This is like a special case of Holt’s linear trend method with  $\beta = 0$  (no evolution of the slope over time)
- These authors also compare the Theta method with Holt’s linear trend method on the data from the M3 forecasting challenge (where the Theta method performed well and was subsequently thrown into the spotlight), and observe that Holt’s linear trend performs competitively
- It is worth knowing about the close connections between Theta and exponential smoothing, as much of the literature on the Theta model does not seem to emphasize this aspect. There has been more recent work on the Theta model (which may further distinguish it from exponential smoothing—we cannot say, because we have not followed it) that you may be interested in reading up on

## 1.2 Prophet model

- Next on our list is the *Prophet model*, proposed by [Taylor and Letham \(2018\)](#), from Facebook. This has become popular for large-scale forecasting enterprises, and the popular opinion seems to be that its advantages over traditional ARIMA or ETS models are twofold: flexibility and speed. But make sure to read on, especially to the end of this subsection, for further discussion of this
- The Prophet model is a particular type of signal plus noise model,

$$x_t = g_t + s_t + h_t + \epsilon_t$$

where  $\epsilon_t$ ,  $t = 1, 2, 3, \dots$  is a white noise sequence, and:

- $g_t$  represents a trend component
- $s_t$  represents a seasonal component
- $h_t$  captures holiday/calendar effects

- This can be seen as a particular type of smoother, based on a particular model for the trend (which we will describe shortly; the seasonal and holiday components are fairly generic). We could also refer to it as a particular type of additive model, where the regressor is time
- The seasonal component is parametrized by a Fourier (cosine and sine) basis at given fixed, known periods chosen by the user. For frequencies  $\omega_j$ ,  $j = 1, \dots, p$  (equivalently, periods  $1/\omega_j$ ,  $j = 1, \dots, p$ ), recall, this is:

$$g_t = \sum_{j=1}^p \left( a_j \cos(2\pi\omega_j t) + b_j \sin(2\pi\omega_j t) \right)$$

for coefficients  $a_j, b_j$ ,  $j = 1, \dots, p$

- The holiday/calendar component is simply parametrized using indicator variables

$$h_t = \sum_{j=1}^m \alpha_j \cdot 1\{t \in D_j\}$$

where  $\alpha_j$ ,  $j = 1, \dots, m$  are coefficients and each  $D_j$  is a set of dates representing a particular holiday or calendar event (e.g., Christmas, Thanksgiving, etc.)

- Finally, the trend component is modeled in one of two ways. The first way is for *saturating* trends. For this, [Taylor and Letham \(2018\)](#) propose to model  $g_t$  using a sigmoid function with a piecewise growth rate:

$$g_t = \frac{C(t)}{1 + \exp \left( c_0 + c_1 t + \alpha \sum_{j=1}^r \beta_j \cdot (t - t_j)_+ \right)}$$

where  $u_+ = \max\{u, 0\}$  denotes the positive-part of  $u$ . Here  $C(t)$  is a (possibly) time-varying capacity, which it appears [Taylor and Letham \(2018\)](#) recommend be set externally (e.g., based on market size considerations)

- For *non-saturating* trends, [Taylor and Letham \(2018\)](#) propose to model  $g_t$  using a piecewise linear trend directly:

$$g_t = c_0 + c_1 t + \alpha \sum_{j=1}^r \beta_j \cdot (t - t_j)_+$$

- In either case (saturating or non-saturating),  $\beta_j$ ,  $j = 1, \dots, r$  are coefficients to be estimated. Also,  $t_j$ ,  $j = 1, \dots, r$  are knots (locations where the slope changes), which, in the simplest case, could be fixed ahead of time. Instead, [Taylor and Letham \(2018\)](#) recommend that knots be selected using  $\ell_1$  penalization from a large initial set of locations. That is, they use the  $\ell_1$  penalty

$$\sum_{j=1}^r |\beta_j|$$

when fitting the model, which is like a special type of lasso regression. (In fact, though it may not be obvious at first pass, placing an  $\ell_1$  penalty on  $\beta_j$ ,  $j = 1, \dots, r$  here is actually equivalent to reparametrizing the entire sequence as  $g_t = \theta_t$ , and then using an  $\ell_1$  penalty on second differences of  $\theta_t$ ; recall, this is the penalization scheme used in *trend filtering*, which you learned earlier in the course when we covered smoothing)

- Altogether, the Prophet model is fit by minimizing the sum of squared errors to the observed data, over all parameters  $a_j, b_j, \alpha_j, c_0, c_1, \beta_j$  that determine the decomposition, with a squared  $\ell_2$  (ridge) penalty on the parameters  $a_j, b_j, \alpha_j$  for the seasonal component  $s_t$  and holiday component  $h_t$ , and an  $\ell_1$  (lasso) penalty on the parameters  $\beta_j$  for the trend component  $g_t$
- Forecasts are generated by extrapolating the fitted components forward in time. For  $s_t$  and  $h_t$ , this is straightforward, because they are periodic in nature. For  $g_t$ , this is done by holding the slope (i.e., growth rate in the saturating model) constant from its last value, as we move forward in time

- (Though unimportant for our purposes here, [Taylor and Letham \(2018\)](#) actually phrase all of this in the context of a hierarchical Bayesian model, with normal and Laplace priors that serve the purpose of regularization; this Bayesian machinery also provides added stochasticity in computation of prediction intervals)
- So, how does the Prophet model compare to ARIMA and ETS? It depends on who you ask. In their original paper, [Taylor and Letham \(2018\)](#) find that ARIMA and ETS models are too rigid in their motivating examples—take a look at Figure 3 in their paper, and compare Figure 2, which displays Prophet forecasts on the same data

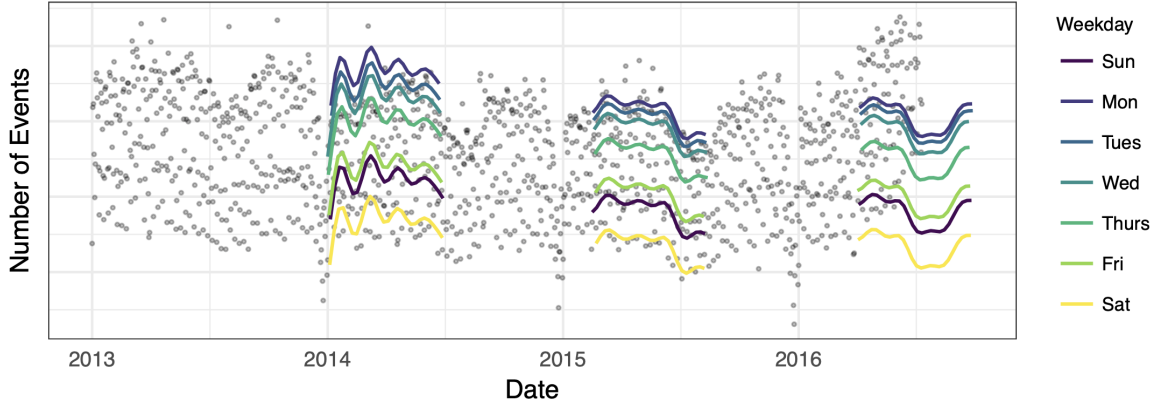


Figure 2: Prophet forecasts, from [Taylor and Letham \(2018\)](#).

- Indeed, in their traditional forms, ARIMA and ETS models lack the flexibility of the Prophet model, particularly the flexibility exhibited in the latter’s trend component. However, both ARIMA and ETS can be extended to accommodate more sophisticated trends. With ARIMA, you have actually already seen a way to do this: we can phrase the problem as *regression with correlated errors* (where we use an ARIMA model for the errors), and can then use time for the regressor and invoke a flexible basis representation to model trends just like Prophet does
- Hyndman and Athanasopoulos (HA) call this a *dynamic regression model* and study it in Chapter 10 of their book. The advantage this has over Prophet is that it is able to capture auto-correlations in the errors, which can lead to narrower prediction intervals. The advantage Prophet has is speed: it is usually more efficient to fit the Prophet model, since its error model (white noise) is simpler and this makes the optimization a version of penalized least squares

### 1.3 Neural network autoregression

- A *neural network* describes a class of models that make real-valued predictions  $f(x) \in \mathbb{R}$  from an input feature vector  $x \in \mathbb{R}^p$  of the form:

$$\begin{aligned}
 f_1(x) &= \rho(W_1 x + b_1) \\
 f_\ell(x) &= \rho(W_{\ell-1} f_{\ell-1}(x) + b_{\ell-1}), \quad \ell = 2, \dots, L \\
 f(x) &= f_L(x)
 \end{aligned}$$

- Here each  $W_\ell \in \mathbb{R}^{d_\ell \times d_{\ell-1}}$  is a matrix of weights that maps from the dimension  $d_{\ell-1}$  of layer  $\ell - 1$  to the dimension  $d_\ell$  of layer  $\ell$ . Note that  $d_0 = p$ , and  $d_L = 1$  (for real-valued predictions)
- Each  $b_\ell \in \mathbb{R}^{d_\ell}$  is a vector of intercepts (often called *biases* in the deep learning community). Generically, the parameters  $W_\ell, b_\ell$  are all learned by minimizing the sum of squared errors of predictions on the training data

- The function  $\rho$  is a nonlinear *activation function* that is interpreted as being applied componentwise. It is user-chosen; common choices are  $\rho(u) = u_+$  and  $\rho(u) = 1/(1 + e^{-u})$
- Lastly,  $L$  here is the number of layers, also a design choice, and often called the *depth* of the network
- For time series, one of the simplest things that can be done with neural networks is just to form input features by taking lags of the given response variable. We can use both nonseasonal and seasonal lags (as in ARIMA). This gives rise to what we call a *neural network autoregressive* (NNAR) model
- What we described above is actually just a particular type of neural network architecture, and indeed, the simplest kind, called a *feedforward* neural network. Many other architectures are possible, and some more appropriate for time series data, such as the *long short term memory* (LSTM) network, a type of recurrent neural network
- A popular time series forecaster based on LSTMs is called *DeepAR*, proposed by [Salinas et al. \(2020\)](#), from Amazon. Relative to all other methods you have learned thus far, DeepAR is quite complicated to describe precisely (and to train). Figure 3 gives a schematic: the left column for training, and the right for forecasting

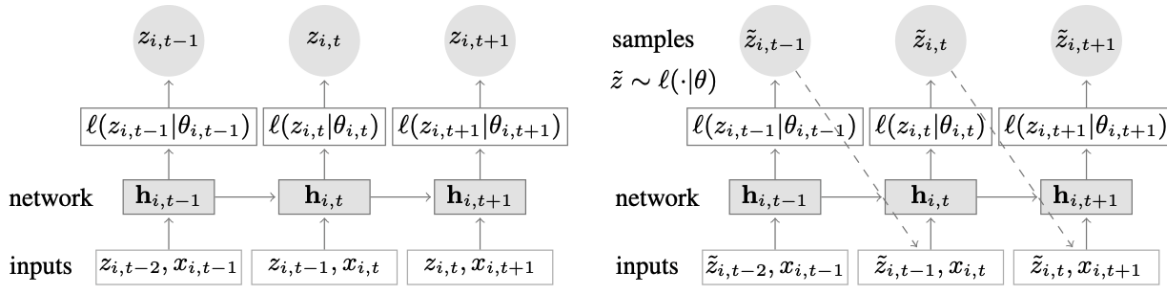


Figure 3: Schematic of DeepAR network, from [Salinas et al. \(2020\)](#), where each  $h_{i,t}$  is the output of a recurrent neural network with LSTM cells.

- However, like many deep learning methods, it can work very well in data-rich prediction problems that have a high signal-to-noise ratio. Another strength it is often cited as having is that it can learn rich, joint correlation structures between *multiple* time series—think of spatiotemporal data, where we have one time series per location, and many locations—and it can and produce forecasts which respect this correlation structure
- Perhaps not surprisingly, some authors are now re-purposing transformers in order to turn them into time series forecasters. As deep learning continues to move forward, we will continue to see spillover into time series forecasting

## 2 Ensembling

- So, you have lagged regression, ARIMA, ETS, Theta, Prophet, and DeepAR in your forecast toolkit. You have data in front of you, and you're unsure which model(s) will work best. What do you do?
- The answer we have given all semester: *use time series CV*. That is, move the time index back to  $t = t_0$ , train each model on the past, make forecasts, compare to the unseen target values, march the time index forward sequentially, and repeat. In doing so, we can compute our chosen accuracy metric (MAE, MASE, etc.) for each model, and choose the one that emerges as the most accurate, perhaps using Occam's razor to help decide between models that perform similarly
- Here is an alternative that can often be advantageous: *combine* the models, which can be informed by their out-of-sample predictions from time series CV, rather than using CV to select one of them

- The combination of prediction models is often called *ensembling*, or *aggregation*. It has a long history in statistics and machine learning, including in forecasting. Still, it seems perhaps people don’t quite appreciate enough just how effective it can be
- Roughly speaking, there are two types of ensembles: *untrained* and *trained* ones. An untrained ensemble simply combines a given set of base models (also called constituent models) without consideration of their individual past performance. Meanwhile, a trained ensemble combines the base models in a way that reflects their performance, typically upweighting more accurate base models
- At a high level, an ideal ensemble model would satisfy the following two properties:
  1. *compete-with-best*: the ensemble should have accuracy that is competitive with the best individual constituent model
  2. *robustness-over-all*: the ensemble should have greater robustness (make fewer large errors) than any individual constituent model
- Interestingly (though perhaps not surprisingly), these two are somewhat at odds. Untrained ensembles can be *very* robust, but typically cannot compete with the best constituent model. On the other hand, trained ensembles can compete with the best constituent model (and even outperform it, if the ensemble is flexible enough), but “heavier” training schemes tend to hurt robustness

## 2.1 Untrained methods

- The simplest untrained ensemble method is just to take a straight average of all constituent model forecasts. Denote these by  $\hat{x}_{t+h|t}^j$ , for  $j = 1, \dots, p$ . Then the simple average ensemble forecast is

$$\hat{x}_{t+h|t}^{\text{avg}} = \frac{1}{p} \sum_{j=1}^p \hat{x}_{t+h|t}^j$$

- Another option is to take a median of point forecasts from the constituent models,

$$\hat{x}_{t+h|t}^{\text{med}} = \text{median} \left\{ \hat{x}_{t+h|t}^j : j = 1, \dots, p \right\}$$

- Despite their simplicity, these methods can be very effective, especially when we have a diverse set of constituent models (which have a healthy degree of diversity in their errors), as this can lead to very robust average or median ensembles
- Figure 4 shows a striking example of this, from 1- though 4-week ahead Covid-19 death forecasting. The COVIDhub-ensemble model was (for most of the pandemic) a simple median ensemble of all forecasts submitted to the US Covid-19 Forecast Hub which met a certain very basic set of inclusion criteria (ruling out truly wild or unrealistic forecasts). This ensemble model was the basis for all official CDC communications on short-term Covid-19 forecasting. The number of constituent models fluctuated up and down over time, in between about 25 to 50
- For state-level forecasts made over a 1.5 year period during the pandemic, the figure considers a notion called *standardized rank* that is defined as follows. For a given forecast task (given state, given target date, and given horizon), we rank all available forecasts using an accuracy measure called WIS (a generalization of absolute error for quantile-based forecasts) figure. We assign a standardized rank of 1 for the most accurate forecast of all available forecasts for that task, and 0 to the least accurate. Then for each forecaster, we average these standardized ranks over all forecast tasks, and plot a histogram of their standardized rank distribution
- As we can see in Figure 4, the distribution of the ensemble is *very* different from all others—in particular, look at its thin left tail. It is the only forecast model that was in the top half of forecasters (standardized rank  $\geq 0.5$ ) for 75% of the forecast tasks. Interestingly, we can also see that it is not in the top quartile of forecasters (standardized rank  $\geq 0.75$ ) as often as the best constituent models, hinting at the tradeoff described above

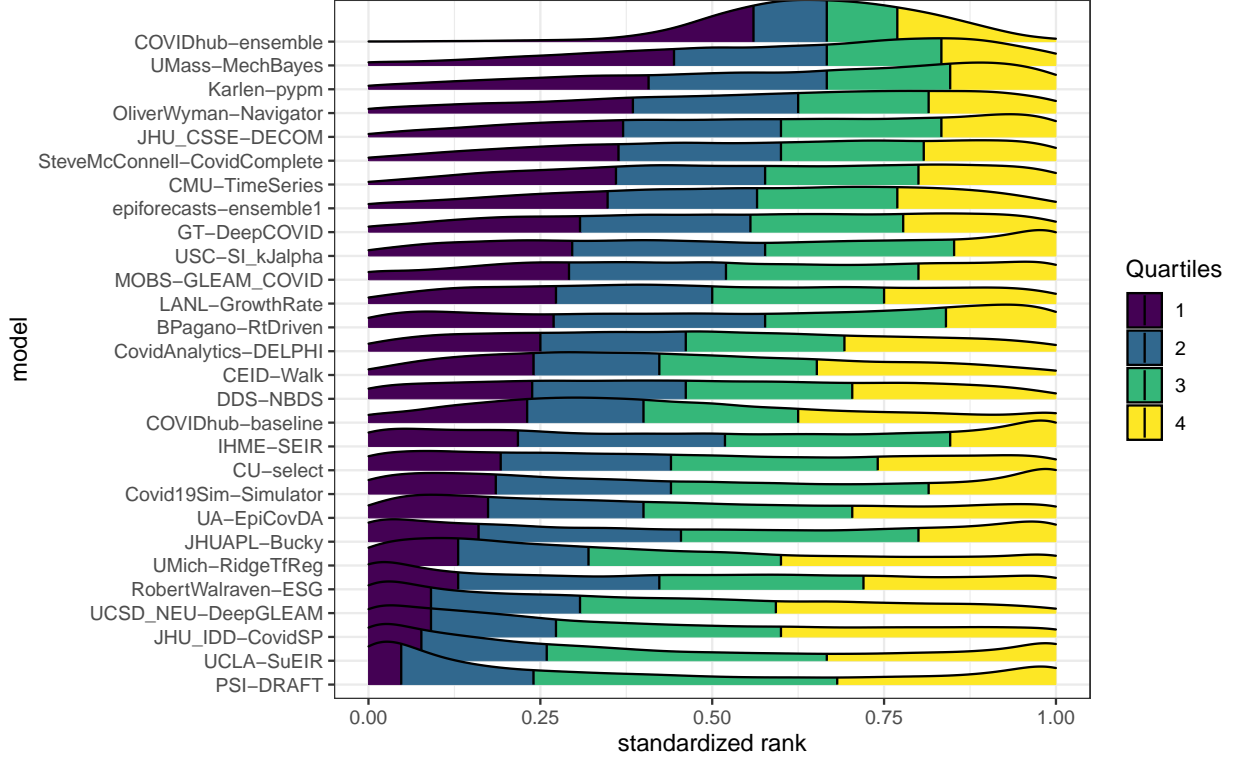


Figure 4: Comparison of Covid-19 death forecast models, from [Cramer et al. \(2022\)](#).

## 2.2 Trained methods

- One of the simplest trained ensemble methods is to directly fit a weighted combination of constituent forecasts in order to optimize accuracy (MSE, MAE, etc.). For example, going by MSE, and fixing  $h = 1$  for simplicity, to form the ensemble at time  $t$  (for a forecast of the target value at time  $t + 1$ ), we would first solve:

$$\begin{aligned} \min_{w \in \mathbb{R}^p} \quad & \sum_{s=t_0+1}^t \left( x_s - \sum_{j=1}^p w_j \cdot \hat{x}_{s|s-1}^j \right)^2 \\ \text{subject to} \quad & \sum_{j=1}^p w_j = 1, \text{ and } w_j \geq 0, j = 1, \dots, p \end{aligned}$$

Note that this is simply a regression of the target values (response) on the forecasts made by constituent models (features), where we constrain the coefficients to be nonnegative and sum to 1

- After obtaining the solution  $\hat{w}_j^t, j = 1, \dots, p$ , we then form the weighted ensemble to make forecasts

$$\hat{x}_{t+1|t}^{\text{stack}} = \sum_{j=1}^p \hat{w}_j^t \cdot \hat{x}_{t+1|t}^j$$

- This is sometimes called *linear stacking*: here “stacking” refers to the fact that we have stacked the predictions made by our constituent models into features for a second-level model, and “linear” refers to the fact that our second-level model is linear in the features (it is a linear regression subject to constraints)
- Instead of re-solving the optimization problem above at each forecast time  $t$ , we could also incrementally update the learned weights over time. A method called (online) *mirror descent* applied to the

above optimization problem yields the following intuitive weight updates, at time  $t$ :

$$\tilde{w}_j^t = \hat{w}_j^{t-1} \exp \left\{ \eta \cdot \hat{x}_{t|t-1}^j \left( x_t - \sum_{j=1}^p \hat{w}_j^{t-1} \cdot \hat{x}_{t|t-1}^j \right) \right\}, \quad j = 1, \dots, p$$

$$\hat{w}_j^t = \tilde{w}_j^t / \sum_{\ell=1}^p \tilde{w}_\ell^t, \quad j = 1, \dots, p$$

Here  $\eta > 0$  is a parameter called the learning rate. In words, if a forecaster’s latest prediction  $\hat{x}_{t|t-1}^j$  is correlated with the ensemble’s latest residual  $x_t - \sum_{j=1}^p \hat{w}_j^{t-1} \cdot \hat{x}_{t|t-1}^j$ , then it is underrepresented in the ensemble, and we increase its weight. This is related to a general class of online learning algorithms called *exponential weights* algorithms; and there is a huge literature on related schemes

- Stacking—especially more flexible second-level models that are nonlinear, or use weights that depend on features—can lead to impressive accuracy in practice. Figure 5 shows a nice example of this (albeit in a batch prediction context, rather than time series). The figure shows a neural network ensemble model called *deep quantile aggregation* (DQA), that is trained as a second-layer model on top of many constituent models, including some very flexible ones (such as deep neural networks). Across 35 data sets, ordered by their signal-to-noise ratio along the x-axis, the WIS (recall this is a generalization of absolute error for quantile-based predictions) of all of the constituent models and other ensemble methods is compared to that of DQA. We can see that no method outperforms DQA (nothing below the red horizontal line), and for problems with more signal, towards the right end of the x-axis, DQA improves massively on a lot of the other models

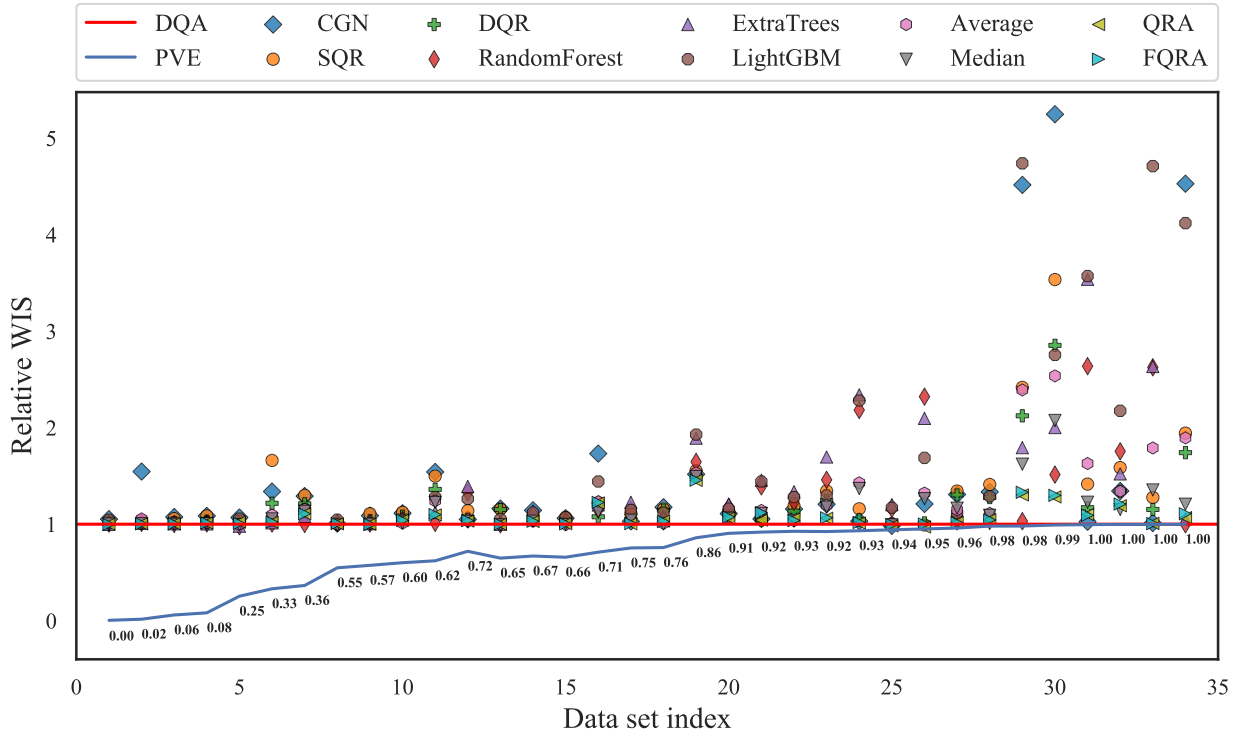


Figure 5: Deep stacking of quantile prediction models, from [Fakoor et al. \(2023\)](#).

### 3 Calibration

- Beyond point forecasts, all of the methods that we have learned thus far produce *prediction intervals*, which reflect uncertainty in the predictions that are made, useful for downstream decision-making



- How can we measure the “fidelity” of such prediction intervals? Fixing a horizon  $h$ , denote by  $I_{t+h|t}^{1-\alpha}$  denote the level  $1 - \alpha$  prediction interval made by our forecaster at time  $t$ , for the target variable  $x_{t+h}$  at time  $t + h$
- Recall, for (say) a 90% prediction interval, this is often (though not necessarily) of the form:

$$I_{t+h|t}^{1-\alpha} = [\hat{x}_{t+h|t} - \hat{\sigma}_h q_{0.95}, \hat{x}_{t+h|t} + \hat{\sigma}_h q_{0.95}]$$

where  $\hat{x}_{t+h|t}$  is a point forecast of  $x_{t+h}$  made at time  $t$ ,  $\hat{\sigma}_h^2$  is an estimate of the variance of the  $h$ -step ahead forecast distribution, and  $q_{0.95}$  is the 0.95 quantile of the standard normal distribution

- Given such intervals, we can measure their empirical *coverage*, in a time series CV sense:

$$\text{Coverage} = \frac{1}{n - t_0} \sum_{t=t_0+1}^n 1\{x_t \in I_{t|t-h}^{1-\alpha}\}$$

In words, this is the fraction of times that our prediction intervals cover their targets. Of course, we want the coverage of our prediction intervals to roughly match the nominal level,  $\text{Coverage} \approx 1 - \alpha$

- Sadly, traditional methods for producing prediction intervals will not always yield empirical coverage that hovers around the nominal level. This is because the methods for producing these intervals rely on assumptions that are often violated in practice
- See Figure 6 for an example of the empirical coverage of the COVIDhub-ensemble model for forecasting cases, hospitalizations, and deaths. These are very difficult forecasting problems (especially cases and hospitalizations)

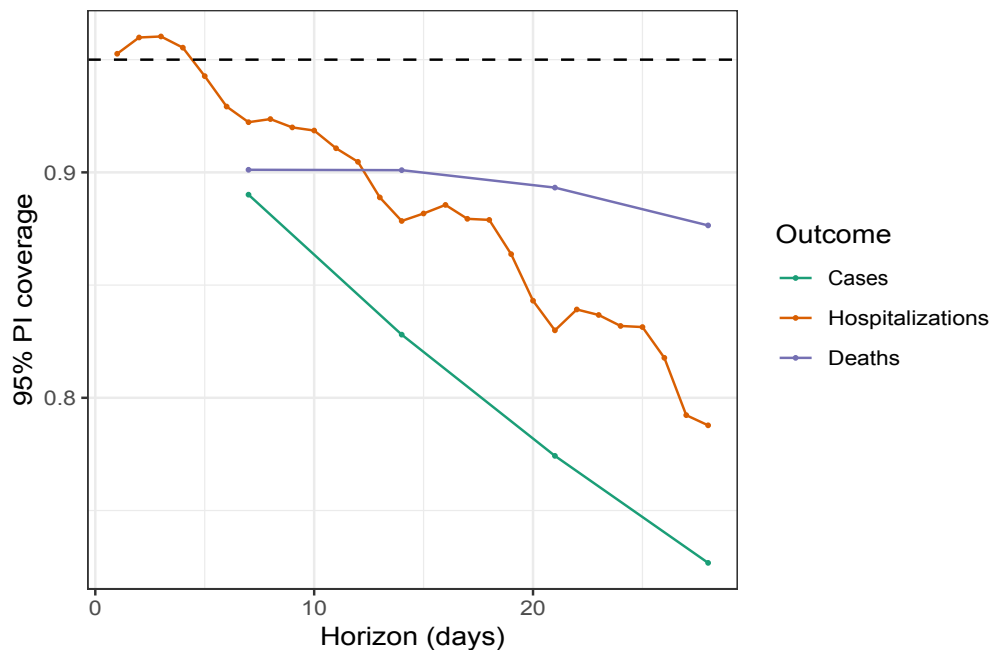


Figure 6: Empirical coverage of Covid-19 ensemble forecasts, from <https://delphi.cmu.edu/blog/2021/09/30/on-the-predictability-of-covid-19/>.

- In what remains of this lecture, we’ll learn about a simple and generic method for adjusting prediction intervals online, so that their empirical coverage tracks the nominal level more faithfully. Then we’ll learn about how to generalize the concept of coverage (which applies to a prediction interval) so that we can measure the fidelity of an entire predicted distribution. This brings us to a topic called *calibration*

### 3.1 Quantile tracking

- We briefly describe a method due to [Angelopoulos et al. \(2023\)](#), called *quantile tracking* (also called conformal P-control, in that paper). Fix  $h = 1$  for simplicity, and denote by  $\hat{q}_t^{1-\alpha}$  a predicted level  $1 - \alpha$  quantile of the distribution of  $e_t$ , the absolute forecast error at time  $t$ :

$$e_t = |x_t - \hat{x}_{t|t-1}|$$

- Note that a level  $1 - \alpha$  prediction interval for  $x_t$  could then be formed via:

$$I_{t|t-1}^{1-\alpha} = [\hat{x}_{t|t-1} - \hat{q}_t^{1-\alpha}, \hat{x}_{t|t-1} + \hat{q}_t^{1-\alpha}]$$

- The motivation is as follows:  $e_t \leq \hat{q}_t^{1-\alpha} \iff x_t \in I_{t|t-1}^{1-\alpha}$ , so if  $\hat{q}_t^{1-\alpha}$  were a proper level  $1 - \alpha$  quantile in the long-run frequency sense:

$$\frac{1}{T} \sum_{t=t_0+1}^{t_0+T} 1\{e_t \leq \hat{q}_t^{1-\alpha}\} \approx 1 - \alpha$$

for large  $T$ , then we would indeed have empirical coverage:

$$\frac{1}{T} \sum_{t=t_0+1}^{t_0+T} 1\{x_t \in I_{t|t-1}^{1-\alpha}\} \approx 1 - \alpha$$

for large  $T$

- [Angelopoulos et al. \(2023\)](#) propose the following online quantile update:

$$\hat{q}_{t+1}^{1-\alpha} = \begin{cases} \hat{q}_t^{1-\alpha} + \eta(1 - \alpha) & \text{if } x_t \notin I_{t|t-1}^{1-\alpha} \\ \hat{q}_t^{1-\alpha} - \eta\alpha & \text{if } x_t \in I_{t|t-1}^{1-\alpha} \end{cases}$$

where  $\eta > 0$  is a parameter called the learning rate. In words, if the latest interval does not cover, then we *increase* the quantile by  $\eta(1 - \alpha)$  (make the next interval *wider*), whereas if the latest interval covered, then we *decrease* the quantile by  $\eta\alpha$  (make the next interval *narrower*). This is very intuitive, and it turns out to be equivalent to online gradient descent run on a particular loss function (which sums the quantile loss between  $e_t$  and  $\hat{q}_t^{1-\alpha}$  over all time  $t$ ). See also [Gibbs and Candès \(2021\)](#) for an earlier, related idea

- [Angelopoulos et al. \(2023\)](#) prove that if the forecast errors are bounded,  $e_t \leq b$  for all  $t$  and for some  $b < \infty$  (note that we do not need to know the bound  $b$  in order to run the algorithm), then quantile tracking results in long-run coverage:

$$\frac{1}{T} \sum_{t=t_0+1}^{t_0+T} 1\{x_t \in I_{t|t-1}^{1-\alpha}\} \rightarrow 1 - \alpha, \quad \text{as } T \rightarrow \infty$$

for any  $t_0$  and  $\eta$ . Importantly, this holds *with no assumptions* on the distribution of the data or the forecast errors (or the forecaster itself)

- You can read the paper for many interesting examples, extensions to unbounded scores, and prospective corrections to forecast errors (driven by a connection to control theory)

### 3.2 PIT calibration

- Last but not least, we arrive at calibration. Unfortunately, this is a term that is often used ambiguously, to mean different things. Here we use it to mean a precise way of measuring the fidelity of an entire *predicted distribution*, rather than just a prediction interval

- That is, suppose that our forecaster produces an entire distribution  $F_{t|t-h}$  for the target  $x_t$  at time  $t$ . Generally, we think of  $F_{t|t-h}$  as a cumulative distribution function (CDF); note that, from  $F_{t|t-h}$ , we could obviously produce prediction intervals for  $x_t$  (at any levels that we desire)
- For notational simplicity, we will hide the forecast time  $t - h$ , and simply denote  $F_{t|t-h}$  by  $F_t$  in what follows
- Now, there are numerous different definitions of what it means for a probabilistic forecaster (which produces  $F_t$ ) to be calibrated, and these definitions are not generally equivalent. (This contributes to the frequent ambiguity in the use of the term “calibration”.) See, e.g., [Gneiting and Resin \(2022\)](#) for a recent nice overview of different definitions and their relationships
- We will stick with one particular definition. The forecaster that produces the sequence of predicted CDFs  $F_t$  is said to be *PIT calibrated* provided:

the empirical distribution of  $F_t(x_t)$ ,  $t = 1, \dots, T$  converges  
to the standard uniform distribution,  $\text{Unif}(0, 1)$ , as  $T \rightarrow \infty$

- Equivalently, this means:

$$\frac{1}{T} \sum_{t=1}^T 1\{F_t(x_t) \leq \tau\} \rightarrow \tau, \quad \text{as } T \rightarrow \infty, \text{ for all } \tau \in [0, 1]$$

- Note that this is even stronger than coverage of prediction intervals: assuming, as we will do throughout for simplicity, that each  $F_t$  is continuous and hence invertible, we have:

$$\frac{1}{T} \sum_{t=1}^T 1\{x_t \leq F_t^{-1}(\tau)\} \rightarrow \tau, \quad \text{as } T \rightarrow \infty, \text{ for all } \tau \in [0, 1]$$

In other words, each  $F_t^{-1}(\tau)$  is a proper level  $\tau$  quantile in the long-run frequency sense, and we can use  $[F_t^{-1}(\alpha/2), F_t^{-1}(1 - \alpha/2)]$  to form a valid  $1 - \alpha$  prediction interval

- To get intuition as to where the definition of PIT calibration comes from (including its name), recall that for a random variable  $X$ , with CDF  $F$  (assumed continuous and hence invertible), its *probability integral transform* (PIT) is  $F(X)$ . A key property of the PIT:  $F(X) \sim \text{Unif}(0, 1)$ . The proof of this fact is elementary: for any  $\tau$ ,

$$\mathbb{P}(F(X) \leq \tau) = \mathbb{P}(X \leq F^{-1}(\tau)) = F(F^{-1}(\tau)) = \tau$$

Hence the random variable  $F(X)$  has the CDF of a standard uniform, and must itself be uniformly distributed

- We can now think of PIT calibration as seeking the uniform property of the PIT transform, when we apply this idea along the sequence  $t = 1, 2, 3, \dots$ . Somewhat more precisely, if  $F$  denotes a *random* CDF, drawn uniformly at random from the collection of CDFs  $F_t$ ,  $t = 1, 2, 3, \dots$ , and  $X$  is a random target variable, again drawn uniformly at random from the collection of target values  $x_t$ ,  $t = 1, 2, 3, \dots$ , then PIT calibration requires that  $F(X) \sim \text{Unif}(0, 1)$
- The observed PIT values  $F_t(x_t)$ ,  $t = 1, 2, 3, \dots$  can be used as a diagnostic tool. These values always lie in  $[0, 1]$ , and we can form an empirical density (or a histogram) of these PIT values as we collect them. If the forecaster is PIT calibrated, then these will look uniform, i.e., the PIT density will be flat. If the forecaster is *underconfident*, then the PIT density will tend to look U-shaped. Meanwhile, if the forecaster is *overconfident*, then the PIT density will tend to have an upside-down U-shape. This behavior can be confirmed with simple examples with normal densities, see Figure 7. An example of real PIT densities from short-term Flu forecasters is given in Figure 8
- Being more general than coverage, it is also harder to achieve PIT calibration in practice (certainly in a distribution-lean sense, without many assumptions). However, this is an active area of research, and we will likely see new advances in the next few years

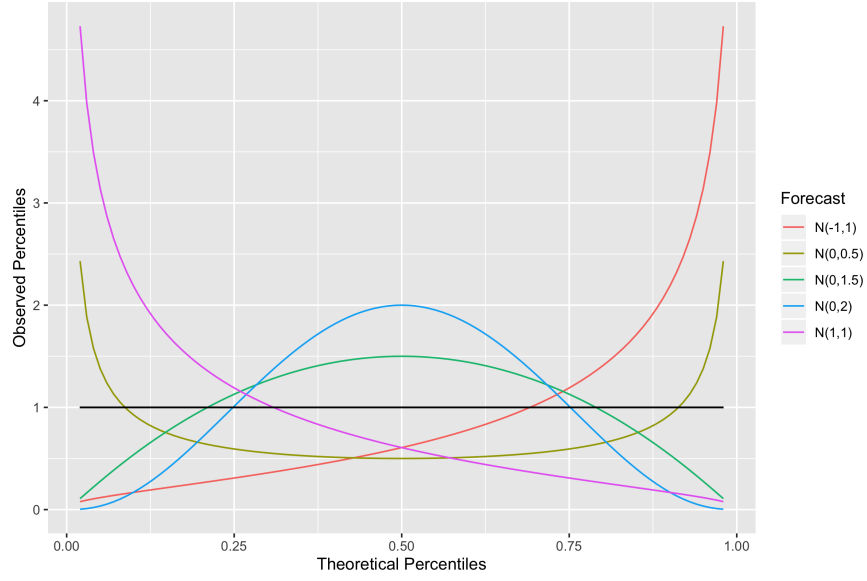


Figure 7: Densities of PIT distributions for several simple normal forecasters, when the true target distribution is  $N(0,1)$ , from [Rumack et al. \(2022\)](#).

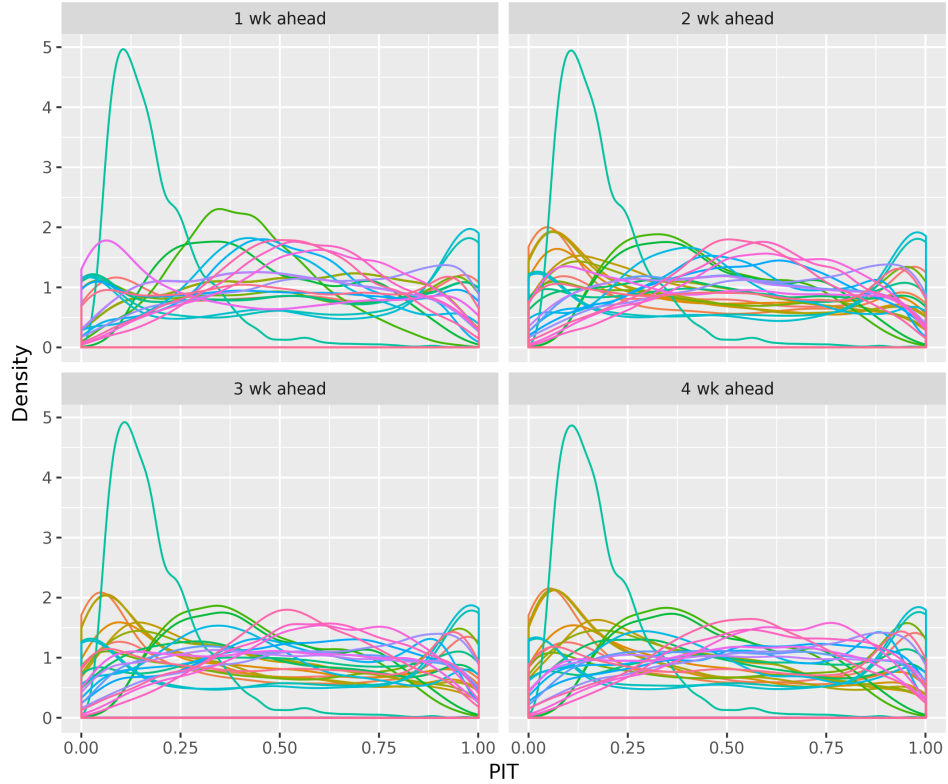


Figure 8: Densities of PIT distributions from 27 forecasters submitted to the annual FluSight challenges, from [Rumack et al. \(2022\)](#). These are seasonal influenza forecasting challenges held by CDC, spanning 9 seasons (2010-11 to 2018-19). The PIT densities fall mostly into one of two categories: overdispersed with a peak around 0.5, and underdispersed with peaks at around 0 and 1. (The outlier with a peak at 0.1 is the PIT density of a simple baseline forecaster.).

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