

Nowcasting with signature methods

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Abstract

Key economic variables are often published with a significant delay of over a month. The nowcasting literature has arisen to provide fast, reliable estimates of delayed economic indicators and is closely related to filtering methods in signal processing. The path signature is a mathematical object which captures geometric properties of sequential data; it naturally handles missing data from mixed frequency and/or irregular sampling – issues often encountered when merging multiple data sources – by embedding the observed data in continuous time. Calculating path signatures and using them as features in models has achieved state-of-the-art results in fields such as finance, medicine, and cyber security. We look at the nowcasting problem by applying regression on signatures, a simple linear model on these nonlinear objects that we show subsumes the popular Kalman filter. We quantify the performance via a simulation exercise, and through application to nowcasting US GDP growth, where we see a lower error than a dynamic factor model based on the New York Fed staff nowcasting model. Finally we demonstrate the flexibility of this method by applying regression on signatures to nowcast weekly fuel prices using daily data. Regression on signatures is an easy-to-apply approach that allows great flexibility for data with complex sampling patterns.

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1 Introduction

Households, businesses, and policymakers need up-to-date economic information to make decisions. However, information is often incomplete or delayed, as

collection and compilation of the data underlying key economic indicators takes time. For example, UK monthly GDP is first published with an approximately 6-week lag. At the same time, there is a wealth of information about various aspects of the economy that can be gleaned from related indicators that are released with less publication lag or on a more frequent basis, for example the Office for National Statistics publishes a wide range of weekly indicators of economic activity. As a result, there has been an increasing demand for modelled early estimates of the economy utilising these more frequent observations. These early estimates are called nowcasts.

The economic literature on nowcasting has grown dramatically in the past 20 years (e.g. Stock and Watson, 2002; Giannone et al., 2006; Bok et al., 2018), and has focused on three main issues. First, when incorporating a large number of predictors into a model, how one imposes structure to reduce dimensionality. Second, how one incorporates data into a model when there are missing observations, which may be caused by mixed or irregular sampling frequencies (Kapetanios et al., 2018; Ghysels and Marcellino, 2018). Third, how one allows for time-varying parameters or for nonlinearities. We discuss these in more detail in the following subsections.

In this paper, we introduce a new nowcasting method, regression on signatures (*“the signature method”*), that addresses the issue of missing observations and works in a general setting so that no additional effort is required to infer nonlinear relationships and standard dimension reduction tools can be applied. We provide motivation and implementation details for the signature method in Sections 3 and 4, before applying this method on both simulated and real data (Sections 5–7). All experimental results can be reproduced with our [Github repository](#). In addition, we have created a Python package [SigNow](#), so that the signature method can be applied to other nowcasting applications,

1.1 Summary of contributions

Our primary contribution is a new nowcasting method, *regression on signatures*, that addresses some of the challenges associated with nowcasting. The path signature is a mathematical object to describe time-series-like data, arising from the theory of rough paths, see Friz and Hairer (2020). We shall refer to methods that utilise the path signature for prediction tasks as *signature methods*. Signature methods naturally allow for missing data, and mixed frequency and irregular sampling, by modelling time series processes in continuous time. They have been used to great success in a range of applications including Chinese handwriting recognition (Graham, 2013), sepsis detection (Morrill et al., 2020), and malware detection (Cochrane et al., 2021).

Advantages of signature methods

The use of signature methods gives a variety of practical benefits over other modelling approaches. We give a brief overview of some of these advantages here.

- *Simplicity of modelling.* A key advantage of working with signature methods is the simplicity of modelling. When compared with existing methodologies, the signature approach reduces almost entirely to two steps: (i)

the computation of signatures from observed paths (continuous time series in the context of nowcasting), and (ii) standard linear regression. The former can be done through existing python libraries such as `esig` or `iisignature` (Reizenstein and Graham, 2020). The latter step does not require significant development effort, and it is easy to include additional predictors. Essentially, using signatures as features eliminates some of the traditional complexity of time series modelling.

- *Ease of workflow.* Using signatures reduces time series modelling to linear regression, and can effortlessly be combined with existing techniques for dimension reduction, variable selection, and various non-standard regression models such as generalized linear models (with signature features). The difficulty in using signatures is simply in building and understanding effective models. We have created `SigNow` a packaged pipeline that is publicly available on Github, which should be easy-to-implement for other nowcasting problems.
- *Interpretability.* There has recently been increased attention given to the use of various machine-learning techniques for estimation. However, these typically suffer from a lack of interpretability – it is difficult to explain what the estimated model is doing, and hence to monitor and assess its risks and performance. As signature models are based on linear regression, which is easily interpretable, the only difficulty is in interpreting the signature terms themselves. While this is not completely straightforward, they have geometric interpretation and work has been done to illustrate this (Yang et al., 2017).
- *Flexibility and robustness.* Signature methods naturally allow for missing data due to mixed frequency and irregular sampling by modelling time series processes in continuous time. Signatures capture the interactions between dimensions in a multi-dimensional path, and the ordering of events is preserved (Gyurkó et al., 2013; Levin et al., 2013).

Essentially, our method is as follows: take the observations available, and compute their signature (treating them as discrete observations of continuous paths). Then, use these signatures as regressors for the target variable of interest. This simple method coupled with the properties of the signature, is sufficient for a wide variety of nowcasting tasks. As in state-space models, signatures can be constructed and used in non-stationary settings and in settings where the underlying dynamics are nonlinear (see Section 2.3).

Signatures on their own do not solve the problem of dimension reduction (see Section 2.1). However, signatures can be combined with dimension reduction methods. As a second contribution, we show how to combine the flexible dynamics of signatures with standard dimension reduction techniques like principal component analysis.

1.2 Outline of paper

The rest of the paper is organised as follows: Section 2 gives an overview of a selection of challenges that nowcasting methods may address; Section 3 details

nowcasting from a mathematical perspective using signatures, this include background on discrete and continuous time state-space models, Kalman filters, and proves that the Kalman filter can be equivalently written as a linear regression on the signature space; Section 4 summarizes how to use regression on signatures in nowcasting, in terms of the pipeline and the practical implementation; Section 5 demonstrates that it is possible to replicate the performance of the Kalman filter by using regression on signature in a simulated problem; Section 6 applies the signature model to nowcast US GDP growth drawing comparisons with a dynamic factor model based on the New York Fed staff nowcast model; Section 7 applies the signature method to fuel data; finally, Section 8 concludes.

2 Challenges of traditional nowcasting

Nowcasting in economics has some commonly occurring challenges that we discuss in this section.

2.1 Dimension reduction

In most nowcasting exercises, a large number of variables are used to predict a single target. For example, the Federal Reserve Bank of New York staff nowcasting model uses 37 variables to nowcast US GDP (FRBNY, 2016). The literature has tackled dimensionality issues in a variety of ways. The leading approaches in the macroeconomic nowcasting literature include dynamic factor models (DFM), Bayesian vector auto-regressions (BVAR), and penalised estimation methods like the LASSO.¹

Dynamic factor models were introduced into economics by Geweke (1977) and are reviewed in Stock and Watson (2017) and Bai and Ng (2008). These models are based on the idea that most of the time series variation in a large set of economic variables is driven by the dynamics of a small number of unobserved common factors. These factors can be estimated from the complete set of economic variables using singular value decomposition of the data matrix. The dynamics of the factors are analysed using standard time series methods such as vector autoregression. There has been rapid development of DFM methods in applied macroeconomic analysis, and they have been used extensively in economic forecasting and nowcasting, see e.g., Giannone et al. (2006), Doz et al. (2006), and Bok et al. (2018).

Bayesian vector auto-regressions, developed in macroeconomics in Litterman (1979) and Sims (1980) (see Karlsson (2013) for a review), impose restrictions on and regularise high-dimensional models by specifying prior beliefs on the parameter space and then using Bayesian methods to estimate the posterior distribution of the parameters. The posterior distribution is used to nowcast the economic variables of interest. These methods are generally restricted to linear models, and so depend on careful choice of parameterization to ensure linear models are appropriate.

¹Recent macroeconomic papers have also investigated the use of other machine learning methods including random forests and artificial neural networks (Richardson et al., 2021). Furthermore, outside of economics, there is a large literature on dynamic estimation across a range of disciplines including statistics, meteorology, computer science, engineering, etc.

Penalised estimation methods impose restrictions on and regularise high-dimensional models by including a penalty term in the estimation objective function. For example, the LASSO estimator, popularised in statistics by Tibshirani (1996), adds a penalty based on the sum of the absolute values of the parameters (L1 penalty). This shrinks parameter estimates towards zero and, due to the non-smooth objective, selects a subset of variables with nonzero coefficients. The ridge estimator (Hoerl and Kennard, 1970) instead adds a penalty based on the sum of squares of the parameters (L2 penalty). This also has the effect of shrinking parameter estimates towards zero. The Elastic Net (Zou and Hastie, 2005) combines the L1 and L2 penalty. Many other penalty functions are possible. Penalized estimation allows researchers to include very large numbers of predictors in nowcasting models and, in many settings, results in nowcasts that have lower error than unpenalised alternatives, by reducing overfitting. These methods have been used in nowcasting in Babii et al. (2021), for example.

2.2 Missing observations and mixed frequency or irregular sampling

Due to varying publication lags of each data series, often some variables in a dataset have missing values for recent time periods. This issue is often referred to as the “ragged-edge” problem (Wallis, 1986). In many cases, one would like to predict low frequency variables (such as monthly or quarterly GDP) using high frequency ones (for example daily, weekly, and monthly prices or expenditure).

Several solutions to this problem have been developed in the macroeconomics literature. One option is to use the classic discrete time Kalman filter (e.g. Kalman, 1960; Stock and Watson, 2002; Bańbura and Modugno, 2014). For this, we assume a linear time series structure and impute missing values using the Kalman filter.

A second option is the bridge method (Schumacher, 2016). This approach uses a two-equation system, first aggregating a set of high frequency predictors to produce low frequency predictors and then using the low frequency predictors to predict the low frequency target.

Consider an example in which one wants to use a single monthly variable z_t to predict a quarterly target y_t . The bridge system can be written as

$$y_t = \sum_{s=1}^p \alpha_s y_{t-3s} + \beta_0 + \beta_1 x_t + \varepsilon_t \quad (1)$$

$$x_t = \sum_{s=0}^2 \gamma_s z_{t-s}. \quad (2)$$

Equation (1) is the autoregressive equation used to predict y_t , where p indicates the lag in the autoregressive process of the quarterly target, ε_t is an exogenous noise process. The parameters in (1) are typically estimated from the data. Equation (2) is the “bridge” equation that aggregates the high frequency variable across time. The parameters in the bridge equation typically are chosen ex ante. For example, if x_t is the quarterly average of z_t , then $\gamma_s = \frac{1}{3}$ for all s .

The bridge equation addresses the mixed frequency problem. To overcome the ragged edge problem, missing values of z_t are imputed using an auxiliary

forecasting model such as an AR(p) process

$$z_t = \delta_0 + \sum_{s=1}^p \delta_s z_{t-s} + \eta_t,$$

where η_t is a zero mean white-noise process, δ_0 and δ_s are parameters to be fitted.

For simple patterns of mixed frequency data (e.g. monthly predictors and quarterly targets), the bridge method is easy to use and doesn't require users to specify in detail the high frequency dynamics of the system. However, the aggregation method employed in the bridge equation itself is ad hoc and may need to be adapted for different contexts. Moreover, adapting the method to more complicated patterns of mixed frequency data (e.g. combining weekly, monthly or irregularly sampled data), can be challenging.

A third approach that directly relates high frequency indicators to a low frequency target is the mixed-data sampling (MIDAS) model (Ghysels et al., 2004). MIDAS accommodates data sampled at different frequencies by incorporating higher frequency variables using distributed lag terms with fractional lags. Mathematically, it can be written in a form that is very similar to the bridge equation, the distinction is that MIDAS employs a more general functional form for (2) and its parameters are estimated rather than specified ex-ante. An example of the MIDAS equation for a quarterly target y_t is given by:

$$y_t = \sum_{s=1}^p \alpha_s y_{t-3s} + \beta_0 + \beta_1 x_t + \varepsilon_t \quad (3)$$

$$x_t = \sum_{s=0}^p \gamma(s, \theta) z_{t-s} \quad (4)$$

where

$$\gamma(s, \theta) = \frac{\exp(\theta_1 s + \theta_2 s^2)}{\sum_{j=0}^p \exp(\theta_1 j + \theta_2 j^2)}.$$

The logistic functional form, also known as the exponential Almon lag, is used when p in (4) is large to avoid parameter proliferation.

To date, most empirical studies on nowcasting using the methodologies above deal with regularly sampled data with missing observations. This is because nowcasting typically utilises official published data which are released on a fairly regular schedule. However, in recent years there has been an increase in the use of alternative data sources, e.g. web-scraped data and supermarket scanner data. These alternative sources can be high-frequency and have complicated/irregular missingness patterns. Therefore, there is a need for a nowcasting methodology which can handle these cases.

2.3 Time-varying parameters and nonlinearities

Classic time series methods, such as the autoregressive integrated moving average (ARIMA) model, assume that the dynamics of a low-dimensional variable, after suitable differencing to ensure stationarity, can be modelled with a linear model with autoregressive and moving average components. That is, after differencing, the data are stationary, the model is linear, and the parameters of

the model are assumed to be constant. In nowcasting settings, all three of these assumptions can be restrictive.

State-space models offer one approach to relax these restrictions (Durbin and Koopman, 2012). While the linear Kalman filter is the best known, these models are highly flexible and can allow for non-stationarity, nonlinearity, and time-varying parameters. Many papers, including Hamilton (1994), Nielsen and Berg (2014), Carter and Kohn (1994), Kim et al. (1999), Kim (1994), also allow for time-varying parameters.

3 Nowcasting from a mathematical perspective

In this section, we outline some of the underlying mathematical theory for nowcasting and illustrate its connection to the path signature. We start with a recap of the discrete time Kalman filter and its continuous time extension, then introduce the theory of signatures, and finally show how regression on signatures subsumes the Kalman filter.

3.1 Discrete time state-space models

We begin by introducing the notation that we use throughout the paper and recapping the classic discrete time Kalman filter (see Bertsekas (2012) for an overview).

Suppose that we have a hidden process $Y \in \mathbb{R}^d$ that we cannot directly measure, but would like to infer from an observed process $X \in \mathbb{R}^m$. At each time t , the Kalman filter splits into two stages. The first is the *prediction* stage: given our previous estimate of Y_{t-1} , how can we model its evolution to Y_t ? The second is the *correction* stage: as more measurements are made, and we obtain information X_t , how can we use this to update the estimate of Y_t ?

To be precise, let us assume that the ground truth is given by

$$Y_t = AY_{t-1} + W_t, \quad X_t = CY_t + V_t,$$

with the starting distribution $Y_0 \sim N(\mu_{0|0}, P_{0|0})$. Here W, V are white noise processes in \mathbb{R}^d and \mathbb{R}^m respectively, with $W_t \sim N(0, \Gamma)$ and $V_t \sim N(0, \Sigma)$ for all t (all values independent). Note A and Γ are in $\mathbb{R}^{d \times d}$, $C \in \mathbb{R}^{m \times d}$, and $\Sigma \in \mathbb{R}^{m \times m}$. We assume all parameters are known, and write $\mathcal{X}_t = (X_1, \dots, X_t)$ to describe our observations up to time t .

For the prediction stage, we know that $Y_t | \mathcal{X}_t \equiv Y_t | (X_1, X_2, \dots, X_t)$ is normally distributed (and similarly $Y_t | \mathcal{X}_{t-1}$), so we write $Y_t | \mathcal{X}_s \sim N(\mu_{t|s}, P_{t|s})$. Using the dynamics of Y and X , we can obtain the prediction equations

$$\begin{aligned} \mu_{t|t-1} &\equiv E[Y_t | \mathcal{X}_{t-1}] = A\mu_{t-1|t-1}, \\ P_{t|t-1} &\equiv \text{var}(Y_t | \mathcal{X}_{t-1}) = AP_{t-1|t-1}A^\top + \Gamma. \end{aligned}$$

Before computing the Kalman correction step, let us define the ‘‘innovation’’ process η and its variance S

$$\begin{aligned} \eta_t &\equiv X_t - E[X_t | \mathcal{X}_{t-1}] = X_t - C\mu_{t|t-1}, \\ S_t &\equiv \text{var}(\eta_t | \mathcal{X}_{t-1}) = \text{var}(X_t | \mathcal{X}_{t-1}) = CP_{t|t-1}C^\top + \Sigma. \end{aligned}$$

The innovation process η tells us what information we “learn” from X_t . For the correction, we compute the new mean estimate to be

$$\mu_{t|t} \equiv E[Y_t|\mathcal{X}_t] = E[Y_t|X_t, \mathcal{X}_{t-1}] = \mu_{t|t-1} + K_t\eta_t,$$

where K is the “Kalman gain” process, which allows us to optimally incorporate new information,

$$K_t = P_{t|t-1}C^\top S_t^{-1} = (S_t^{-1}CP_{t|t-1})^\top.$$

Finally the variance correction equation is given by

$$P_{t|t} \equiv \text{var}(Y_t|X_t, \mathcal{X}_{t-1}) = (I - K_tC)P_{t|t-1}.$$

This method also generalises to the case where the parameters are time-varying.

3.2 Continuous time state-space models

In order to treat variable timings of observations, it is natural to embed the discrete-time model in a continuous-time framework. As pointed out elsewhere, “[a]lthough missing observations can be handled by a discrete time model, irregularly spaced observations cannot. Formulating the model in continuous time provides the solution. Furthermore, even if the observations are at regular intervals, a continuous time model has the attraction of not being tied to the time interval at which the observations happen to be made.” (Harvey (1990), p.479. See also Bergstrom (1984).) The most natural continuous-time extension of the Kalman state-space model is the Kalman–Bucy filter². As in the discrete time setting, we assume a hidden process Y and observed process X . We assume a ground truth of the form

$$\begin{aligned} dY_t &= (F_tY_t + f_t)dt + \sigma_t dV_t, \\ dX_t &= (H_tY_t + h_t)dt + dW_t, \end{aligned}$$

where $F \in \mathbb{R}^{d \times d}$, $\sigma \in \mathbb{R}^{d \times p}$, $f \in \mathbb{R}^d$, $H \in \mathbb{R}^{m \times d}$, $h \in \mathbb{R}^m$, so Y is d -dimensional, X is m -dimensional. The processes V and W are Brownian motions, and W is independent of Y .

The filter, which estimates the current state of the underlying process Y , is then characterised by the pair of stochastic differential equations

$$d\hat{Y}_t = (F_t\hat{Y}_t + f_t)dt + R_tH_t^\top(dX_t - (H_t\hat{Y}_t + h_t)dt), \quad (5)$$

$$\frac{dR_t}{dt} = \sigma_t\sigma_t^\top + F_tR_t + R_tF_t^\top - R_tH_t^\top H_tR_t. \quad (6)$$

Given our observations up to time t , the hidden process Y_t has conditional distribution $N(\hat{Y}_t, R_t)$. We can see that (6) is a first order ordinary differential equation which is quadratic in R_t – a Riccati equation – and therefore solve it to find R_t . We note that in particular that R_t is deterministic and does not depend on the observations X .

²For a detailed study of these equations, and derivation of the filter, see for example Bain and Crisan (2008) (whose notation we broadly follow) or Cohen and Elliott (2015, Chapter 22).

3.3 Paths and signatures

The path signature (Chen, 1958) is a property of continuous paths. It is motivated by the theory of rough paths, and has been shown to be an effective description of intertemporal information in prediction tasks. In this section, we introduce the terminology and definitions related to paths and signatures. For a more detailed introduction see Chevyrev and Kormilitzin (2016); Fermanian (2021); Lyons et al. (2007).

Formally, a path in \mathbb{R}^d is defined as a continuous function $X : [a, b] \rightarrow \mathbb{R}^d$, where each component is a one-dimensional path $X^k : [a, b] \rightarrow \mathbb{R}$.

We define the signature through an iterative construction. Let

$$S^1(X^k)_{a,t} \equiv \int_a^t dX_s^k = X_t^k - X_a^k,$$

and define the double iterated integral as

$$S^2(X^k X^l)_{a,t} \equiv \int_a^t S^1(X^k)_{a,s} dX_s^l = \int_a^t \int_a^s dX_r^k dX_s^l.$$

More generally define the n-fold iterated integral by

$$S^n(X^{k_1} X^{k_2} \dots X^{k_{n-1}} X^{k_n})_{a,t} \equiv \int_a^t S^{n-1}(X^{k_1} X^{k_2} \dots X^{k_{n-1}})_{a,s} dX_s^{k_n}.$$

Note that the superscript in $S^2(X^k X^l)_{a,t}$ denotes the number of iterated integrals we take, the subscript indicates the limits of the outermost integral, and the terms inside the brackets give the order of integration, starting with the innermost integral. We refer to the number of iterated integrals taken as its “level”. The signature of the path is the ordered infinite collection of all such terms

$$S(X)_{a,b} \equiv \left(1, S^1(X^1)_{a,b}, \dots, S^1(X^d)_{a,b}, S^2(X^1 X^1)_{a,b}, S^2(X^1 X^2)_{a,b}, \dots\right).$$

The path signature captures geometric information, such as the order of events. The first level signature terms give the change/increment in each dimension between the start and end of the path. The second level terms are linked to areas bounded by the path. See Figure 1 for a visual representation of the intervals and areas represented by the signature terms for a two-dimensional path. In particular, cross terms, $S^2(X^i X^j)_{a,b}$, $i \neq j$, are used to compute the Lévy area, which captures which dimensions tend to change before others. For a two-dimensional path, the Lévy area A is given by

$$A = \frac{1}{2} (S^2(X^1 X^2)_{a,b} - S^2(X^2 X^1)_{a,b}),$$

a positive value of A indicates X^1 is typically followed by X^2 . See Yang et al. (2017) for further discussion.

There is a Stone–Wierstrass theorem which gives universality of this approximation class (Levin et al., 2013). That is, any continuous function on paths can be approximated arbitrarily well through a linear combination of signature terms. Intuitively, linear functions of signatures behave like “polynomial functions of paths”.

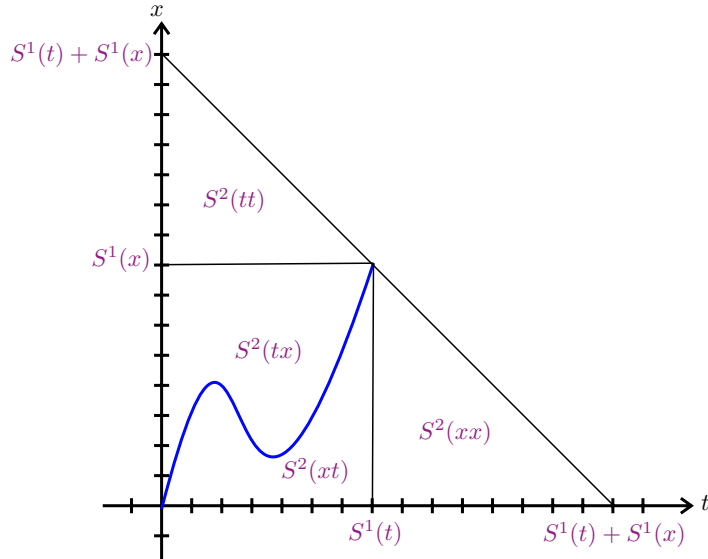


Figure 1: Illustration of the first 2 levels of the signature terms of a two-dimensional path (with variables t and x).

As signature terms are iterated path integrals, they also inherit their invariance properties, that is, the terms are invariant to translation of the path and to any monotonic increasing reparameterisation in time.

The theory to support applications of signatures is well-developed. For example, Chen’s identity (Lyons et al., 2007) gives us a way to compute the signature of a concatenated path as an algebraic product of individual paths, and yielding efficient computation of signatures for moving windows.

3.4 Practical signature computation

In the previous section, we saw that the signature is a natural object/basis to consider for paths and time-series data in particular. When working with real data, we only observe variables at discrete intervals. As a result, we need to interpret new data releases as coming from some underlying continuous time series and exploit connections to model the target variable. Using signatures as features is a natural way to make sense of data releases continuously in real-time and sidesteps data issues discussed below.

A common feature of datasets is that different variables are collected at different frequencies. Some things are easy to measure and we may get frequent updates, for example electricity and oil prices. Other data may be more expensive or difficult to measure, for example the unemployment rate or personal income. Another issue is that there may be missing data. Finally, the data collection can be very irregular and sporadic. By interpreting the time series as a path object, we can nevertheless use data collected in these settings.

The iterated integrals defining the signatures can be efficiently computed with the `iisignature` (Reizenstein and Graham, 2020) and `esig` packages in Python. Due to the invariance to reparameterisation property of signatures

mentioned in Section 3.3, it is common to manually add time as one of the input variables if the rate at which things occur is relevant (as is typical in prediction problems). Adding time as one of the variables is referred to as *time-augmentation*. This also ensures that signatures are unique (Hambly and Lyons, 2010).

The number of signature terms at each level is dependent on the dimension of the path, therefore provided the number of explanatory variables is fixed, the number of signatures at each level is fixed too. However, it is important to note that the number of terms is increasing exponentially with the level of signatures. For a d -dimensional path if we truncate at the k th level, then there are $\sum_k d^k$ number of signature terms. So whilst signatures can universally approximate any path objects, we need to be wary of the exponentially increasing number of terms, which would increase model complexity, and may result in over-parameterisation.

A classic result in rough paths is the factorial decay of the magnitude of signature terms with increasing levels (Lyons et al., 2007). Let $X : [0, T] \rightarrow \mathbb{R}^d$ be a path with finite one-variation (total variation), that is

$$\|X\|_{1,[0,T]} \equiv \sup_{P \in \mathcal{P}} \sum_{i=0}^{n_P-1} |X_{i+1} - X_i| < \infty,$$

where \mathcal{P} is the set of all partitions of $[0, T]$ and n_P is the number of points in partition P . Then, for each level $k \geq 1$,

$$S^k(X \dots X)_{[0,T]} = \int_0^T \dots \int_0^{t_1} dX_{t_0} \dots dX_{t_{k-1}} \leq \frac{\|X\|_{1,[0,T]}^k}{k!}. \quad (7)$$

Therefore, as we increase the level of signature terms, since the terms are decreasing factorially (which is faster than exponential), we still obtain a reasonable approximation by truncating the signature at a sufficiently high level (certain lower level terms may be increasing in the expanding window case as discussed previously).

There are two different ways to consider incoming data. We can either consider an *expanding window*, appending the new information to our existing data, or we can take a *rolling window*: fixing a window size and discarding older data.

In the case of the expanding window, we note that certain terms in the signature may be increasing. For example, the first level signatures terms are just the differences/increments between the start and the end of the path, and therefore the signatures corresponding to time can increase as more observations become available.

As signatures describe continuous paths, we interpolate our discrete observations before computing the signature. The choice of interpolation can matter, but if sufficiently similar paths are generated, we do not expect the signature to be greatly affected by the frequency of observation/sampling. There are many choices we can make to construct the path, and different methods are suitable depending on when we want to make a prediction. Choices of interpolation have been explored (Morrill et al., 2022). If we are making predictions only once, when we have all the predictive data, we can consider using a smooth interpolation method such as natural cubic splines. If data is measured at different times and frequencies, and we may want to make a prediction at any

point, then rectilinear interpolation is suitable. Since in nowcasting, we have an online problem where we want to make inferences at given points, then the interpolation method employed should be either continuously online or discretely online (see Morrill et al. (2022)). In practice, we typically use forward filling (discretely online) and rectilinear interpolation (continuously online).

It can be useful to add additional information by providing e.g. time since the last observation or counting the number of measurements observed in certain variables. These have been observed to boost performance in practice as the data may not be missing at random (Morrill et al., 2022).

In the last few years, signatures have been utilised in a range of applications with great success, these include Chinese handwriting recognition (Graham, 2013), malware detection (Cochrane et al., 2021), and time series generation (Ni et al., 2021).

3.5 Using signature as features in prediction tasks

One very useful property of truncated signatures is that if two paths are “close” in some sense (to be precise, under the 1-variation norm), then the truncated signatures of these paths will also be close (Giusti and Lee, 2020). Combined with the universal approximation property (by the Stone–Weierstrass Theorem (Levin et al., 2013)), signatures are a natural feature set to explore for prediction or classification tasks. Further, because linear combinations of signatures can effectively capture nonlinear relationships, even a simple regression model is theoretically sufficient. We briefly review the use of signatures as features and highlight some variations of “signature methods” in other applications.

Regression on signatures has previously been used in high-frequency financial time series (Lyons et al., 2014; Levin et al., 2013). More recently Fermanian (2020) looked at functional linear regression where information on predictors over time is available. This is similar to the setting of economic nowcasting. Fermanian (2020) showed that signature regression is competitive with traditional functional regression methods such as functional PCR and B-splines.

Often the truncation level is chosen through a hyperparameter search (Morrill et al., 2021b). Several works have discussed a more systematic approach to choosing the truncation depth. For example Fermanian (2020) assumes that the true model is linear in some truncated signature space, that is the model can be replicated with some m^* . An estimator for the truncation level \hat{m} is designed to minimise the sum of the mean square error of the model and a penalty based on model complexity. They then obtain a bound on the probability that the estimator \hat{m} differs from the true truncation level m^* , that is, (provided that there exist some bounds on the data, and for sufficiently large sample size n)

$$P(\hat{m} \neq m^*) \leq C_1 \exp(-C_2 n^k) \quad (8)$$

for some $k \in (0, 1)$. This estimator is used with Ridge regression for their functional regression model. See Fermanian (2020) for more details. In addition, results have also been obtained for the LASSO regularized case to show that the truncation error decays exponentially fast with the truncation level (Bleistein et al., 2023).

We note that there is some model tuning to be done in signature regressions, and attempts have been made to systematically categorize the type of

operations and model choices. For example, Morrill et al. (2021a) splits model processes which they called “modifications” into 4 primary groups: augmentation, windows, transforms, and rescaling, and then proceeds to an empirical analysis.

3.6 Signatures generalize (linear) Kalman filters

In this section we argue that the filter (5) can be *equivalently written* as a linear regression problem on the signature space. In other words, we can express \hat{Y} as a linear combination of the iterated integrals of the extended/time-augmented observation process (t, X_t) . For the sake of precision, we formally express this result as a theorem. This result holds in any dimension of observation and filter processes. Therefore, it includes the full range of (continuous time) vector autoregressive and linear state-space models. This suggests that linear regression on signatures has capacity to avoid many of the modelling restrictions needed in order to apply filtering methods in practical situations.

Theorem 1. *The continuous time Kalman–Bucy filter \hat{Y} , as described by (5), can be written as a linear function of the initial estimate \hat{Y}_0 and the signature of the augmented observation process (t, X_t) . In particular, this representation uses only the signature terms where the observation X appears once in the iterated integral.*

Proof. One approach to this result would be to use the generic property that *any* sufficiently continuous functional of an observation path can be represented in terms of the signature (see Lyons et al. (2007)). Here we instead give an explicit construction.

With the notation set out in Section 3.2, write

$$\begin{aligned} A_t &\equiv F_t - R_t H_t^\top H_t, \\ \xi_t &\equiv \int_0^t (f_s - R_s H_s^\top h_s) ds + \int_0^t R_s H_s^\top dX_s, \end{aligned} \quad (9)$$

which gives us the simplified expression for the filter

$$d\hat{Y}_t = A_t \hat{Y}_t dt + d\xi_t. \quad (10)$$

Denote the iterated integrals of A and ξ by

$$\begin{aligned} \mathbb{A}_t^n &= \int_0^t \int_0^{t_1} \cdots \int_0^{t_{n-1}} (A_{t_1} A_{t_2} \cdots A_{t_n}) dt_n \cdots dt_1 \\ \Xi_t^n &= \int_0^t \int_0^{t_1} \cdots \int_0^{t_{n-1}} (A_{t_1} A_{t_2} \cdots A_{t_n} \xi_{t_n}) dt_n \cdots dt_1, \end{aligned}$$

with the conventions $\mathbb{A}^0 = I_d$ (the identity matrix) and $\Xi_t^0 = \xi_t$. Observe that these are both formally solutions of the recurrence relation $Q_t^n = \int_0^t A_s Q_s^{n-1} ds$ with different values for Q^0 (and in different dimensions). We note that the n th iterated integral is super-polynomially small (see (7)), and in particular the infinite sums $\sum_{n \geq 0} \mathbb{A}_t^n$ and $\sum_{n \geq 0} \Xi_t^n$ are both well defined.

Writing \hat{Y} in integral form, we have

$$\hat{Y}_t = \hat{Y}_0 + \int_0^t A_s \hat{Y}_s ds + \xi_t.$$

It is then easy to check that the value of \hat{Y} can be expressed through the series solution

$$\hat{Y}_t = \sum_{n \geq 0} \left(\mathbb{A}_t^n \hat{Y}_0 + \Xi_t^n \right).$$

Given the recurrence relation mentioned above, we can see that this is the (unique) solution to the integral equation. This shows that \hat{Y}_t is a (linear) function of its initial value \hat{Y}_0 and the iterated integral processes \mathbb{A}^n and Ξ^n . This expansion is very closely related to the Picard series approximation of the stochastic differential equation (5) defining \hat{Y} .

It remains to show that \mathbb{A} and Ξ can be expressed in terms of the (joint) signature of the time-augmented path (t, X_t) . As A is a continuous deterministic function of time, we know that (over any finite time horizon) it can be approximated arbitrarily well by a polynomial (by the Stone–Weierstrass theorem). As the signature of time is simply the sequence $\mathbb{S}(t) = (1, t, t^2/2, \dots, t^n/n!, \dots)$, we see that A can be written as a (matrix-valued) linear function of the signature of t .

The process ξ in (9) is slightly more delicate, as it depends on both time and the observations X . The first term $\int_0^t (f_s - R_s H_s^\top h_s) ds$ is deterministic, so again has a polynomial expansion in terms of the signature of t . Considering the second term $\int_0^t R_s H_s^\top dX_s$, we see that if H is continuous³, $R_s H_s^\top$ has an expansion in terms of the signature of time, so $\int_0^t R_s H_s^\top dX_s$ has an expansion in terms of the signature of (t, X_t) , with the special form where the only integral with respect to X is the outermost one.

As both A and ξ have expansions in terms of the signatures of t and (t, X) respectively, it follows that their iterated integrals \mathbb{A} and Ξ also have expansions of this type. For our purposes the explicit values of this expansion are not of particular interest (and do not have a simple algebraic form), but the definition of \mathbb{A}^n immediately shows that if A can be written as a polynomial in time, then so can \mathbb{A}^n for each n . Similarly for Ξ^n , but now this will involve iterated integrals with a single X integral included.

□

The absence of iterated integrals with more than one X integral is equivalent to the linear dependence of the filter \hat{Y} on the observations X . We refer to the iterated integrals with only a single X integral as “linear” signature terms.

Lemma 1. *For a continuous path \hat{Y} , and any time $t \in [0, T]$, the value of \hat{Y}_0 can be expressed as a linear function of \hat{Y}_t and the linear signature terms of (t, \hat{Y}) .*

Proof. The iterated integrals of time give polynomials (of the form $t^n/n!$). By

³If H is not continuous, then we need to use the fact that polynomials in time are dense in the $L^2([0, t])$ space, which is the appropriate space to consider given we have an outer integral with respect to the process X . In this case we will still have a polynomial approximation $p_n(s) \approx R_s H_s^\top$, such that the integral $\int_0^t p_n(s) dX_s \rightarrow \int_0^t R_s H_s^\top dX_s$ converges in a mean-square sense as $n \rightarrow \infty$.

using the inner products

$$\begin{aligned}\langle \hat{Y}, t^n/n! \rangle_{L^2} &= \frac{1}{n!} \int_0^T \hat{Y}_s s^n ds \\ &= \hat{Y}_0 \frac{T^n}{n!} + \frac{1}{n!} \int_0^T \left(\int_0^s d\hat{Y}_u \right) s^n ds\end{aligned}$$

and similarly $\langle t^m/m!, t^n/n! \rangle$. A simple application of integration by parts shows that the final integral term of the above equation can be expressed as the sum of two signature terms, of the form $S^n(\hat{y}t..t)$ and $S^n(t...t\hat{y})$.

Suppose we compute a regression expansion of Y_t in terms of $\{t^m/m!\}_{m \leq M}$. As \hat{Y} is a continuous path, and the polynomials are dense in the L^2 space on $[0, T]$, this (theoretical) expansion perfectly approximates Y_t as we take $M \rightarrow \infty$. It follows that \hat{Y}_t can be evaluated by evaluating a linear function of the signature terms and \hat{Y}_0 . Rearrangement gives the desired expression for \hat{Y}_0 in terms of \hat{Y}_t . \square

The next lemma shows that we have great flexibility when constructing our signature expansion for the filter, as we can use the signature over any sufficiently large horizon in our representation of the filter.

Lemma 2. *We can express the Kalman–Bucy filter \hat{Y}_t as a linear function of \hat{Y}_s and the linear signature of X on $[s', T]$, for any $s' \leq s < t$.*

Proof. By translation, we can assume $s' = 0$. This result then follows from Lemma 1 and observing that, as \hat{Y} can be expressed as a linear function of the linear signature of X , the linear signature terms of \hat{Y} can also be expressed as a linear function of the linear signature of X . \square

We conclude with an example where we explicitly compute the optimal filter in terms of the signature.

Example 1. *Consider the situation where $d = p = m = 1$, $f = h = 0$, $F = -1$, $H = 1$ and $\sigma = \sqrt{3}$. Then our filter equations simplify to*

$$\begin{aligned}d\hat{Y}_t &= -\hat{Y}_t dt + R_t(dX_t - \hat{Y}_t dt), \\ \frac{dR_t}{dt} &= 3 - 2R_t - R_t^2.\end{aligned}$$

For simplicity, assume the initial variance is at the steady state $R_0 = R_t = 1$, so we have

$$d\hat{Y}_t = -2\hat{Y}_t dt + dX_t,$$

which can be solved as

$$\hat{Y}_t = e^{-2t}\hat{Y}_0 + \int_0^t e^{-2(t-s)} dX_s.$$

Assuming $X_0 = 0$, the signature expansion of \hat{Y} can also be computed, as was

done for (10), with the identity $A_t = -2$ and $\xi_t = X_t$, to give⁴

$$\begin{aligned}\mathbb{A}_t^n &= (-2)^n \frac{t^n}{n!} = (-2)^n S^n(t \cdots t) \\ \Xi_t^n &= (-2)^n S^{n+1}(xtt \cdots t),\end{aligned}$$

and hence

$$\begin{aligned}\hat{Y}_t &= \sum_n \left(\mathbb{A}_t^n \hat{Y}_0 + \Xi_t^n \right) \\ &= \hat{Y}_0 + \sum_{n \geq 1} \left([(-2)^n \hat{Y}_0] S^n(t \cdots t) + [(-2)^{n-1}] S^n(xtt \cdots t) \right) \\ &= \hat{Y}_0 \left(1 - 2t + 2t^2 - \frac{8}{6}t^3 + \frac{16}{24}t^4 + \dots \right) \\ &\quad + X_t - 2 \int_0^t X_s ds + 4 \int_0^t \int_0^{t_1} X_s ds dt_1 \\ &\quad - 8 \int_0^t \int_0^{t_1} \int_0^{t_2} X_s ds dt_1 dt_2 + \dots\end{aligned}$$

In particular, if t is small, the first few terms of this series provide a good approximation for the value of \hat{Y}_t , in terms of a linear function of the signature. Note that $\hat{Y}_t \approx \hat{Y}_0 - 2\hat{Y}_0 t + X_t$, the single-step Euler–Maruyama approximation of (5), appears as the first terms in this expansion.

The simple structure we obtain here is due to the assumptions we have made on our state-space model (in particular, the absence of signature terms of the form $S^n(t \dots txt \dots t)$ is due to the assumption the variance is in its steady state).

The key advantage of this approach, even while restricting our attention to the Kalman–Bucy state-space model, is that we now have an expansion which is valid, in theory for all t , and in practice for all t not too large. This simplifies dramatically the problem of working with data at mixed frequencies, as we can evaluate the filter state at any t , in terms of the corresponding signature terms, rather than having to compute (as is done, for example, in a MIDAS model) a version of the filter which depends on the timing of observations.

Nonlinear examples

A further advantage of signature methods is that they allow us to easily incorporate nonlinearity in our state-space models. For example, if our observations X were replaced in the Kalman–Bucy setting by $\bar{X} = \exp(X)$. Then, by Itô’s chain rule, we know that

$$d\bar{X}_t = \bar{X}_t dX_t + \frac{1}{2} \bar{X}_t dt \iff dX_t = \frac{1}{\bar{X}_t} d\bar{X}_t - \frac{1}{2} dt.$$

Substituting into (5), we see that the Kalman–Bucy filter can be written in terms of the modified observation process \bar{X} , and by very similar arguments to

⁴Here we write $S^n(\dots)$ for the n -fold iterated integral with respect to the sequence indicated (which must be of length n , with the innermost integral listed first); i.e. $S^3(xtt) = \int_0^t \int_0^{t_1} \int_0^{t_2} 1 dX_{t_3} dt_2 dt_1$. Note that in this example all signature terms are taken over the interval $[0, t]$, so we drop the subscript on signatures for convenience.

before, this would have an expression in terms of the signature, but with terms involving multiple integrals with respect to \bar{X} .

This suggests that using the signature expansion is robust to the specification of the observation time series. That is, the traditional issues around the choice of transformations (whether to use a series, its logarithm, differences, etc...) are less significant, as they can be absorbed into the signature expansion. The price that we pay to capture these nonlinear relationships is that there are multiple integrals with respect to \bar{X} , so we lose the ability to write the filter with only the “linear” signature terms and hence all signature terms should be retained in model fitting.

4 Nowcasting via signature regression

Now that we have demonstrated that the Kalman filter can be represented in terms of the signature, we gain an easy approach to nowcasting, using regression on the signatures of our observations. In this section we focus on the practicalities of this method.

Let Y_t be the (low-frequency) target variable we want to nowcast. We write X_t for the (high-frequency) observed explanatory variables at t . Motivated by the approximation of the Kalman filter in terms of signatures, we assume that for some truncation level K we have a model of the form

$$Y_t = \sum_{k=0}^K (\alpha_k + \beta_k Y_{t-}) \psi_{k,t} + \epsilon_t, \quad (11)$$

where

- Y_{t-} is a past observation of the low frequency target variable, available at the time of nowcast. This can correspond to the target variable at the beginning of the lookback window over which the signatures are computed, or the most recent observation;
- ϵ_t is a mean-zero error term which we assume is stationary;
- $\psi_{k,t}$ is a sequence (for each value of t) of signature terms at level k , including iterated integrals of t and the different components of the observed process X , calculated over a lookback window ending at the present time;
- α_k, β_k are vectors of regression coefficients (the dimension of which depends on k).

As outlined in Section 3.6, in order to replicate the Kalman filter, we would have $\beta_k = 0$ whenever ψ_k corresponds to a signature term depending on X , and α_k, β_k nonzero only for those signature terms which depend either purely on t , or on a single component of X appearing once in the iterated integral. Both of these restrictions can be relaxed, leading to a richer class of models than considered by the (linear) Kalman filter.

As discussed in Section 3.5, there are a variety of practices for choosing the signature truncation level K . We follow Morrill et al. (2021b) and select the level through hyperparameter optimisation. We search over a range of levels (in features and time) usually up to level 4 as that is enough to capture the

changes in the area between time-series (see Section 3.3) and higher dimensional equivalents. Hyperparameters are obtained through running the models over a validation period. The best hyperparameters are used to fit the final model on the whole training and validation period, and this calibrated model is applied to the test period.

Remark 1. There is a challenge when performing this regression, as the signature terms are typically highly multicollinear. This can be addressed through any standard technique, e.g. using PCA to capture the most important information, or using only a smaller number of terms in the expansion.

4.1 Regularisation

As the number of signature terms can still be large, particularly if we take a higher truncation level, the regression framework allows for well-known regularisation techniques to be applied to reduce overfitting. In particular, we can frame this with the elastic net (Zou and Hastie, 2005), which includes both L1 penalty and L2 penalty terms, and therefore also performs feature selection. The optimisation problem being solved is given by

$$\min_{\alpha_k, \beta_k} \left\{ \frac{1}{n} \left\| Y_t - \sum_{k=0}^K (\alpha_k + \beta_k Y_{t-k}) \psi_{k,t} \right\|_2^2 + \sum_{k=0}^K \left(\gamma \lambda (\|\alpha_k\|_1 + \|\beta_k\|_1) + \gamma (1 - \lambda) (\|\alpha_k\|_2^2 + \|\beta_k\|_2^2) \right) \right\},$$

where n is the number of time points that data are collected from, γ is the regularisation strength parameter, and $\lambda \in (0, 1)$ is the L1 ratio. The parameters of the elastic net, λ and γ are found through hyperparameter optimisation.

Recall that signature terms are decreasing at a factorial rate with the truncation level (7). If we introduce regularisation, then the signature terms should be standardised/normalised, else we risk eliminating all the higher level terms due to their decreased magnitude. Therefore, as default, we typically standardise the signatures once the truncated terms are obtained (before using them in regression) with the `StandardScaler` from `sklearn`.

4.2 Rolling windows for signatures

In Section 3.6 we saw that we can replicate the Kalman filter with a linear combination of signatures, where the linear signature terms are multiplied by the true value of the hidden process at some past time. In the context of nowcasting, to account for non-stationary behaviour of the time series, we can take a rolling/sliding window as discussed in Section 3.4. Given we want to maximise the use of alternative data sources that are more timely than the target series, our window of information should include at least the time of the last target variable measurement. Therefore, it is reasonable to assume that the “true” value of the target variable at some point in the window that we compute signatures will be available to us, which we can then use as a multiplier. This is denoted by Y_t^- in (11). Alternatively, a past estimate can be used. As explained in Lemma 2, the multiplier can be at any fixed time within the lookback window (different past times will result in different regression coefficients).

4.3 Framework

We set out the proposed framework in two algorithms. Algorithm 1 gives the procedures to fit a single signature regression model on a set of cleaned data. The full proposed signature regression method pipeline is as detailed in Algorithm 2. Note that here we laid out the hyperparameter optimisation in the pipeline with grid search, other hyperparameter search strategies can also be easily incorporated.

The full list of configuration settings and a brief explanation can be found in Appendix B.

Algorithm 1: Fit signature regression model

Input:

- cleaned/pre-processed data available in a wide table, pivot format, which includes the observed data and the target variable;
- target variable name;
- data parameters such as the publication lag in the target variable;
- a set of model parameters associated with the number of signature terms such as the truncation level, lookback window length, whether to use the previous known value of the target variable.

Process:

1. Define the auxiliary dataframe of observations available.
2. Convert the path information of the observations into truncated signatures over the lookback window.
3. Split signatures into training data and test data. The train data can be a proportion of the data available, or it can be the maximum data available (all timepoints where the target is defined).
4. Fit a (regularised) regression model.

Return: fitted model

5 Simulation Exercise

We have seen that the signature method theoretically subsumes the linear Kalman filter in Section 3.6. In this section we show that it is practically possible to almost replicate the performance of the Kalman filter by using regression on signatures (as described in Section 4) for experiments with simulated data.

5.1 Experiment set-up

In Example 1, by assuming that the variance is in a steady state, we derived the coefficients of the optimal filter \hat{Y} with respect to its infinite signature expansion. In particular, due to our assumptions, the only non-trivial terms

Algorithm 2: Signature regression framework

Input:

- data available (observation & target) in a wide table, pivot format;
- target variable name;
- data parameters, e.g. publication lag in the target variable;
- model parameters associated with signature regression, e.g. the truncation level, lookback window length, whether to use the previous known value of the target etc, these may be given as lists to hyperparameter search over.

Impute missing values in the dataset by specified method (defaults to forward fill for all values except those at the beginning which are filled by backward fill).

```
for each set of hyperparameters/configurations do
  for each time over the hyperparameter optimisation period
    (validation set) do
      1. Fit a signature regression model as outlined in Algorithm 1.
      2. Evaluate the hidden data set with the regression model.
      3. Store predictions and errors.
    end
  end
end
Identify the best set of hyperparameters.
if Recursive nowcasts then
  for each time that a nowcast is required do
    1. Fit a signature regression model as outlined in Algorithm 1.
    2. Evaluate the test set (only the time for the current nowcast) with the
      regression model for a nowcast value.
  end
else
  1. Combine the train and validation set and fit a signature regression model
    as outlined in Algorithm 1.
  2. Evaluate the test set with the above trained regression model.
end
Return nowcast values.
```

in the expansion are signatures that we refer to as “linear” (where only one of the iterated integral is with respect to the observed data, the rest are time integrals), and further, the only integral with respect to the data variables is the innermost one.

We simulate data from a 1-dimensional example with dynamics similar to that in Example 1, where $h = f = 0$ and $F = -1$, but now we set $H = 10$ and $\sigma^2 = 2$ for a higher signal-to-noise ratio. To be precise, we suppose each simulated path starts at $Y_0 = 0.1$, and follows the dynamics

$$\begin{aligned} dY_t &= -Y_t dt + \sqrt{2}dV_t, \\ dX_t &= 10Y_t dt + dW_t. \end{aligned}$$

Assuming a steady state variance, we can compute R as

$$\frac{F + \sqrt{F^2 + \sigma^2 H^2}}{H^2}.$$

The end time, T , is randomly sampled as Uniform on $[0.1, 1]$. Let us take the mesh size in time to be $\Delta t = 0.005$. Initially, to better understand the limitations of the signature method, we do not drop any of the data and therefore the time series is regularly sampled.

Here the dynamics are linear and if the parameter values were assumed known, that is, we do not need to learn any parameters from data, then the Kalman filter should give the optimal inference of Y_t (in the sense that the mean square error is minimised). The mean of the filter is given by (5). The discretised optimal filter is then given by

$$\hat{Y}_{t+\Delta t} = \hat{Y}_t + (F - RH^2)\hat{Y}_t\Delta t + RH(X_{t+\Delta t} - X_t).$$

To evaluate regression on signatures against this optimal filter with perfect information on the parameter values, we proceed as detailed in Section 4. Unlike for the Kalman filter, the signature method does not assume any parameter values are known and everything has to be inferred from the data available. We compute signatures of each path, X_t , truncated at depth 6, keeping only the linear signatures of the form $S^n(xtt\dots t)$ as discussed (Example 1). For this reason, we compare with a large training set. We simulate 1000 independent paths (for both X_t and Y_t) of which we take 800 to be in the training set (to calibrate the regression model) and the remaining 200 in the test/evaluation set.

5.2 Results

The residuals after applying the Kalman filter on the test set have a mean of 0.02 and a variance of 0.12, and the signature method residuals have a mean of 0.02 and a variance of 0.14. The performance of the signature method is comparable to the Kalman filter here. The histograms of the residuals can be seen in Figure 2.

We can see the residuals of the Kalman filter plotted against the residuals of the signature method in Figure 3. The line of best fit in this case has gradient 1.00 and a y -intercept of -0.02 . The R^2 of the regression on these residuals is 0.99. Therefore we see excellent alignment of the errors of the signature method (where the parameters are fitted from data) against the ideal filter (given full information).

After learning the linear model on the signature terms, we may apply the fitted model recursively given a new path of observations to infer an estimated

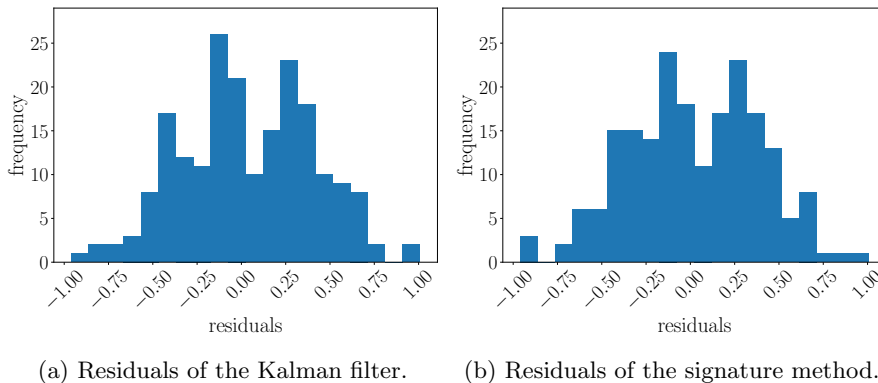


Figure 2: Histograms of residuals for predictions on simulation data.

path for the underlying true values. The inferred paths of the Kalman filter (also applied recursively with the same window length as the signature method for a fair illustration) and the fitted signature model on one particular simulated path are given in Figure 4. We see that the signature method predicts very similarly to the Kalman filter (despite the additional difficulty of fitting parameters from data), and the inferred path is close to the true underlying path, Y_t .

In this case study, we can also compare how close our learnt parameters are against the true infinite expansion coefficients (similar to those derived in Example 1). We find that if we use many terms (higher truncation order) corresponding to only integrals in t , then there are large coefficient differences due to multicollinearity. Further details can be found in Appendix A.1. To mitigate this, we can take different (lower) truncation levels for those signature terms corresponding to only integrals in t . We note further that some differences between the coefficients are still expected, as the best coefficients for the infinite expansion may not correspond to the best coefficients for the finite truncated expansion.

5.3 Irregularly-sampled data

We next downsample to 20% of our data in each path randomly, so that the dataset is effectively composed of irregularly sampled time series, and check the performance of the signature method. We recalibrate our model using this reduced dataset. We see in Figure 5 the plot of the residuals from the signature method against those from the Kalman filter (adjusted for irregular sampling). The Kalman filter residuals have a mean of 0.02 and a variance of 0.13 and the signatures residuals also have a mean of 0.01 and a variance of 0.14. The line of best fit in this case has gradient 0.93 and y -intercept of -0.02 , with an R^2 value of 0.93. If we take the same simulated path used for Figure 4, remove 80% of the data, then the inferred paths from the two methods can be seen in Figure 6. In this subsampled case, we see that the signature method still learns a good approximation for the Kalman filter.

This experiment demonstrates that regression on signatures provides a competitive method to infer parameters from data even in the case that the data is not regularly-sampled, and further, no additional modifications have to be

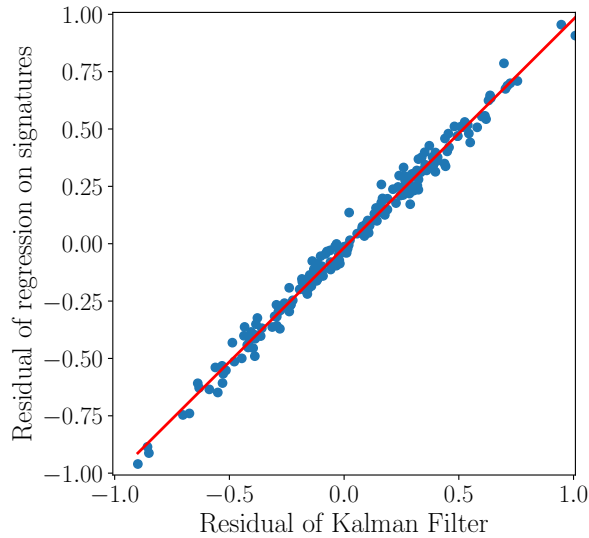


Figure 3: Residuals of the Kalman filter against the signature method on regularly sampled data. The red line is the line-of-best-fit and has gradient 1.00 and a y -intercept of -0.02 .

made to the method to achieve this.

5.4 Modifications for nonlinear transformation

As discussed in Section 3.6, the signature method can also be applied to nonlinear problems. In this subsection, we demonstrate this flexibility by applying a sigmoid transform to the input data of our simulation example, that is, our new “observed” data are given by

$$\bar{X} = g(X) = \frac{1}{1 + e^{-X}},$$

with the derivative of the inverse of g given by

$$\frac{dg^{-1}}{d\bar{X}} =: \bar{g}(\bar{X}) = \frac{1}{\bar{X}(1 - \bar{X})}.$$

The optimal filter is obtained by using $\bar{g}(\bar{X})d\bar{X}$ in the place of dX in (5).

The signature method can be applied to this transformed problem almost off-the-shelf. The only thing to bear in mind is that we lose the ability to represent the filter by only the innermost linear signatures. For this regression problem, we therefore keep all signatures of a certain (specified) truncation level.

Results on regularly sampled data

After applying the sigmoid transform on the observed data X_t , we use all signatures up to level 3 in the regression instead of just the innermost linear signatures. This is so that we have a comparable number of signature features to previous experiments (15 signature features here compared with 13 features

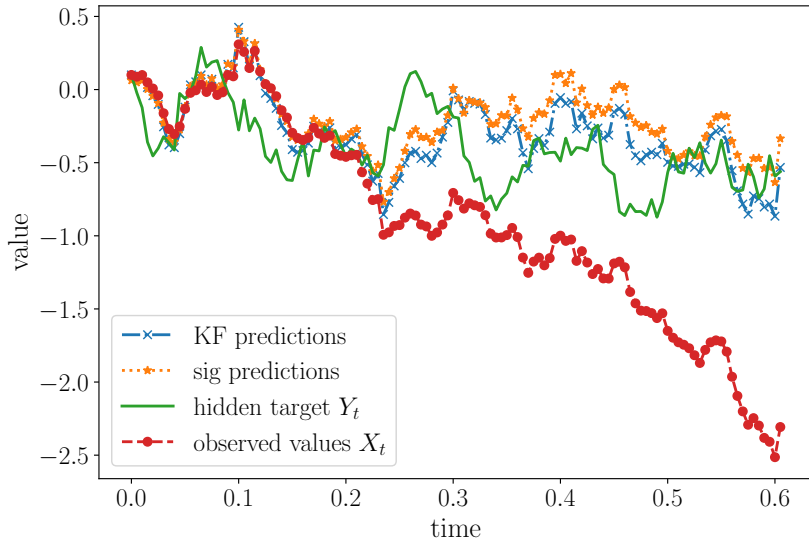


Figure 4: Illustration of one regularly sampled simulated path - the observed values (X_t) along with the true, hidden values that needs to be inferred (Y_t). The trajectories inferred from X_t with the Kalman filter and the signature method are also shown.

previously). The Kalman filter has a mean of 0.005 and a variance of 0.13 and the signatures residuals have a mean of -0.002 and a variance of 0.21. The line of best fit in this case has gradient 0.86 and y -intercept of -0.02 , and the regression fit has R^2 value of 0.50. The correlation has weakened compared with previously, indicating that the nonlinear relationship may be more difficult to learn.

Using the same trajectory data as that for Figures 4 and 6, but now with the observed data being passed through a sigmoid transform, the two methods give inferred paths as seen in Figure 7. Here we see that despite the observed data \bar{X}_t having a nonlinear relationship with the hidden/true values Y_t , we can still apply regression on the signature terms to obtain good predictions closely matching the theoretical optimum.

Results on irregularly sampled data

Finally, we apply the sigmoid transformation and then irregularly subsample the data (removing 80% of the observations in each path). In this case, the Kalman filter residuals have a mean of -0.01 and a variance of 0.29 and the signatures residuals have a mean of 0.001 and a variance of 0.21. The line of best fit in this case has gradient 0.23 and y -intercept of 0.001. The R^2 value is only 0.09. We see that there is very little correlation between the two methods in the presence of both a nonlinear transform and irregular subsampling, but the signature method still attains competitive results.

To illustrate the fitted model, in Figure 8, we plot the results on the same simulated path as in Figure 7 but now the observations \bar{X}_t have been irregularly subsampled. We see that despite a weaker correlation between the signature

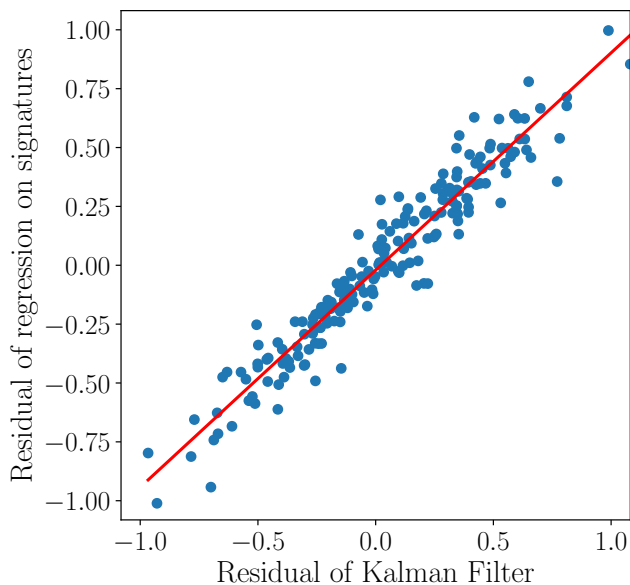


Figure 5: Residuals of the Kalman filter against the signature method on irregularly sampled data. The red line gives the line-of-best-fit, which has gradient 0.92 and a y -intercept of -0.02 .

method residuals and the Kalman filter residuals, the predictions are not dissimilar. Therefore regression on signatures can pick out the key information even when the data are more sparse and nonlinear.

5.5 Summary

In this section, we compared regression on signatures (fitting the parameters from the data) against the Kalman filter (assuming perfect knowledge of the system). We see that the residuals of the signature method are highly correlated with those from the Kalman filter, except in the case of combining a nonlinear transform and irregular subsampling. However, even in that case, the mean and variance of the residuals of the signature method are still competitive with the theoretical optimum.

This highlights the fact that the signatures can handle the combination of nonlinear transformation of the data and irregularly sampled data without the need for further modifications to the method.

6 Nowcasting US GDP growth

Official quarterly GDP estimates are published some time after the reference quarter, posing challenges to economists and practitioners when monitoring the state of the economy in real-time. In order to detect economic fluctuations in a more timely manner, economists have sought alternative data sources/indicators with higher observed frequencies to be adopted in nowcasting methods. These economic time series may have similar trends indicating that they are driven by

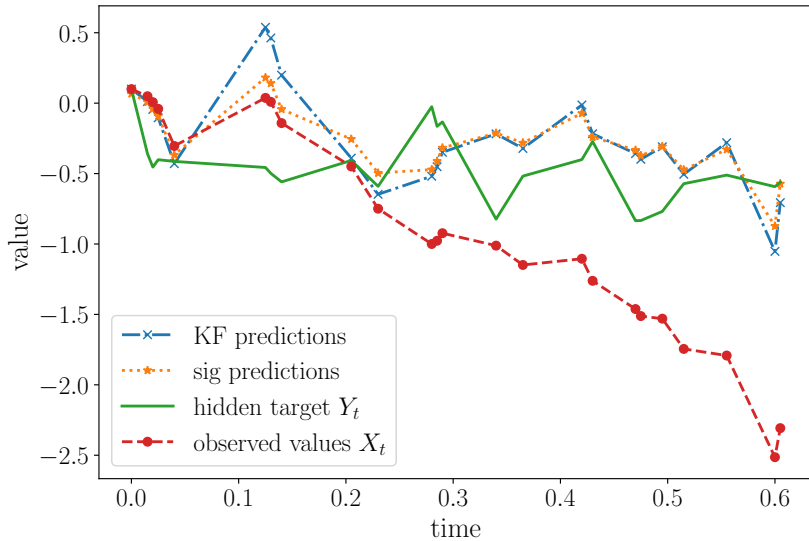


Figure 6: Illustration of the same simulated path as in Figure 4 but now irregularly sampled. The plot gives the observed values (X_t) along with the true, hidden values that needs to be inferred (Y_t), as well as the predicted trajectories from the Kalman filter and the signature method.

only low-dimensional hidden factors, making the dynamic factor model (DFM) (Geweke, 1977; Giannone et al., 2008) a suitable candidate method.

In this section we apply signature regression (as described in Section 4) to nowcast the quarter on quarter (QoQ) GDP growth for the United States, which we henceforth refer to as simply the “US GDP”. This target variable is chosen as there is a well-known DFM described by Bok et al. (2018) that is utilised for producing nowcasts, which are available to view online and can be used to benchmark the signature method. This will be referred to as the “New York Fed Staff Nowcast model” or “NYFed model”. Since not all time series used in the NYFed model are publicly available, we also fit our own dynamic factor model based on the NYFed model (Bok et al., 2018) by utilising broadly the same underlying variables and autoregressive structure, and compare this result with regression on signatures.

6.1 Data

In order to nowcast the US GDP, we use 33 of the 37 variables listed in Bok et al. (2018), since the remaining four variables (“ISM nonmanufactory: NMI composite index”, “ISM mfg.: Prices index”, “ISM mfg.: PMI composite index”, and “ISM mfg.: Employment index”) are not publicly available. We use University of Michigan: Consumer Sentiment (Curtin, 2008) as a replacement for these ISM series, giving us 34 variables in total. The dataset includes a selection of monthly variables that covers housing, income, manufacturing, labour, surveys, trade, and consumption. The indicators are monthly, but due to varying publication lags, new information is obtained each week.

Following Bok et al. (2018), we apply the same factor structure. A “soft”

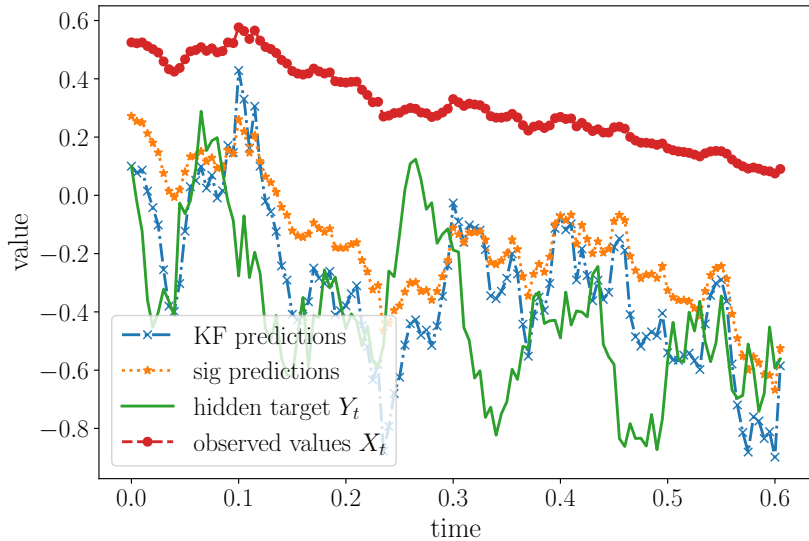


Figure 7: Illustration of the same simulated path as in Figure 4 but now the observed data has been passed through a sigmoid transform. The plot gives the transformed observed values (\bar{X}_t) along with the true, hidden values (Y_t) that needs to be inferred, as well as the predicted trajectories from the Kalman filter and the signature method.

block is included to model information from survey data; “real” and “labour” factors to model real and labour market variables respectively, and finally a “global” factor that affects all the variables. For the exact breakdown details see Appendix C as well as Table 5.1 of Bok et al. (2018).

Caveats

To avoid the complication of data revisions and so that the performance differences can be attributed to the method, we take the latest vintage of each data series. Therefore, the models are run on a different dataset from the real-time NY Fed Staff nowcast predictions. Consequently, we do not draw direct comparisons with Bok et al. (2018). Our baseline model is thus a DFM based on the NYFed model, with the same factor structure, which uses the latest (publicly) available data to learn the latest value of the published GDP.

Further, we note that the NY Fed model recalibrates the parameters every quarter, but our models are recalibrated each time we get new information.

6.2 Method

DFM

The nowcasting problem can be written in the form of a state-space model and hence inference can be done using Kalman filtering techniques as detailed in Section 3. From the discussion in Section 3.6, we see that linear regression on signatures generalises a Kalman filter. As the DFM can also be cast in the

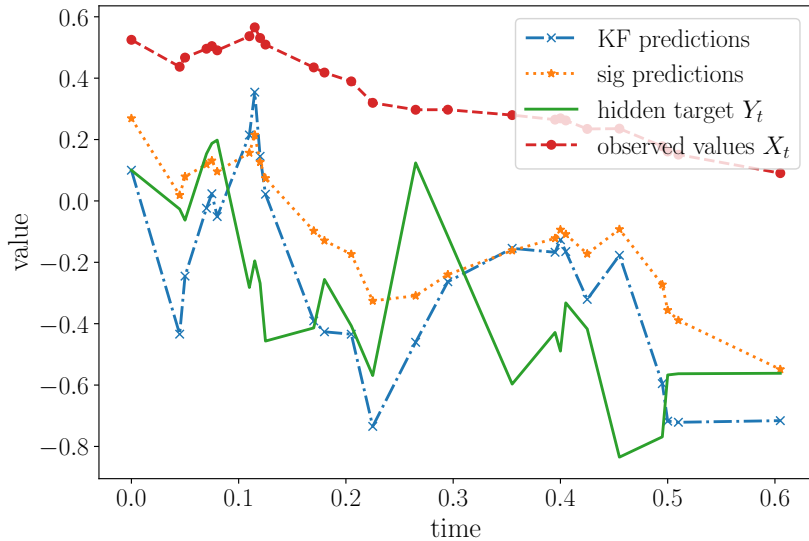


Figure 8: Illustration of the same simulated path as in Figure 7 (that is, after a sigmoid transform) but now irregularly sampled. The plot gives the transformed observed values along with the true, hidden values that needs to be inferred, as well as the predicted trajectories from the Kalman filter and the signature method.

state-space model form, this makes it a natural benchmark against which we compare the signature method.

We briefly outline the DFM and specific choices for NYFed model as discussed in Bok et al. (2018), upon which we base our own DFM. A dynamic factor model assumes that the n observed variables, $x_t \in \mathbb{R}^n$, are driven by r unobserved factors $f_t = (f_{1,t}, \dots, f_{r,t})^\top$. The state-space model is then given by

$$\begin{aligned} x_t &= \Lambda f_t + e_t, \\ f_t &= \sum_{i=1}^p a_i f_{t-i} + v_t, \\ e_t &= \sum_{i=1}^q b_i e_{t-i} + w_t, \end{aligned}$$

where $\Lambda \in \mathbb{R}^{n \times r}$ represents the factor loading, $e_t \in \mathbb{R}^n$ is the idiosyncratic noise process in the observations, and $v_t \in \mathbb{R}^r$ and $w_t \in \mathbb{R}^n$ are independent Gaussian random noise. We have p and q as the orders of the autoregressive processes of the factors and the idiosyncratic noise process respectively. Often $p = q = 1$ is assumed, as is the case in the NYFed model (Bok et al., 2018), which we also adopt for our replication of the model.

A model of this kind was first introduced by Geweke (1977). Subsequently Stock and Watson (1989) used this to extract one-dimensional common factors ($r = 1$) from a set of monthly indicators as new economic indices, which were then extended by others (e.g. Mariano and Murasawa, 2003; Aruoba et al., 2009). Large datasets often pose the issue of dealing with “large n (the number

of predictors), small T (the length of time series)” and how best to infer the hidden factors. Bok et al. (2018) extract the factors from the observed data with the following procedure. The factors are initialised by finding the principal components of each factor group. Then there is a two-step procedure. First, given the estimated factors, apply regression to find the best model parameters. Second, given the estimated parameters, update the common factors using a Kalman smoother and the EM algorithm (see Baum and Petrie (1966); Baum and Eagon (1967); Baum et al. (1970)). Maximum likelihood estimation is obtained by iterating these two steps until convergence.

Signature regression

We compare this model with linear regression on signatures as presented in Section 4. For regression on signatures, we use the same factor structure/groupings as in the dynamic factor model. Each time we receive new information, we use principal component analysis to find the first principal component of each factor group, listed in Appendix C. We proceed to use these 4 components along with the date as predictors (i.e. we have a time-augmented path) and compute signatures on the time series of these “factors” to be used in regression. We also use the latest available GDP value (which has a 30-day publication delay after the end of the quarter) in the analysis so that we are essentially learning the change in GDP growth.

Given the discussion on the increase in number of signature terms at each truncation level, we use Elastic net for regularisation. Since higher order signature terms are decreasing factorially, we also standardise the signature terms so that it has mean 0 and variance 1 before regression.

Given that the signature method is largely agnostic to the observations used, we can also use it as a method to improve existing nowcasts. For example, we can take the mean estimated factors from the DFM and use these as “observations” in the signature method, we refer to these as “filtered DFM factors”. Theoretically, the signature method and the DFM should be comparable, since we have shown in the previous sections how we can replicate the performance of the Kalman filter by using a regression on the signature.

For each time point where new information arrives (i.e. weekly), we use information from previous quarters to fit the model and then apply the model to the current quarter. We use data from 1st Jan 2000 to 31st December 2015 as the training set, which means that they are always included in the dataset to fit the parameters for regression. From 2016 onwards, each week, new data is incorporated into the model (and new factors/principal components are computed) and another nowcast prediction is made for the GDP in the current quarter. The period from 1st January 2016 to 31st December 2017 is used as a validation set to select hyperparameters. The period 1st January 2018 to 31st December 2019 is then used as a “test set” with fixed hyperparameters, but the regression parameters are still tuned at each quarter.

6.3 Results

After a hyperparameter search, we fix the hyperparameters to be a Ridge regression with regularisation strength $\alpha = 2.0$ over a 730-day rolling window,

and we fit an intercept. The full details of the chosen hyperparameters can be found in Appendix D.1.

The results on the validation period can be seen in Figure 9a. From the plot we can see that the signatures returned a more stable prediction compared with the DFM, and attained the lowest root mean square error in this case. The RMSE for signatures, DFM, and NY Fed Staff forecast are 0.57, 0.85, and 0.86 respectively. Over the test period, the RMSE becomes 0.92, 0.98, and 1.03 for signature, DFM, and NYFed respectively. The nowcasts can be seen in Figure 9b.

We also fit a signature regression model on the filtered DFM factors. The results on the validation period can be seen in Figure 10a. From the plot we can see a better fit here compared with the principal components. The root mean square error is also lower in this case at 0.46. Over the test period, the RMSE becomes 0.82. See Figure 10b for the nowcasts. We note that using these mean estimates from the DFM model results in a noticeable improvement in prediction quality (over using the estimates in DFM directly).

	sig (PCA)	sig (filtered)	DFM	NY Fed DFM
validation period	0.57	0.46	0.85	0.86
test period	0.92	0.82	0.98	1.03

Table 1: RMSE for the signature method on PCA and filtered factors, as well as our DFM and the NY Fed published values.

The results are summarised in Table 1. We see that the test period seems to be a more volatile period, resulting in all models giving higher errors. Using the signature method achieved the lowest RMSE values, with using the filtered factor values giving the best performance.

7 Analysing fuel prices at the pump

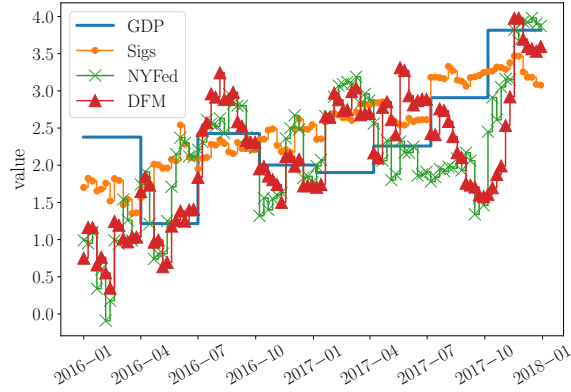
As a demonstration of the flexibility of the signature method onto other, more frequent time series, we give an application of nowcasting the weekly series of fuel prices (“at the pump”), based on the daily close price of an exchange-traded commodity (ETC) on Brent crude oil.

7.1 Data

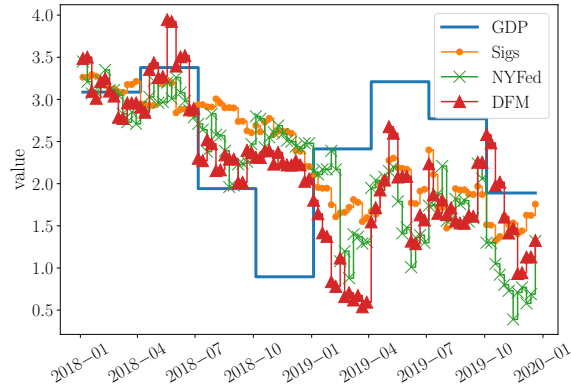
The target data available comes from the “[Weekly road fuel prices](#)” series published by the Department for Business, Energy & Industrial Strategy (BEIS).

To nowcast this series, we use the daily close price of WisdomTree Brent Crude Oil (BRNT), an ETC, at the London Stock Exchange. The data (in USD) is publicly available on [Yahoo](#), and we then use a [daily USD-GBP exchange rate](#) to convert the price of BRNT to GBP. The data is visualized in Figure 11. Although the frequency of the BRNT data series we are using is daily, since no trades occur on the weekend, there is missing data. Hence we have an irregular indicator series.

We note that although the frequency of the data is different to the applications in previous sections, after amending the configuration file with the new



(a) Validation period. The RMSE are sig (0.57), DFM (0.85), NY Fed Staff (0.86).



(b) Test period. The RMSE are sig (0.92), DFM (0.98), NY Fed Staff (1.03).

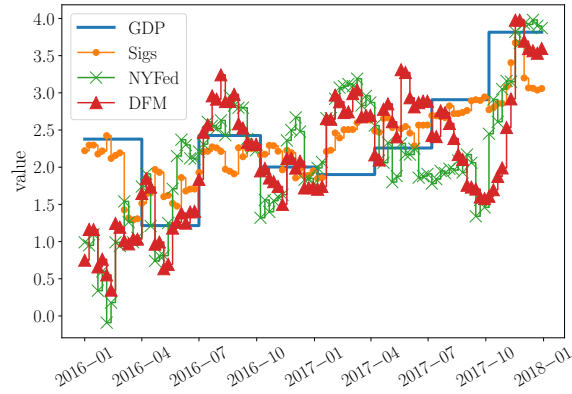
Figure 9: US GDP results when the signature method is applied to the principal components of the factor groups. The plots show comparison of: the signature model, our fitted dynamic factor model, the NY Fed Staff forecast values and the latest released GDP values.

publication lag (of 8 days), no additional modifications have to be made to the methods to handle the different structure. The embedding in continuous time of our method simplifies the pipeline.

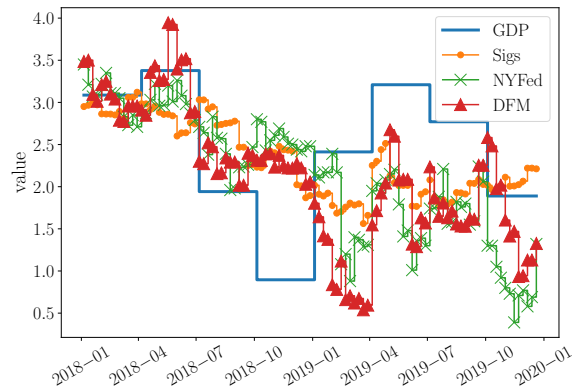
We nowcast the change in weekly fuel prices, and split our data using a train/validation/test proportion of 0.64/0.16/0.20 respectively. The target series is seen in Figure 12.

7.2 Results

In Figure 13, we see the nowcast of the signature method on the change in weekly fuel prices when the model is provided with previous weekly fuel prices and the BRNT price for the out-of-sample test data. The RMSE in this case is 1.132. The full details of the hyperparameters and the settings used can be



(a) Validation period. The RMSE are sig (0.46), DFM (0.85), NY Fed Staff (0.86).



(b) Test period. The RMSE are sig (0.82), DFM (0.98), NY Fed Staff (1.03).

Figure 10: US GDP results when the signature method is applied to the filtered means of the DFM. The plots show comparison of: the signature model, our fitted dynamic factor model, the NY Fed Staff forecast values and the latest released GDP values.

found in Appendix D.2.

In addition, we also look at two baselines. The first is a simple regression on the previous value (corresponding to an AR(1) model with an intercept). The second comes from using the package `pmdarima` to automatically search and select amongst ARIMA models. Since these models require the current value of the target to predict the next value, this means that the models output a new nowcast only when new data arrive on Tuesday of each week. For a fair comparison to signatures which produce a nowcast each day, we take the weekly nowcast produced by the baselines and compute daily errors.

The results can be seen in Figure 14. The AR(1) model has a RMSE of 1.237. The autoARIMA model selected to minimise the Akaike information criterion (AIC) is ARIMA(3,0,2), which has an RMSE of 1.243. The residuals for all three methods are seen in Figure 15. All results are given in Table 2.

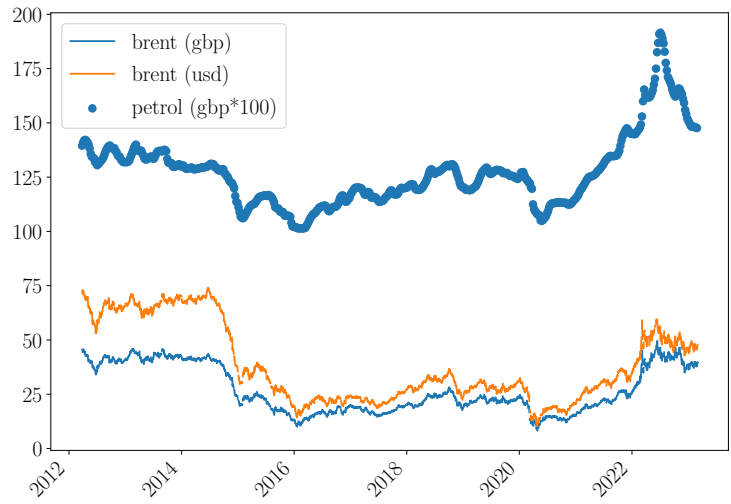


Figure 11: Time series of the price of BRNT before and after currency conversion along with the weekly fuel prices (in pennies) from BEIS.

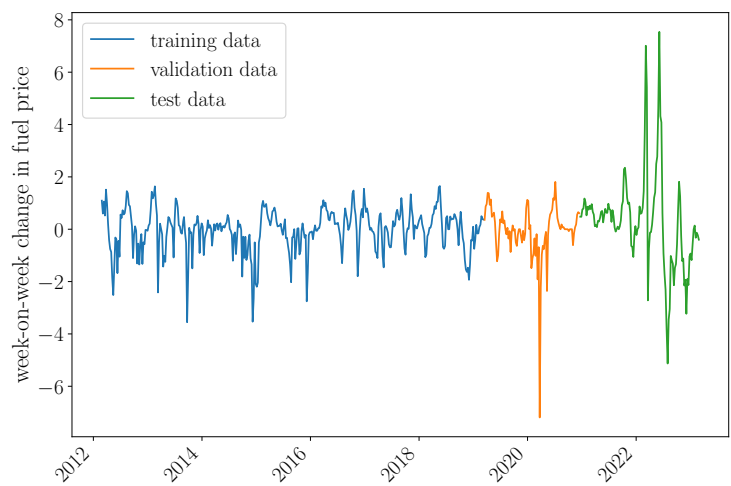


Figure 12: Target series: weekly change in fuel prices.

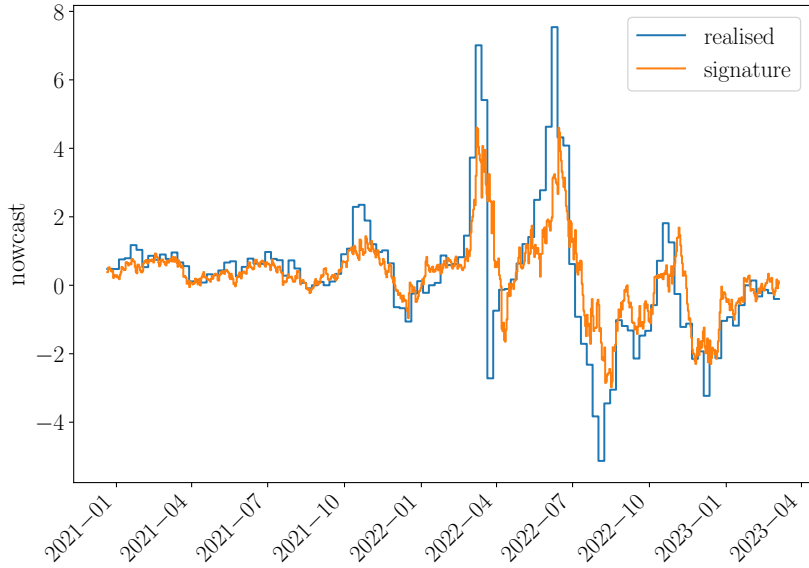


Figure 13: Nowcasts of incremental weekly fuel prices based on daily Brent ETC (BRNT) prices.

Note that the model selected by autoARIMA has slightly higher RMSE than an AR(1) with intercept since the results reported are out-of-sample, and therefore autoARIMA may have overfit to capture relationships specific to the training set. Therefore regression on signatures obtained the lowest RMSE.

	sig	AR(1)	autoARIMA
test period	1.132	1.237	1.243

Table 2: RMSE for the signature method, AR(1), and autoARIMA models for the fuel application.

In Figure 16, we see mean absolute error of the nowcast against the number of days it has been since the last release of the target variable. The nowcast error decreases sharply when a new weekly value is observed, indicating a fairly strong autoregressive relationship. As the week goes by the error increases slightly. This may indicate that the information in the daily Brent price series has limited use when nowcasting the weekly fuel price, perhaps due to the lag in effects of raw crude price on pump prices. Error increases on the weekend as time is elapsing and there is no information in the indicator series (as no trade is occurring on the exchange).

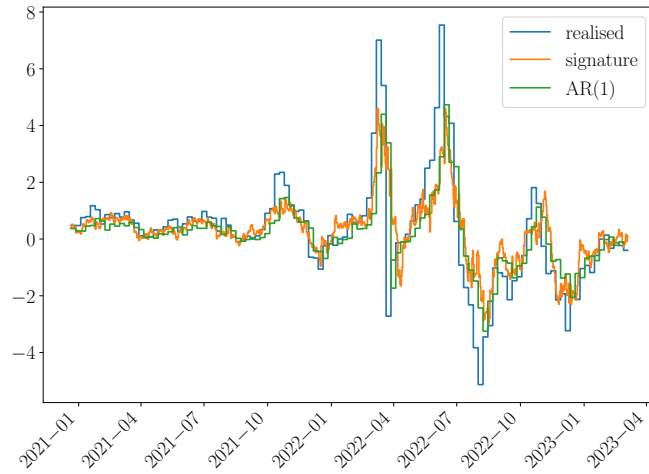
8 Conclusion

To summarise, the path signature, captures relevant geometric properties of sequential data. Signature methods naturally allow for missing data from mixed frequency and irregular sampling, issues often encountered in nowcasting, by embedding the observed data in continuous time.

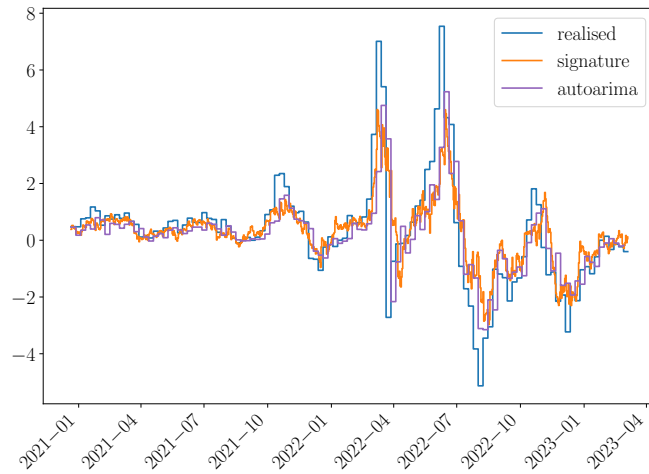
In this paper we demonstrated the application of regression on signatures to nowcasting. First, we introduced the theory of path signatures and illustrated how they are computed. We then showed that regression in the signature space subsumes the linear Kalman filter, which is commonly used in the nowcasting literature. We detailed a methodology for estimating nowcasts via regression on signatures and implemented this in the package [SigNow](#).

Second, we demonstrated that how it is possible to almost replicate the optimal performance of the Kalman filter (when the parameters are known) by using regression on signatures (fitting parameters based on the data) for a simulated experiment.

Finally we applied the model proposed in this paper to a couple of real-world data applications. The first is a well-known empirical exercise: nowcast GDP growth for the US. We have shown that the results obtained can offer improvements compared with a dynamic factor model based on the one published in Bok et al. (2018). We then demonstrated that we can take our methodology to other applications and observation frequencies, by illustrating its performance on inferring fuel prices. We stress that no additional changes are required in this application — embedding the problem to continuous time allows mixed and irregularly sampled problems to be solved by the same method.



(a) Nowcasts of incremental weekly fuel prices using AR(1).



(b) Nowcasts of incremental weekly fuel prices using autoarima.

Figure 14: Baseline models for fuel application. The AR(1) model with intercept has a RMSE of 1.237 and ARIMA(3,0,2), chosen by minimising AIC with autoARIMA, has an RMSE of 1.243.

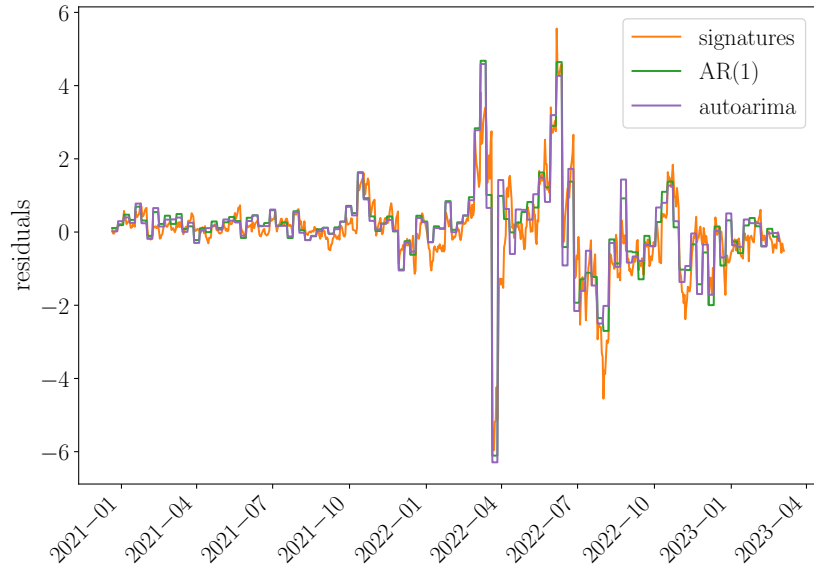


Figure 15: Residuals on the fuel application for signatures

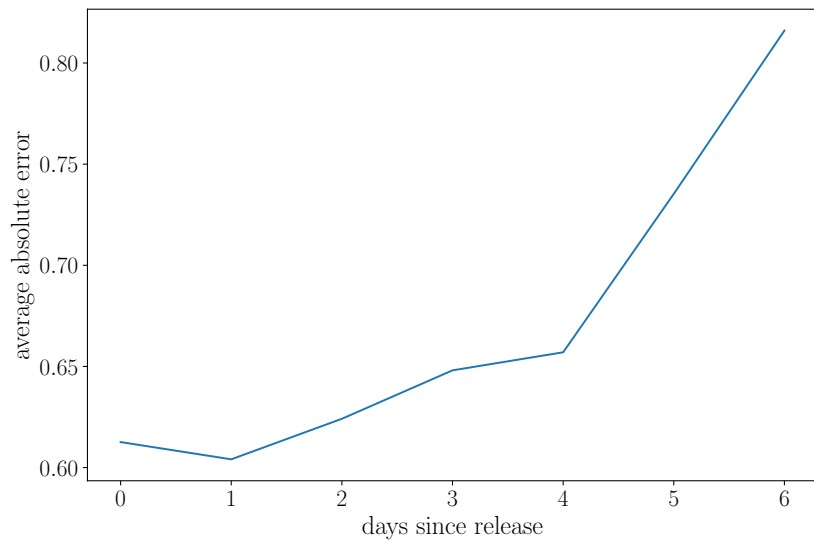


Figure 16: Nowcast error of signature method for weekly fuel prices based on how long it has been since the last release of the target variable.

A Comparing the regression coefficients

A.1 Regularly sampled data

For this exercise, we want to investigate the coefficients obtained from regression on signatures. We refer to the coefficients of the linear model from regression as the “regression coefficients” and the coefficients derived in Example 1 as the “true coefficients”. When we compare the regression coefficients against the true coefficients, there are large differences in value as can be seen in Figure 17.

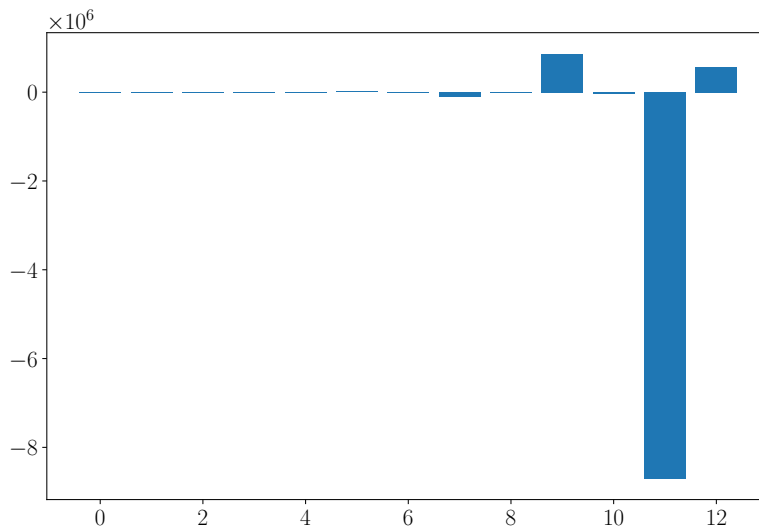


Figure 17: Difference between the theoretical and regression coefficients on linear signature terms up to level 6.

Upon closer inspection, the differences between the learnt regression coefficients against the true coefficients are greatest for those signatures which are composed of iterated integrals with respect to time only, i.e. those terms of the form $\frac{t^k}{k!}$. We note that, because we are working with fairly small values for the end time T , the higher order signature terms for t will be very small. In addition, these terms are clearly not algebraically independent, and this correlation may cause some degree of colinearity.

Therefore, we reduce the depth of the signatures that we take for the t terms to only keep 1 term in t , we also truncate the linear signature terms to 3. The fit of the linear models is still good, with the signatures residuals having a mean of -0.003 and a variance of 0.15 . The line of best fit (when plotted against the same Kalman filter residuals as in Figure 3) has gradient 0.97 and y -intercept of -0.04 , and an R^2 value of 0.86 . Figure 18 shows the difference between the residuals in this case, the coefficients on similar orders now.

We note that the true coefficients are correct for the infinite expansion of all signature terms of the filter. However, for a finite truncation level, we may be seeing alternative coefficients which are better fit for the data. If we take higher truncation level for the signature terms, for example up to level 6 and compute the singular values of the data matrix (of signatures), we find that at least 1 value is negligible. This means that the signature data matrix has a

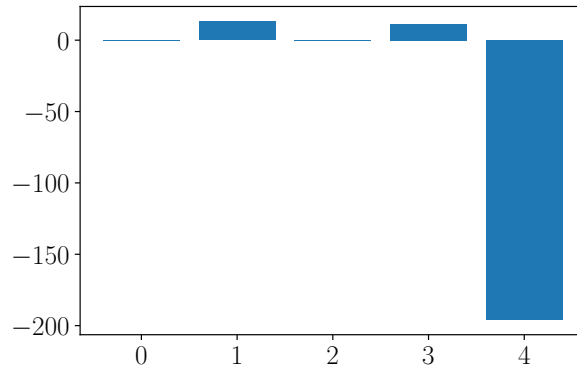


Figure 18: Difference between the theoretical and modified-level coefficients (only level 1 for t and level 3 for observed variables).

lower effective rank. This may also contribute to explaining why the regression coefficients are different even though the comparison against the Kalman filter appears almost perfect.

B Configurations for nowcasting pipeline

A short description of the main configuration settings for our signature nowcast can be found in Table 3.

In addition, the pipeline includes the possibilities to use custom factor structures to find principal components and much more. See the implementation of the experiments in this paper from our [Github repository](#) or refer to the documentation of our package [SigNow](#) for more details.

C Factor structure for the US GDP DFM

As in Bok et al. (2018), we assume that there are 4 factors: soft, real, labour, and global which affect the time series behind the US GDP nowcasting application. In this appendix, we list the groupings. The final list below, “other”, lists all the variables which do not fall under soft, real, and labour, but are also included in the analysis and are affected by the global factor alone.

Soft = {Empire state Mfg. Survey: General business conditions, Philly Fed Mfg. business outlook: Current activity, University of Michigan: Consumer Sentiment}

Real = {Real domestic gross product, Manufacturers new orders: Durable goods, Retail sales and food services, New single family houses sold, Housing starts, Industrial production index, Merchant wholesalers: Inventories: Total, Value of construction put in place, Building permits, Capacity utilization, Inventories: Total business, Real personal consumption expenditures, Manufacturing shipments: Durable goods, Mfrs. unfilled orders: All manufacturing industries, Manufacturing inventories: Durable goods, Real gross domestic income, Real

variable name	data type	description
window_type	str	The type of window used for generating the signatures - possible values are 'days', 'ind' and None. Arguments 'days' and 'ind' behave as moving windows. None acts as a expanding window.
max_length	int	When 'window_type' is set to 'days' or 'ind', 'max_length' is the number of days or rows to index by.
fill_method	str	How missing data should be filled - typical values 'ffill', 'bfill' or 'rectilinear'.
level	int	Truncation level for the variables.
t_level	int	Truncation level for the time parameter.
basepoint	bool	Whether to add a basepoint of zeros before computing signatures to remove lateral invariance.
use_multiplier	bool	Whether to multiply only the signatures corresponding to time only by multiplier based on target series.
keep_sigs	str	The type of signature terms to keep - values 'all', 'innermost', 'all_linear' and None.
regularize	str	Method to regularize, accepted values are 'elastic_net', 'l1', 'l2', and None. Default: 'elastic_net'.
alpha	float	Determines the strength of regularisation, see sklearn .
l1_ratio	float	Weight for the L1 penalty in the Elastic-Net of sklearn .
fit_intercept	bool	Whether to fit an intercept in regression.
target_lag	int	The target variable will have a lag due to the release timetable, this variable creates a shift so models never utilise future information.
use_prev_value	bool	Whether to use the latest value of the target series for the regression.
standardize	bool	Standardize signature terms before regression. Default: True.
training_proportion	float	This is an optional parameter to specify a fixed proportion for the train set.
reduce_dim	bool	Whether to reduce the dimension of the observed variables with principal component analysis.
k	int	The number of principal components to use.

Table 3: Description of the main configs used in the nowcasting pipeline.

disposable personal income, Exports: Goods and services, Imports: Goods and services}

Labour = {All employees: Total nonfarm, Civilian Unemployment rate, ADP nonfarm private payroll employment, Nonfarm business sector: Unit labor cost, JOLTS: Job openings: Total}

Other = {CPI-U: All items, PPI: Final demand, Import price index, PCE less food and energy: Chain price index, CPI-U: All items less food and energy, PCE: Chain price index, Export price index}

D Experimental Details

D.1 US GDP

For the US GDP growth example, there are two models, depending on whether the data used are the principal components of each factor group, or whether they are the final hidden factors from the dynamic factor model.

The target variable has a 30 day publication lag after the end of the quarter. This corresponds to a lag of about 124 days from the start of the reference quarter. All signature terms are standardised as discussed in Section 4.1.

For the data obtained using PCA, the final fitted model used all the linear signature terms truncated at level 3 over a lookback window of 730 days. This is a ridge model with a regularisation parameter strength of 2.0. We use the previous value of the target in the regression and allow an intercept. We use the target series as a multiplier but we do not use a basepoint when computing the signatures. Missing data is filled by forward fill. A summary of these hyperparameters can be found in Table 4.

Parameter	Value
level	3
t_level	3
fill_method	ffill
max_length	730
keep_sigs	all_linear
regularize	elasticnet
alpha	2.0
l1_ratio	0
standardize	True
fit_intercept	True
use_multiplier	True
basepoint	False
target_lag	124
use_prev_value	True

Table 4: Table of hyperparameters used to obtain the US GDP results with PCA “factors”.

For the data obtained using DFM means as observations, the final fitted model again used all the linear signature terms truncated at level 3 over a look-

back window of 730 days. This is a ridge model with a regularisation parameter strength of 1.0. We use the previous value of the target in the regression and allow an intercept. We use the target series as a multiplier but we do not use a basepoint when computing the signatures. Missing data is filled by rectilinear interpolation. A summary of these hyperparameters can be found in Table 5.

Parameter	Value
level	3
t_level	3
fill_method	rectilinear
max_length	730
keep_sigs	all_linear
regularize	elasticnet
alpha	1.0
l1_ratio	0
standardize	True
fit_intercept	True
use_multiplier	True
basepoint	False
target_lag	124
use_prev_value	True

Table 5: Table of hyperparameters used to obtain the US GDP results with filtered DFM factors.

D.2 Fuel analysis

For the fuel analysis the data for fuel price on a particular Monday is published the following day on Tuesday. Since we nowcast the next weekly change in price, this means that there is an 8-day delay in the target variable. As per discussion in Section 4.1, all signature terms are standardised.

The final fitted model used all the linear signature terms truncated at level 4 aside from the terms in time (which were truncated at level 3) over a lookback window of 17 days. This is an ridge model with a regularisation parameter strength of 0.5. We use the previous value of the target in the regression and allow an intercept. We use the target series as a multiplier but we do not use a basepoint. Missing data is filled by rectilinear interpolation. A summary of these hyperparameters can be found in Table 4.

Parameter	Value
training_proportion	0.80
level	4
t_level	3
max_length	17
fill_method	rectilinear
keep_sigs	all_linear
regularize	elasticnet
alpha	0.5
l1_ratio	0
reduce_dim	False
standardize	True
fit_intercept	True
use_multiplier	True
basepoint	False
target_lag	8
use_prev_value	True

Table 6: Table of hyperparameters used to obtain the fuel analysis results.

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