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Shapley regressions: a framework for statistical inference on machine learning models

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Abstract

Machine learning models often excel in the accuracy of their predictions but are opaque due to their non-linear and non-parametric structure. This makes statistical inference challenging and disqualifies them from many applications where model interpretability is crucial. This paper proposes the Shapley regression framework as an approach for statistical inference on non-linear or non-parametric models. Inference is performed based on the Shapley value decomposition of a model, a pay-off concept from cooperative game theory. I show that universal approximators from machine learning are estimation consistent and introduce hypothesis tests for individual variable contributions, model bias and parametric functional forms. The inference properties of state-of-the-art machine learning models — like artificial neural networks, support vector machines and random forests — are investigated using numerical simulations and real-world data. The proposed framework is unique in the sense that it is identical to the conventional case of statistical inference on a linear model if the model is linear in parameters. This makes it a well-motivated extension to more general models and strengthens the case for the use of machine learning to inform decisions.

Key words: Machine learning, statistical inference, Shapley values, numerical simulations, macroeconomics, time series.

JEL classification: C45, C52, C71, E47.

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1 Introduction

Model families from machine learning, like support vector machines, tree ensembles and artificial neural networks, often excel in the accuracy of their predictions (Fernandez-Delgado (2014)) but are opaque due to their complex structure. More generally, many models make a trade-off between simplicity and accuracy. Accuracy provides confidence that a model’s predictions are close to actual outcomes, while simplicity facilitates understanding and communication. On a technical level, this usually boils down to a statistical inference analysis, e.g. the estimation of a coefficient associated with a variable in the model and its confidence levels with respect to a hypothesis (mostly the null). This approach is largely limited to linear parametric models or generalised linear models (Greene (2017)).

On the other hand, machine learning models are mostly non-parametric, built around producing accurate predictions (Friedman et al. (2009)). For example, artificial neural networks, which are driving current advances in artificial intelligence in the form of deep learning (Goodfellow et al. (2016)), have long been known to have universal approximator properties (Portnoy (1988)). They can approximate almost any unknown function given enough training data. However, this directly leads to the black-box critique of machine learning models, because it is not straightforward to understand a model’s input-output relations or perform a statistical inference analysis on them. This causes not only practical obstacles for their application, but also ethical and safety concerns more generally which are increasingly reflected in legal and regulatory frameworks (European Union (2016)).

Despite these important concerns, machine learning models could provide substantial benefits in the context of prediction policy problems (Kleinberg et al. (2015)). These are situations where the precise prediction of outcomes is important to inform decisions. Examples include the forecasting of economic developments (Garcia et al. (2017)), modelling the soundness of financial institutions (Chakraborty and Joseph (2017)), consumer credit scoring (Fuster et al. (2017)), policy targeting based on uncertain outcomes (Andini et al. (2017)), the prediction of extreme weather events in the face of climate change (Racah et al. (2016)), medical image analysis and diagnosis (Litjens et al. (2017)) or aiding expert judgement (Kleinberg et al. (2018)). Institutional transparency is an additional aspect from a public policy point of view, such as the decision processes of central banks, regulators and governments (Bernanke (2010)). On the one hand, decision makers need to understand the driving factors of the quantitative models

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1 There is also an active area of research into simple but accurate models, e.g. via the use of decision heuristics or fast-and-frugal trees (see for example Aikman et al. (2014); Simsek and Buckmann (2015)).

2 This property also applies to other non-parametric models often used in machine learning, see e.g. Scornet et al. (2014); Christmann and Steinwart (2008).

3 As soon as we need to consider the change in outcome due to any action taken as a response to a prediction, we enter the area of a causal inference or mixed policy problem.
they rely on, and, on the other hand, also be able to communicate them clearly. Again, the
opaqueness of machine learning models hinders their application with regard to both points.
Finally, the need for machine learning models is likely to be aggravated by the current prolif-
eration of large and granular data sources. For instance, data from social media, smart phone
usage, ubiquitous sensors or the ‘internet of things’ may allow for the modelling of human
behaviour or the dynamics of autonomous machines in complex environments on an unprece-
dented level. Such capabilities may provide large benefits for technological advancement or
societal development more generally. Again, a detailed understanding of the deployed models
will be needed to fully utilise this potential.

Two approaches to address the interpretability issue of machine learning models\textsuperscript{4} are variable
attributions via the decomposition of individual predictions (local attribution) and importance
scores for the model as a whole (global attribution). A well-motivated local decomposition
is provided by model Shapley values (\textcite{Strumbelj2010}; \textcite{Lundberg2017}), a pay-off concept from cooperative game theory (\textcite{Shapley1953}; \textcite{Young1985}). It
maps the marginal contribution coming from a variable within a set of variables to individual
model predictions. However, model decomposition is only one part of model interpretability.
An equally important part is statistical inference in the form of hypothesis testing to assess the
confidence we can have in specific model outputs.

This paper proposes a general statistical inference framework for non-parametric models based
on the Shapley decomposition of a model, namely Shapley regressions. This framework trans-
fers the model inference problem into a locally linear space. This simultaneously opens the
toolbox of econometrics, or parametric statistics more generally, to machine learning and vice
versa. Model inference consists of three steps. First, model calibration and fitting (training).
Second, model testing and Shapley value decomposition on a hold-out dataset. Finally, infer-
ence based on a surrogate regression analysis using the Shapley decomposition as its inputs. For
the known case of a linear model, this approach reduces to the standard least-squares case\textsuperscript{5} In
this sense, Shapley regressions can be seen as a natural extension of regression-based inference
to the general non-linear model. The main distinction is that inference is often only valid on
a local level, i.e. within a region of the input space due to the potential non-linearity of the
model plane. A consequence of this is that the concept of a regression coefficient as a standard
way of measuring and communicating effects is not directly applicable. I propose a generalised
coefficient concept suited for the non-linear case which is close to its linear parent. It allows
for similar assessment and communication of modelling results. On a deeper level, the current

\textsuperscript{4}I only discuss supervised learning in this paper. However, the proposed methodology can be applied more
generally in situations where a model delivers a score which needs to be evaluated based on its inputs.

\textsuperscript{5}Shapley values have been used in linear regression analysis before to address collinearity issues (\textcite{Lipovetsky2001}). I do not see scope for confusion with the current application.
work builds on seminal work in non-parametric statistics (Stone (1977)) and connects it with recent developments in the machine learning literature.

The remainder of this paper is structured as follows. Section 2 discusses model interpretability more widely and the nascent literature on statistical inference using machine learning models in econometrics. Section 3 introduces the concept of Shapley values and Shapley regressions for model inference. A slightly modified null hypothesis is introduced to test the statistical significance of variables in a model. Shapley share coefficients are defined as a summarising concept to assess individual variable contributions akin to linear regression coefficients. In Section 4, the theoretical estimator properties of machine learning model are investigated. General estimation consistency is shown for the large class of piecewise analytic functions. I present a test to assess model bias for more general model decompositions which are based on Shapley values and introduce robust component estimates. The validity conditions for inference within the Shapley regression framework are stated. Particularly, valid asymptotic inference depends on sample splitting for model training and testing which is a common procedure when building a machine learning system. Section 5 considers applications. First, empirical estimation properties of commonly used machine learning models - like artificial neural networks (NN), support vector machines (SVM) and random forests (RF) - are investigated using numerical simulations. Second, the Shapley regression framework is applied to modelling long-run macroeconomic time series for the UK and US. Machine learning models are mostly more accurate than either regularised (biased) and unbiased linear benchmark models. Inference from the Shapley regression framework is robust against model choice and richer than that of benchmark models pointing to the importance of non-linearities in modelling these data generating processes. Differences in results are in line with analytical model properties and can be used for model selection. The main drawback of using the Shapley regressions framework is the computational cost of calculating Shapley value decompositions. Depending on the application, this can be addressed via appropriate approximations or sampling procedures. Section 6 concludes.

An inference recipe for machine learning models is summarised in Box 1 in the Appendix together with figures, tables and proofs of theoretical results. The code and data for the numerical and empirical analyses alongside supplementary results are available on Github.com/Bank-of-England/Shapley_regressions.

2 Literature

Approaches to interpretable machine learning come from different directions: General issues around model interpretability, technical approaches from within machine learning research and approaches from econometrics and statistics. I will primarily focus on the latter two.
The highest level of discussion relates to reasons why models should be interpretable and well-communicated, despite good comparative numerical performance. Especially in the context of informing decisions, these are intertwined ethical, safety, privacy and increasingly legal concerns about the application of opaque models (Crawford (2013); European Union (2016); Fuster et al. (2017)). Lipton (2016) discusses desirable properties of interpretable research in general (trust, causality, transferability, informativeness) and models we use (transparency, e.g. via local decomposability) and interpretability (e.g. via visualisations and relatedness). He argues that a complex machine learning model does not need to be less interpretable than a simpler linear model if the latter operates on a more complex space. This is in line with Miller (2017), who provides a comprehensive discussion of explainable artificial intelligence (often referred to as XAI) from a social science perspective. One take-away message is that humans prefer simple explanations, i.e. those citing fewer causes and explaining more general events, are generally preferred, though they may be biased. Shallow tree models from machine learning, or derived fast-and-frugal-trees, may thus offer accurate models while also providing satisfactory transparency (Aikman et al. (2014); Simsek and Buckmann (2015)).

Approaches in computer science have focused on model decomposition by means of variables attribution techniques. That is, scores of importance are given to each input variable for single observations or the full model. Gini importance for tree-based classifiers is an example of a model score. It is a measure for how much a variable contributes to the optimisation of the objective function (Kazemitabar et al. (2017); Friedman et al. (2009)). Local attributions decompose individual predictions assigning scores to each input variable. Here, one approach is to construct approximate surrogate models which allow for model decomposition. Examples are LIME (Ribeiro et al. (2016)), DeepLIFT (Shrikumar et al. (2017)) and Shapley values (Strumbelj and Kononenko (2010)). Lundberg and Lee (2017) demonstrate that Shapley values offer a unified framework of previous attribution schemes with appealing properties. These are also the reason for their use in the current paper.

The literature of inference using machine learning models from an econometrics point of view is just at its beginning and also the main area this paper talks to. I distinguish three approaches. First, one can construct a correspondence between an econometric and a machine learning model where possible. Mullainathan and Spiess (2017) present the simple but intriguing idea to treat a not-too-deep tree model as a regression model with multiple interaction terms, one per leaf node. Similar to the tree model, overfitting is an emerging issue. This can be addressed via regularisation, and the estimation of unbiased coefficient on the regularised model corresponding to a pruned tree when shrinking coefficients to zero.

6Local Interpretable Model-agnostic Explanations.
7Deep Learning Important FeaTures for NN.
The second approach is double or debiased machine learning (Chernozhukov et al. (2018)). It deals with the issue of parameter regularisation bias using machine learning, e.g. when estimating a partially linear model in the presence of a high-dimensional nuisance parameter. This bias is avoided via the construction of orthogonal score functions for the estimation of a low-dimensional target parameter. The procedure is model independent and allows for the well-defined inference on causal parameters. The main difference to the current paper is that I do not allow parameters of interest to be part of the model optimisation stage but rather recover those from an a posteriori decomposition which may involve a particular parametric form or not.

A third approach has been to use a priori modified models which have well-defined statistical properties, e.g. for the estimation of treatment effects. Wager and Athey (2018) introduce a type of RF for the estimation of heterogeneous treatment effects. The idea is based on the notion that small enough leaf nodes provide uncorrelated sub-samples as though they had come from a randomised experiment. Intuitively, trees in a forest act as a form of matching algorithm which is more flexible than conventional techniques due to the adaptive nature of tree models. For the construction of these causal forests, they introduce the concept of honest trees as a modification of the original algorithm. These now have an asymptotically Gaussian and centred sampling distribution. The idea of using specific characteristics of machine learning models to improve on existing techniques is again intriguing. The present paper is complimentary to this approach. RF from honest trees are still open to the black-box critique, which can be addressed by the Shapley regression framework.\footnote{The same applies with respect to Chernozhukov et al. (2018), meaning the current paper is not a substitute but a complement to preceding work.}

3 The Shapley regression framework

3.1 Notation and definitions

This paper considers the common case where \( f(x; \beta) : D \subset \mathbb{R}^m \rightarrow \mathbb{R}^p \) is the data generating process (DGP) of interest with domain \( D \). We only consider the case \( p = 1 \) (the extension to \( p > 1 \) is straightforward). The data \( x \in \mathbb{R}^{n \times m} \) with \( m \) being the number of features or independent variables and \( n \) the number of observations. Features are assumed to be independent from each other, while observations need not be (column-wise independence). Consequences of this restriction and ways to address it, if too stringent, will be discussed.

The vector \( \beta \in \mathbb{R}^{m+1} \) describes the parameterisation of the DGP, such as the set of coefficients of a linear model with \( \beta_0 \) being the intercept. The parameters \( \beta \) represent the effects we are interested in studying. The DGP \( f \) is assumed to be piecewise continuous and differentiable.
on finite sub-domains of \( D \) and to have finite moments, i.e. \( \mathbb{E}_D[f^d] < \infty \), with \( d \geq 1 \). Regions within \( D \) are labelled \( \Omega \subset D \).

The non-parametric model is \( \hat{f}(x; \theta) : D \subset \mathbb{R}^m \mapsto \mathbb{R}^q \) with \( \theta \in \mathbb{R}^q \) where \( q \to \infty \) as \( m \to \infty \) is allowed. It represents our machine learning models of interest, such as NN, SVM or RF. In these cases, \( \theta \) represents the network weights, support vector coefficients and split points, respectively. The model parameters \( \theta \) are slightly different to their usage in semi-parametric statistics, where \( \theta \) often describes a high-dimensional nuisance parameter, which may be present or not. The model \( \hat{f} \) is assumed to have finite moments but no other regularity conditions are imposed. The linear model is parameterised by \( \hat{\beta} \).

The used index convention is that \( i, j \in \{1, \ldots, n\} \) refer to individual observations and \( k, l \in \{1, \ldots, m\} \) to feature dimensions. No index refers to the whole dataset \( x \in \mathbb{R}^{n \times m} \). An index \( c \in \{1, \ldots, C\} \) refers to components of linear decompositions of either a DGP or a model, e.g. \( f = \sum_{c=1}^{C} \psi_c \equiv \Psi \). \( \Phi^S \) refers to the Shapley decomposition of a model (see below). Superscripts \( S \) refer to “Shapley-related” quantities which will be clear from the context. Estimated quantities are hatted, except \( \Phi/\phi \) for simplicity.

### 3.2 The linear model as a guiding principle

Statistical inference can be local or global. The linear model \( \hat{f}(x_i) = x_i \hat{\beta} = \sum_{k=0}^{m} x_{ik} \hat{\beta}_k \) is special in the sense that it provides local and global inference at the same time. The coefficients \( \hat{\beta} \) describe local effects via the sum of the product of variable components and coefficients. At the same time, the coefficient vector \( \hat{\beta} \) determines the orientation of the global model plane with constant slope in each direction of the input space. As long as the number of co-variables in a model is modest, the linear model is widely accepted to provide good inference properties and is the workhorse of econometric analysis.

The linear model belongs to the class of additive variable attributions. For an observation \( x_i \in \mathbb{R}^m \) we define the model decomposition \( \Phi \) as

\[
\Phi(\hat{f}(x_i)) \equiv \phi_0 + \sum_{k=1}^{m} \phi_k(x_i) \overset{\text{lin.model}}{=} \hat{\beta}_0 + \sum_{k=1}^{m} x_{ik} \hat{\beta}_k, \tag{1}
\]

where \( \phi_0 = \hat{\beta}_0 \) is the intercept. The standard approach to test for the importance of a certain variable is to test against the null hypothesis \( \mathcal{H}_k^0 : \{\beta_k = 0\} \). The goal of this paper is to arrive at a similar hypothesis test valid for more general models \( \hat{f} \).

### 3.3 Shapley values

A more general class of additive attribution is given by model Shapley values \( \Phi^S \), a pay-off concept from cooperative game theory (Shapley (1953)). Making the analogy between players
of a multi-player game cooperating to generate a pay-off and variables \( x_k \) within a model to generate predictions \( \hat{f}(x) \), the marginal contribution from variable \( k \) is defined in the form of its Shapley value (Strumbelj and Kononenko (2010))

\[
\phi_k^S(\hat{f}, x) = \sum_{x' \subseteq C(x) \setminus \{k\}} \frac{|x'|!(n - |x'|-1)!}{n!} \left[ \hat{f}(x' \cup \{k\}) - \hat{f}(x') \right],
\]

where \( C(x) \setminus \{k\} \) is the set of all possible coalitions of \( m - 1 \) model variables when excluding the \( k^{th} \) variable. \(|x'|\) denotes the number of included variables. Eq. 2 is the weighted sum of marginal contributions of variable \( k \) accounting for the number of possible coalitions for a certain \( x' \).

Intuitively, the above definition of a Shapley value is similar to the regression anatomy of a coefficient \( \hat{\beta}_k \), i.e. the bivariate slope coefficient after partialling out all other regressors in a multi-variate model (Angrist and Pischke (2008)). This will be formalised below.

Shapley values are the unique class of additive value attribution with the following properties (Shapley (1953); Young (1985); Strumbelj and Kononenko (2010)).

**Property 1: Efficiency.** The attribution model \( \Phi^S \) matches the original model \( \hat{f} \) at \( x_i \),

\[
\Phi^S(x_i) \equiv \phi_0^S + \sum_{k=1}^{m} \phi_k^S(x_i) = \hat{f}(x_i).
\]

In a modelling context, this property is called *local accuracy*. A model’s Shapley decomposition always sums to the predicted value. The intercept \( \phi_0 \) is the expected or average model value.

**Property 2: Missingness (null player).** If a variable is missing from a model, no attribution is given to it, i.e. \( \phi_k^S = 0 \) (dummy player).

**Property 3: Symmetry.** If \( k \) and \( k' \) are two variables which are equivalent, such that

\[
\hat{f}(x' \cup \{j\}) = \hat{f}(x' \cup \{k\})
\]

for all possible \( x' \) not containing \( j \) or \( k \), then \( \phi_j^S = \phi_k^S \).

**Property 4: Strong monotonicity.** Variable attributions do not decrease if an input’s contribution to a model increases or stays the same regardless of other variables in the model. That is, for any two models \( \hat{f} \) and \( \hat{f}' \), if

\[
\hat{f}(x') - \hat{f}(x' \setminus k) \geq \hat{f}'(x') - \hat{f}'(x' \setminus k)
\]

for all possible \( x' \), then \( \phi_k^S(f, x) \geq \phi_k^S(f', x) \). \( x' \setminus k \) indicates the set of variables excluding \( k \).

In the context of variable attribution, this property is also called *attribution consistency*. It is

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Footnote 9: For example, assuming we have three players (variables) \( \{A, B, C\} \), the Shapley value of player \( C \) would be

\[
\phi_C^S(\hat{f}) = 1/3[\hat{f}(\{A, B, C\}) - \hat{f}(\{A, B\})] + 1/6[\hat{f}(\{A, C\}) - \hat{f}(\{A\})] + 1/6[\hat{f}(\{B, C\}) - \hat{f}(\{B\})] + 1/3[\hat{f}(\{C\}) - \hat{f}(\{\emptyset\})].
\]
an innovation to previous approaches, such as Gini importance of decision trees (Lundberg et al. 2018).

**Property 5: Linearity.** For any two independent models \( \hat{f} \) and \( \hat{f}' \), i.e. where the outcome of the one does not depend on the inputs or outcome of the other, the joint Shapley decomposition for a variable \( k \) can be written as

\[
\phi^S_k (a(\hat{f} + \hat{f}')) = a\phi^S_k (\hat{f}) + a\phi^S_k (\hat{f}')
\]

for any real number \( a \). A consequence of these properties is the following proposition.

**Proposition 3.1.** The Shapley decomposition \( \Phi^S \) of a model \( \hat{f} \) linear in parameters \( \hat{\beta} \), \( \hat{f}(x) = x\hat{\beta} \), is the model itself. The proof is given in the Appendix.

Hence, the Shapley value decomposition of the linear model is well known.

Regarding the computation of model Shapley values (2), most models cannot handle missing variables to evaluate “variable coalitions”. If missing from a coalition, the contribution of a variable is integrated out via conditional expectations relative to a representative background sample. Particularly, we evaluate \( \mathbb{E}_{x \setminus \{C\}} [\hat{f}(x)|x_C] \), where \( C \) is the set of non-missing variables in a coalition. For this to be exact, one has to assume feature independence to avoid model evaluations at unreasonable inputs. This can be a strong assumption for many applications. I will demonstrate a way to quantify errors made based on this assumption in Section 5.2.4.

The computation of Shapley decompositions is challenging due to the exponential complexity of (2). Two approaches have been proposed in the machine learning literature which preserve the properties of Shapley values, Shapley sampling values (Strumbelj and Kononenko (2010)) and Shapley additive explanations (SHAP, Lundberg and Lee (2017)). The latter provides an improvement on the former and will be the basis for the calculation of Shapley decompositions in this paper. The background dataset is taken to be the training set of a model which contains the information the model parameters \( \hat{\theta} \) are based on from the optimisation process. The calculation of model Shapley values is probably the biggest drawback in their usage. Appropriate approximations or sampling procedures may be used and tested depending on the situation.

### 3.4 Shapley regressions

Having a well-defined measure for variable attributions, we next turn to hypothesis testing, e.g. to assess the significance of individual variable contributions. For this, one can reformulate an inference problem in terms of a model’s Shapley decomposition. That is, one estimates the

\[10\] This corresponds to linear Shap in Lundberg and Lee (2017).

\[11\] For high-dimensional data, such as images or text, it is often more practical and intuitive to work with lower dimensional representations, such as super-pixels/objects or topics, respectively.
Shapley regression

\[ y_i = \Phi_i^S \hat{\beta}_S^j + \hat{\epsilon}_i \equiv \sum_{k=0}^{m} \phi_k^S (\hat{f}, x_i) \hat{\beta}_k^S + \hat{\epsilon}_i , \quad (7) \]

where \( k = 0 \) corresponds to the intercept and \( \hat{\epsilon}_i \sim \mathcal{N}(0, \sigma^2) \). The surrogate coefficients \( \hat{\beta}_k^S \) are tested against the null hypothesis

\[ \mathcal{H}_0^k(\Omega) : \{ \beta_k^S \leq 0 \mid \Omega \} . \quad (8) \]

The key difference to the linear case is the regional dependence on \( \Omega \), i.e. only local statements about the significance of variable contributions can be made. This is related to the potential non-linearity of a model whose hyperplane in the input-target space may be curved compared to that of the linear model \([1]\).

The following proposition provides further justification for the use of Shapley regressions for inference on machine learning models.

**Proposition 3.2.** The Shapley regression problem of Eq. \([2]\) for a model \( \hat{f} \) linear in parameters \( \hat{\beta} \) is identical to the least-square problem related to \( \hat{f}(x) = x \hat{\beta} \), i.e. \( \hat{\beta}_S^j = 1 \). The proof is given in the Appendix.

Proposition 3.2 provides practical guidelines and intuition regarding the the coefficients \( \hat{\beta}_k^S \). Geometrically, they describe the alignment of the model hyperplane spanned by the Shapley decomposition and the target variable, in the same way as the coefficients of a linear model in the original input space. Notionally this is not different from a variables transformation. One expects coefficient values of unity, i.e. \( \hat{\beta}_S^j = 1 \) if the machine learning model generalises well.\(^{12}\) Deviations from unity are caused by the best-fit hyperplane being tilted in certain directions and provide insight about the generalisation properties of the model. Values greater than unity indicate that \( \hat{f} \) underestimates the effect of a variable. Values smaller than one indicate the opposite. Particularly, statistical significance will drop as \( \hat{\beta}_k^S \) approaches zero as there is no clear alignment between Shapley components \( \phi_k^S \) and the target \( y \). We reject negative coefficients, as they are opposed to the alignments of attributed effects \( \phi_k^S \). These can occur when \( \hat{f} \) is not a good fit itself.

Having derived a test against the null hypothesis, it is not yet clear how to communicate inference results. The coefficients \( \hat{\beta}_S^j \) are only partially informative, as they to not quantify the components of \( \Phi^S \) but rather their alignment with the target independent of their actual magnitude. I propose the following generalised coefficient.

\(^{12}\)A formal derivation of this statement is given in the next section.
3.4.1 Shapley share coefficients

The Shapley share coefficients (SSC) of variable $x_k$ in the Shapley regression framework is defined as

$$\Gamma_{k}^{S}(\hat{f}, \Omega) \equiv \left[ \text{sign}(\hat{\beta}_k) \left\langle \frac{|\phi_k^S(\hat{f})|}{\sum_{l=1}^{n} |\phi_l^S(\hat{f})|} \right\rangle_{\Omega_k} \right]^{(*)} \in [-1, 1],$$

(9)

$$f(x)=x^\beta \hat{\beta}_k^{(*)} \left\langle \frac{|(x_k - \langle x_k \rangle)|}{\sum_{l=1}^{n} |\hat{\beta}_k(x_l - \langle x_l \rangle)|} \right\rangle_{\Omega_k},$$

(10)

where $\langle \cdot \rangle_{\Omega_k}$ stands for the average over $x_k$ in $\Omega_k \in \mathbb{R}$. The SSC $\Gamma_{k}^{S}(\hat{f}, \Omega)$ is a summary statistic for the contribution of $x_k$ to the model over a region $\Omega \subset \mathbb{R}^m$.

It consists of three parts. The first is the sign, which is the sign of the corresponding linear model. The motivation for this is to indicate alignment of a variable with the target. The second part is coefficient size. It is defined as the fraction of absolute variable attribution allotted to $x_k$ across the range of $x$ considered. The sum of absolute value of SSC is one by construction. It measures how much of the model output is explained by $x_k$. The last component $(*)$ is used to indicate the significance level of Shapley attributions from $x_k$ against the null hypothesis (8) and, thus, the confidence one can have in information derived from that variable.

Eq. (10) provides the explicit form for the linear model. The main difference to the conventional case is a normalising factor accounting for localised properties of non-linear models. Given the definition over a range $x_k$, it is important to also interpret them in this context. For example, contributions may vary over the input space such that $\hat{\beta}_k$ takes on different values at different points or times.

More generally, a coefficient is a constant factor multiplying some quantity of interest. This is a concept from linear models which does not directly translate to the non-linear case. Eq. (9) is constructed in such a way to provide comparable information and structure. A key property and further difference to the linear case of this generalisation is that (9) does not make assumptions about the functional form of the DGP, hence it may be called a “non-parametric coefficient”.

3.4.2 SSC standard errors

Given the conditions we required from $\hat{f}$, the classical central limit theorem applies to the sampling distribution of Shapley values $\phi_k^S(\hat{f})$, tending to a multivariate normal distribution. This can be used to construct standard errors and confidence intervals for $\mathbb{E}_\Omega[\phi_k^S]$. However, the information derived from this may be hard to interpret given the lack of a scale in components $\phi_k^S$. Not so for the SSC (9), which are normalised.

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The normalisation is not needed in binary classification problems where the model output is a probability. Here, the a Shapley contribution relative to a base rate can be interpreted as the expected change in probability due to that variable.
Let $\mu_k = |\Gamma^S_k(\hat{f}, \Omega)| \in [0,1]$ be the absolute value of the $k$-th SSC. The upper bounds on the variance of $\mu_k$ and its sampling standard error of the mean are given by\footnote{One will generally be interested in the expected explanatory fraction $\mu_k$ of a variable, while the sign of the SSC is fixed. Accounting for the sign, the bound on the RHS of (11) needs to be multiplied by four.}:

$$\text{var}(\mu_k) \leq \mu_k(1 - \mu_k) \leq \frac{1}{2} \implies \sigma_k^\phi \equiv \text{se}(\mu_k) \leq \frac{1}{\sqrt{2|\Omega|}}.$$

The sampling distribution of $\mu_k$ will also approach a Gaussian with increasing sample size $|\Omega|$. Thus, $\sigma^\phi$ provides a well-defined measure of the variability of $\Gamma^S$ within $\Omega$. We have now assembled tools for statistical inference on machine learning models regarding the direction, size, significance and variability of variable contributions. Next, I provide the theoretical underpinning of the proposed framework.

### 4 Machine learning estimator properties

Focusing on regression problems\footnote{A regression model in machine learning refers to fitting a continuous target or dependent variable. Problems which describe categorical variables, e.g. a binary target, are called classification problems. All results presented here can be applied in the classification case.} it is common to minimise the mean squared error (MSE) between a target $y$ from a DGP $f(\beta)$ and a model $\hat{f}(\theta)$ over a dataset $x$. The expected MSE can be decomposed as

$$\mathbb{E}_x[(y - \hat{f}(\theta))^2] = \left( f(\beta) - \mathbb{E}_x[\hat{f}(\theta)] \right)^2 + \left( \hat{f}(\theta) - \mathbb{E}_x[\hat{f}(\theta)] \right)^2 + \sigma^2,$$

where $\sigma^2$ is the irreducible error component of the DGP corresponding to the variance of $y$. Eq. (12) distinguishes between external model parameters $\theta$ and internal parameters $\beta$ of the DGP. This separation is important, because machine learning models are often subject to regularisation as part of cross-validation procedures (model calibration) and the training process. This directly affects the model parameters $\theta$ (if present) when minimising (12). Thus, if $\hat{\beta}$ would explicitly be part of the training process, its values would be biased as was investigated in Chernozhukov et al. (2018). It is at the heart of machine learning to generalise to $\beta$ from $(y, x)$ by the means of $\theta$. This generalisation can made explicit, i.e. by recovering $\beta$, using Shapley values and regressions.

#### 4.1 Estimator consistency

Statistical inference on machine learning models requires two steps. First, the control of bias and variance according to (12) and, second, the extraction of and inference on $\hat{\beta}$. Regarding the
former, most non-parametric estimators for regression problems are consistent in the sense that the squared error tends towards $\sigma^2$ as the training data size tends to infinity. This property can be called error consistency, that is

$$p \lim_{n \to \infty} |y - \hat{f}(\theta|x)| = 0,$$

i.e. that expected divergence of $\hat{f}$ from the true value $y$ converges towards zero in probability as the sample size increases (assuming $\sigma = 0$). Eq. 13 defines the universal approximator property of machine learning models. It is ultimately based on the consistency of non-parametric regressions according to Stone (1977)\textsuperscript{16} That is, universally consistent machine learning models can be interpreted as generating a local weight distributions which mimic the DGP. This does not necessarily imply estimator consistency\textsuperscript{17} i.e. if

$$p \lim_{n \to \infty} \beta_k - \hat{\beta}_k(\theta|x) = 0.$$

That is, if universal approximators learn the correct parameterisation of a DGP.

In many applications of interest, $f$ can be locally approximated by a polynomial regression. For a polynomial DGP the following result holds.

**Theorem 4.1.** (polynomial consistency of machine learning estimator): Let $f$ be a DGP of the form $f(\beta, x) = \sum_{k=0}^{m} \beta_k p_k^d(x) \equiv P_k(x)$, where $p^d(x)$ is a polynomial of order $d$ of the input features on a subspace $x \in \Omega$ and $\Omega \subseteq D \subseteq \mathbb{R}^m$. If for each $x' \subseteq \Omega$, a model $\hat{f}(\theta)$ is error consistent, then the estimator $\hat{\beta}(\theta)$ is also estimator consistent in the sense of (14) as long as $\hat{f}$ does not explicitly depend on $\beta$. The proof is given in the Appendix.

Theorem 4.1 can be used to make a more general statement about non-linear parameter dependencies.

**Corollary 1** (universal consistency): Let $f(\beta)$ be a DGP on $\Omega \subseteq D \subseteq \mathbb{R}^m$ and $\hat{f}(\theta)$ a model not involving $\beta$. If $f$ can be approximated by a polynomial $\hat{f}_p(\theta')$ arbitrarily close and $\hat{f}$ is error consistent, then $\hat{f}$ is estimator consistent for any $f(\beta)$. Particularly, the effect $\beta$ is locally approximated by $\hat{f}_p(\theta')$ arbitrarily precise. The proof is given in the Appendix.

Corollary 1 tells us that an error consistent model will learn most functional forms of interest and their true parameters provided sufficient data\textsuperscript{18} This property can be called implicit estimation consistency where we do not allow parameters of interest to enter the estimation

\textsuperscript{16}Particularly Proposition 5 on page 609.
\textsuperscript{17}The term consistency carries three different meanings in this paper. Namely, consistency of model variable attributions (e.g. Shapley values), error consistency for universal approximator in machine learning and estimator consistency with respect to $\beta$ (see also Zhao and Yu (2006); Munro (2018)).
\textsuperscript{18}An intuitive illustration of how a SVM with radial kernel can approximate almost any function is given in Appendix.
stage. The Shapley decomposition can now be used make the functional form explicit and to test parameterisations of the DGP.

4.2 Estimator bias

When has a model sufficiently converged for well informed inference, e.g. judged by its Shapley share coefficients from Eq. 9? Before addressing this question, let us connect model Shapley values to parametric functional forms which have a finite decomposition.

Lemma 1 (model decomposition): There exists a decomposition \( \hat{\Psi}(x) = \hat{\psi}_c(x) = \hat{f}(x) \) if the equation \( \hat{\Psi}(x) = \Phi^S \) is solvable for each \( x_k \), with \( k \in \{1, \ldots, m\} \). The proof is simple as \( x(\Phi^S) \) can be used to construct \( \hat{\Psi} \).

A decomposition \( \hat{\Psi} \) can be called an additive functional form representation of \( \hat{f} \) with the Shapley decomposition \( \Phi^S \) being the trivial representation. That is, \( \hat{\Psi} \) is a parameterisation of \( f \) for which the following results holds.

Theorem 4.2. (composition bias): Let \( f \) be a DGP and \( \Psi^*(x) = \sum_{c=1}^{C} \psi^*_c(x) = f(x) \) the true local decomposition of \( f \). Let \( \hat{f} \) be an error consistent model according to Theorem 4.1 with a local decomposition \( \hat{\Psi}(x) = \sum_{c=1}^{C} \hat{\psi}_c(x) = \hat{f}(x) \), e.g. its Shapley decompositions \( \Phi^S \). Applying the Shapley regression (7), \( \hat{\Psi} \) is unbiased with respect to \( \Psi^* \) if and only if \( \hat{\beta}_c = 1 \), \( \forall c \in \{1, \ldots, C\} \). Particularly, there exists a minimal \( m_u \) for which \( \hat{\beta}_c = 1 \), \( \forall c \in \{1, \ldots, C\} \) at a chosen confidence level. The proof is given in the Appendix.

Theorem 4.2 implies that \( \hat{\beta}^S \to 1 \), as \( m \to \infty \) for either \( \Phi^S \) or \( \hat{\Psi} \). Having \( \Phi^S \), the mapping \( \Phi^S \mapsto \hat{\Psi} \) can be used to test the functional form of \( f \). Corollary 1 extends this to local approximations of any form, i.e. for those to which Lemma 1 does not apply but a local decomposition can be formulated.

For a linear model, \( \hat{\beta}_c = 1 \) is nothing else as the unbiasedness of coefficients if the model is well specified. This can be seen from Proposition 3.2 and shows again that Shapley regressions reduce to the standard case in this situation.

For a general non-linear model, unbiasedness can only be assessed if \( \hat{\beta}_c = 1 \), \( \forall c \in \{1, \ldots, C\} \) due to the accuracy condition (3) required from each decomposition \( \hat{\Psi} \). The Shapley regression (7) tests linear alignment of \( \hat{\Psi} \) with the dependent variable, while the level of individual components \( \hat{\psi}_c \) may shift until \( \hat{\beta}_c = 1 \), \( \forall c \in \{1, \ldots, C\} \) for sample sizes smaller than \( n_u \). Consistency implies that such a shift happens towards the true level \( \psi^*_c \).

Robust component estimates

This leads to the definition of a robust component estimate: \( \hat{\psi}_c \) is said to provide a robust
estimation of $\psi_c^*$ within $\Omega \subseteq D \subseteq \mathbb{R}^m$, if $\mathcal{H}_0(\Omega) : \{\beta^S_c = 0|\Omega\}$ is rejected and

$$\mathcal{H}_1(\Omega) : \{\beta^S_c = 1|\Omega\} \text{ is not rejected}$$

at a chosen confidence level. That is if the chosen confidence bounds for $\hat{\beta}^S_c$ exclude zero but include one. Regarding the test for $\mathcal{H}_1^c$, one may set the confidence level to $\alpha_1 = 1 - \alpha_0$ for $\alpha_0$ being the desired confidence level to test against $\mathcal{H}_0$.\footnote{Practically, one can impose restrictions of the form $\hat{\beta}^S_c = 1$ and use a Wald test and not reject $\mathcal{H}_1^c$ with high probability $\alpha_1$ imposing tight confidence bounds on $\hat{\beta}^S_c$ which need to include one. The fully restricted test with $\sum_{c=1}^C \beta^S_c = C$ may be too lenient as loosely determined coefficients allow for greater flexibility for fulfilling that constraint, especially if Shapley regression coefficients are located on both sides of one. Instead one should require $\hat{\beta}^S_c = 1$, $\forall c \in \{1, \ldots, C\}$ individually.} Alternatively, one may define an acceptable range for $\hat{\beta}^S_c$ provided $\mathcal{H}_0$ can be rejected, e.g. $\hat{\beta}^S_c \in [0.9, 1.1]$ admitting a small amount of bias.

Both conditions are necessary to guarantee meaningful information content in $\hat{\psi}_c$. This can be seen by considering a linear model with a pure noise variable. The best least-squares fit will return $\hat{\beta}^S_c = 1$ by construction, but $\mathcal{H}_0$ is almost certain not to be rejected. The practicality of robust component estimates is that they can provide useful information despite a failing test for a model being unbiased, i.e. biases in component levels.

For instance, changes between different points in a region $\Omega$ are independent from the actual level of $\hat{\psi}_c$ if the model and target are well aligned. For example, the change of $\psi_c^*$ between two points $x_1, x_2 \in \Omega$ can be approximated by $\hat{\psi}_c = \psi_c^* + b_c$ with $b_c$ the component bias, if $\hat{\beta}^S_c \approx 1$,

$$\Delta \psi^*_c(x_1, x_2) \equiv \psi^*_c(x_2) - \psi^*_c(x_1) = \hat{\beta}^S_c(\hat{\psi}_c(x_2) + b_c) - \hat{\beta}^S_c(\hat{\psi}_c(x_1) + b_c) \approx \hat{\psi}_c(x_2) - \hat{\psi}_c(x_1) = \Delta \hat{\psi}_c(x_1, x_2).$$

### 4.3 Validity conditions for the Shapley regression framework

Eq. 7 is an auxiliary model in a linearised space of generated regressors (Pagan (1984)), minimising the log-likelihood

$$l(\beta^S, \theta; y, x) \sim -\frac{1}{2\sigma^2} \left[ (y - \Phi^S(\hat{\theta})\beta^S)^T (y - \Phi^S(\hat{\theta})\beta^S) \right].$$

Inference with regard to $\beta^S$ is valid under two conditions. First, the cross terms of the Fisher information must vanish, i.e. $\mathcal{I}(\beta^S, \hat{\theta}) = 0$.\footnote{Inference with regard to $\beta^S$ is valid under two conditions. First, the cross terms of the Fisher information must vanish, i.e. $\mathcal{I}(\beta^S, \hat{\theta}) = 0$. This is achieved by the two-step approach and sample splitting, such that the optimisation processes for $\theta$ and $\beta^S$ are independent from each other. Particularly, $\phi^S_k(\hat{\theta})$ are independent random variables when estimating $\hat{\beta}^S$. Sample splitting between a training dataset on which the model is fitted and a hold-out set for testing is common in machine learning, so this does not impose a constraint.} This is achieved by the two-step approach and sample splitting, such that the optimisation processes for $\theta$ and $\beta^S$ are independent from each other. Particularly, $\phi^S_k(\hat{\theta})$ are independent random variables when estimating $\hat{\beta}^S$. Sample splitting between a training dataset on which the model is fitted and a hold-out set for testing is common in machine learning, so this does not impose a constraint.
Second, the non-parametric part, $\Phi^S$ in our case, is required to be $\sqrt{n}$-consistent. The accuracy property of Shapley values (3) relates this to the error consistency of a model and its convergence rate $r_e \sim n^{-\xi}$, which may depend on the sample size. However, non-parametric techniques, including machine learning models, often converge slower, i.e. $\xi < \frac{1}{2}$. In this case, the relative rates of convergence can be accounted for via appropriate sample splitting between the training and the test set. Specifically, the condition for the maximal size of the test is

$$n_{\text{test}} \leq n_{\text{train}}^{2 \min_k \xi_k} \xi_k^{\text{lin.conv.}} = n_{\text{train}}^{2\xi}.$$  

(18)

The convergence rate of individual Shapley components $\phi_k^S$ are labelled $\xi_k$. If the $\xi_k$ are different, the smallest $\xi_k$ will dominate $r_e$ at some point, leading to a non-constant rate $r_e(n_{\text{train}})$ and the most conservative condition for $n_{\text{test}}^{\text{max}}$. In the case of equal and constant $\xi_k$, $r_e$ is constant, setting the maximally permissible test set for asymptotic inference. Rates of convergence usually depend on the data, model and algorithm used and are an active area of research. In practice, the rate of convergence $r_e$ and individual $\xi_k$ can be determined empirically by fitting model learning curves, i.e. the error and its component dependence on the sample size.

Condition (18) affects the asymptotic behaviour of $\hat{\beta}^S = \sqrt{n}(\hat{\beta}^S - \beta^S) \to_p \mathcal{N}(0, \mathcal{I}^{-1}(\hat{\beta}^S, \hat{\beta}^S))$, as $n \to \infty$ with $\xi = \min_k \xi_k$. If $\xi < \frac{1}{2}$, this quantity diverges resulting in an asymptotically biased estimator. Practically this means that confidence intervals from $\mathcal{I}$ will not overlap (or will fail to do so at some point) with one if $\beta^S = 1$. Thus, tests for the robustness of a component using $\mathcal{H}_1(\Omega)$ and model bias will fail despite the model being consistent. However, we do know $\lim_{m \to \infty} \hat{\beta}^S = 1$, meaning we can quantify the bias in $\hat{\beta}^S$ at any point, e.g. for deciding if a component estimate is sufficiently robust for practical purposes. Importantly, this does not impose restrictions on $n_{\text{test}}$ for tests against $\mathcal{H}_0(\Omega)$. Asymptotic inference on $\mathcal{H}_0(\Omega)$ is still valid without sample splitting because $\mathcal{I}(\beta^S, \hat{\theta}) = 0$ if $\beta^S = 0$ (Pagan [1984]), but not for other hypotheses.

We see from the above discussion that the only possible true values for $\beta^S$ are $\beta^S \in \{0, 1\}^m$. Provided estimator consistency holds according to Theorem 4.1 this can be understood as follows. If there is a relationship between the target $y$ and $x_k$ (or $\psi_k$ more generally), then $\beta^S_k = 1$ otherwise $\beta^S_k = 0$. Intuitively this means that either there is a signal or not. 

21 See for example Andoni et al. (2014); Sutskever et al. (2013); Biau (2012); Scornet et al. (2014); Steinwart and Scovel (2007); Christmann and Steinwart (2008) and references therein.
5 Applications

5.1 Numerical simulations

Let us first showcase the Shapley regression framework in a controlled experiment. I investigate the statistical inference properties of NN, SVM and RF for learning low-order polynomials, considering the following DGP

\[ \Psi^*_1 \equiv f_1(\gamma; x; \beta) = \beta_0 + \beta_1 x_1^\gamma + \beta_2 x_2 + \beta_3 x_3 + \sigma \epsilon, \quad \gamma \in \{2, 3\} \quad (20) \]

\[ \Psi^*_2 \equiv f_2(\gamma; x; \beta) = \beta_0 + \beta_1 x_1^2 + \beta_2 x_1 x_2 + \beta_3 x_1 + \beta_4 x_2 + \beta_5 x_3 + \sigma \epsilon, \quad (21) \]

Each \( x_k, k \in \{1, 2, 3\} \) is i.i.d.-sampled from the standard normal distributions \( \mathcal{N}(0, 1) \). The last term in both processes is an independent noise term, where \( \sigma \) is the noise level and \( \epsilon \) is also sampled from \( \mathcal{N}(0, 1) \). The intercept and coefficients \( \beta \) are set to the vectors \( \beta f_1 = (0, 2, 4, 0.5) \) and \( \beta f_2 = (2, 2, 1, 1, 0.5) \) for \( f_1, \gamma \) and \( f_2 \), respectively. Process \( f_1, \gamma \) can be seen as the simple case with a single non-linearity featuring strong (\( x_2 \)) and weak (\( x_3 \)) controls. Process \( f_2 \) is more complex with multiple non-linearities (polynomial and interaction), controls of different strength and an intercept. The noise level \( \sigma \) is set to zero (no noise) and 10% of the standard deviation of each DGP, i.e. \( 0.1 \sigma_{\text{std}}(y) \).

Sample sizes are equally spaced on a logarithmic scale between one hundred and ten thousand, i.e. \( n_q = 10^q \) for \( q \in \{2, 2.5, 3, 3.5, 4\} \). Using only the raw feature values \( x_k, k \in \{1, 2, 3\} \), NN, SVM and RF are calibrated, trained and tested on three independently generated datasets \( x_{cv}, x_{train} \) and \( x_{test} \), respectively, for each sample size. That is, the models are not given the functional forms (20) and (21). Rather they have to infer them from \((x, y)_{\text{train}}\). After fitting each model using \( x_{\text{train}} \), the Shapley value decomposition (3) on \( x_{\text{test}} \) is used together with the knowledge of the DGP \( (\Psi^*) \) to estimate \( \hat{\beta} \) learned by each model. The mappings \( \hat{\Psi}(\Phi^S) \) are given in the Appendix. Each configuration of DGP, model, noise level and sample size is simulated 50 times for numerical robustness.

Fig. 1 shows the averaged error learning curves for all cases. The NN and SVM achieve almost perfect fits from a sample size of \( n = 10^{2.5} \approx 300 \) on in the noiseless case, and stagnate close to the minimally achievable error in the noisy case. Process \( \Psi^*_{1,3} \) is a slight exception as the cubic non-linearity needs more data to learn. The RF has higher test errors overall. There are two reasons for this. One is its hierarchical structure. Even though individual trees are randomised, they still prefer certain features at higher split points (if more than one feature

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22 Hyper-parameter tuning via cross-validation.

23 Cross-validation is limited to \( n_{cv} \leq 1000 \) for the SVM due to the computational costs related to the underlying non-sparse quadratic programming problem.

24 All simulations have been done on the cloud using about \( 10^4 \) computing hours on standard 2.4GHz CPU nodes.
is allowed at each split), learning different aspects of the target function sequentially as more training data become available. Second, RF do axes-aligned splits of the input space in hyper-rectangles. This makes the learning of smooth functions, such as polynomials, more difficult as more hyper-rectangles are needed compared to fitting step-like functions.

These differences in learning can now be investigated in the Shapley regression framework. Fig. 2 shows the convergence towards the true $\beta$ for DGP 2 (21) with noise. Convergence is measured against the blue dotted unit line for normalised coefficients ($\hat{\beta}_u = \hat{\beta}/\beta$). The black lines are average $\hat{\beta}_u$ with 90% confidence intervals given by the blue shaded areas.

The differences in the error learning curves of Fig. 1 are reflected in the coefficient learning curves of Fig. 2. Across most of the sample size range, the NN and the SVM can be said to have learned unbiased representations of DGP 2 judged by the criterion $\hat{\beta}_u \in [0.9, 1.1]$ (green lines). All individual components have been estimated robustly according to this definition. To the contrary, the RF adapted to some parts of $f_2$ quickly (e.g. $\beta_2$), while taking substantially more data for others, particularly $\beta_5$. These differences make explicit the property of regression and classification trees by which they differentiate between variables according to their contribution to the loss function. The interaction with strong coupling $\beta_2$ features stronger in $f_2$ and is learned before the weak control $\beta_5$.

A rigorous way to test for robustness is to test for $H_1$ (15). We have to take the convergence rates $r_e$ of individual components of $\hat{\Psi}$ into account for this. I consider estimates of linear convergence rates of $n^{-\xi}$ as a first-order approximation. These are shown in Tab. 1. Good linear approximations are indicated by a high $R^2$. This is mostly the case. Where it is not, convergence slows with increasing sample size.

We have $\xi \geq \frac{1}{2}$ in the majority of cases, needing no further adjustment with regard to $H_1$. For the current case of DGP 2 with noise, I set $\xi = \frac{1}{4}$ (about the rate of the NN) for all models for demonstration. Tests for $H_1$ setting $\alpha_1 = 1 - \alpha_0 = 90\%$ are shown by the 10% confidence intervals (red shaded areas) in Fig 2 after adjusting for degrees of freedom according to (18). This test for component robustness is more stringent than the one based on the condition $\hat{\beta}_u \in [0.9, 1.1]$. The estimates of all models and parameters converge despite noise, but not all components can be said to be estimated robustly even for large samples sizes.

This simulation study has showcased the main aspects of statistical inference in the Shapley regression framework. Let us turn to a real-world example.

5.2 Macroeconomic time series modelling

I present a comparative analysis using NN, SVM and RF within the Shapley regression framework to model quarterly UK and US macroeconomic time series between 1955-2017 and 1965-25. Flat or negative $\xi$ may suggest even lower values. However, low rates and poor linear fits are more associated with problems in learning in the presence of noise (a general problem). Furthermore, using too high a rate leads to more conservative estimates when testing for robustness.

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25 Flat or negative $\xi$ may suggest even lower values. However, low rates and poor linear fits are more associated with problems in learning in the presence of noise (a general problem). Furthermore, using too high a rate leads to more conservative estimates when testing for robustness.
2017, respectively. I particularly look at year-on-year percentage changes in output, unemployment and inflation on a one-year horizon using a simple lead-lag setting. Features lag the respective target variable by one year,

\[ y_{t(q)} = \hat{f}(x_{t(q)-4}, y_{t(q)-4}; \theta) + \hat{\epsilon}_{t(q)}, \tag{22} \]

where also one yearly lag of the dependent variable is included. This exercise can provide a starting point for narratives around driving factors behind changes in the target variables taking non-linearities from machine learning models into account, complimentary to more structural approaches. A summary of both datasets is given in Tab. 2. All series are standardised to a mean of zero and unit variance (z-scores). This makes results comparable for different target variables, across models and time. However, the current approach is not suitable for forecasting because of the built-in look-ahead bias through the variable standardisation and the model training process resulting in information leakage.

5.2.1 Cross-validation and training

Nested cross-validation is used for training and testing. This also addresses the relative smallness of both datasets. An outer loop randomly splits the data into ten (90%,10%) folds for training and testing, respectively. A model is evaluated based on its out-of-sample prediction performance on all outer test sets. Within each such training set, an inner loop of ten (90%,10%) splits for the calibration (cross-validation) of model hyper-parameters and training is used. This procedure is repeated over 50 bootstrap iterations for numerical stability. The baseline for comparison is a non-penalised linear regression model fitted and tested on the same folds as the machine learning models. The time dimension of our data is addressed by including a lag of each dependent variable and by accounting for potentially remaining autocorrelation in the error terms at the regression stage using heteroskedasticity and autocorrelation robust standard errors.

5.2.2 Test performance

The key motivation for using machine learning models in the first place is that they are expected to deliver more accurate predictions, i.e. a smaller test error. I test this by comparing the out-

\(^{26}\)Two additional benchmark models have been evaluated. A linear model with elastic net regularisation and a vector autoregressive model (VAR) with one lag. The VAR has been fitted on a quarterly frequency and evaluated on its in-sample forecast performance on a one-year horizon. One lag provides approximately the same information content as either machine learning models or the other linear benchmarks receive. The inclusion of more lags would provide it with additional information of the time series structure of the data which the other models did not have access to, while also increasing the risk of overfitting. Out-of-sample and in-sample test performance for the elastic net and VAR have only been marginally better on average than the baseline regression, while introducing substantial bias in the parameter estimates and potentially overfitting the data, respectively.
of-sample root-mean-square errors (RMSE) of all models. Comprehensive error statistics are given in Tab. 3. Machine learning models outperform the benchmark (Reg) in the majority of cases, often by a wide margin. The two exceptions are the SVM for modelling UK inflation and unemployment where both perform similarly.

Comparing the UK and US, all models perform better for the US in absolute terms. This suggests that we are missing some important factors for the UK. Two aspects not very well represented in our data are the ‘smallness and openness’ of the UK economy relative to the US and its much larger financial sector compared to its real economy. Thus, taking more external and financial factors into account may improve overall model performance for the UK. Moreover, machine learning models provide larger performance gains for the US than the UK relative to the linear baseline. This suggests a greater importance of non-linearities when investigating the US economy.

The data cover two major episodes of macroeconomic stress, the stagflation period of the 1970s and the global financial crisis (GFC) 2008 and its aftermath. These are associated with sharp turning points in economic indicators, likely associated with non-linear dynamics. Machine learning models generally cope better with these situations, partly explaining their better test performance.

5.2.3 Model inference

I focus on modelling unemployment. The graphical decomposition of out-of-sample test predictions for all models and both countries is shown in Figures 3–6. The contributions attributed to the each variable according to (2) sum up to the model prediction at each observation (accuracy property of Shapley values). Only the six largest components by absolute values of the SSC (9) are shown explicitly. These mostly constitute the great majority of model output. The remaining four variables are grouped in the light gray contributions.

All models consistently attribute importance to the same variables, mostly changes in GDP and prices, but also the policy rate, the money supply and private debt. The current account balance also features in all models for the US, though with a relatively small share. Considering the NN (Fig. 3) and the SVM (Fig. 4), the largest deviations from the target lines are at turning points. Particularly, during the 2008 GFC where the spike in unemployment in either countries is largely missed by both models. With the exception of the NN in the US, which did, however, not capture its full magnitude. This is much so for the UK, where changes in unemployment have a more volatile profile overall.

27 Variables are ordered column-wise from left to right from the largest to the smallest absolute SSC. ‘Others’ always comes last.

28 It is not surprising that neither model is able to capture the crisis dynamics well. There was no comparable event in the data from which they could have been able to generalise, i.e. have learned. Assuming that future crises are similar in their dynamics, they may provide better guidance.
Assuming that turning points are somewhat non-linear phenomena and that they are better captured for the US, one expects both models to exhibit some non-linear variable dependencies and that these are stronger for the US than for the UK. This is qualitatively confirmed in Fig. 7. It plots the features dependence learned by all models for the UK (LHS) and the US (RHS) for changes in GDP, private debt and broad money in terms of their Shapley contributions for each observation. Approximate functional forms are traced out by best-fit degree-three polynomials (dashed lines). It is reassuring that all models learned comparable functional dependencies, which are considerably non-linear29 The exception to this is the SVM for the UK, which has an almost flat structure. The SVM is actually very similar to the linear benchmark in this case (as for modelling UK inflation) which explains their comparable performances. Another difference between models is the relative magnitude of model output allocated to GDP for the US, especially for negative feature values. We would expect these to be comparable for an unbiased model. However, for practical purposes it may be enough for these components to be estimated robustly. Both aspects will be discussed below.

Considering the NN (upper part of Fig. 7), the learned functional dependencies are comparable for both countries. As expected, relations tend to be more non-linear in the US case indicated by the higher curvature of the best fit lines. This is particularly so for broad money. Here, the RHS endpoints correspond to the end of the Bretton Woods system, the “oil shocks” of the 1970s and the stagflation period. The latter is highlighted by vertical dashed lines in Fig 3-6. For the UK, this endpoint corresponds to the first oil shock (1973) and the onset of the stagflation period. This means that the NN learned a functional form connecting different economic regimes, suggesting it internalised structural changes in the underlying DGP to some degree30

The discussion has been qualitative so far. The Shapley regression framework allows for rigorous statistical inference analysis on each model. The results of this exercise for unemployment, inflation and GDP are summarised in Tab. 4-6 respectively. The SSC Γ from (9) are shown for the UK (LHS) and the US (RHS) for all three target variables, respectively. Estimated standard errors31 p-values regarding tests against $H_0(\Omega)$ (8) and values of the Shapley regression coefficients $\hat{\beta}^S$ from (7) are shown in parentheses. Square brackets indicate robustness of variable components based on $\hat{\beta}^S \in [0.9, 1.1]$, excluding the benchmark (Reg) where components are robust by definition.

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29 The turning points of the polynomial fits at the limits of input spaces are artefacts of the polynomial approximation. Nevertheless, these extrema indicate saturation beyond a certain threshold which also is a non-linear phenomenon interesting in itself.

30 One has to be careful with structural interpretations however. Machine learning models fit observed patterns in the data, i.e. they can be interpreted as a form of reduced form estimation.

31 The variance estimator from Ōkui (2014) has been used to account for time series autocorrelations with a maximal lag of $\sqrt{|\Omega|}$. 
The discussion focuses again on unemployment (Tab. 4). Taking GDP for the NN as an example, its SSC $\Gamma_{GDP}^S$ is interpreted as follows. About 21% of model predictions for the UK are significantly, robustly and negatively attributed to changes in output with estimated 95% confidence bounds of $\pm 1.7\%$ around that level. GDP is also the dominant variable in this model measured by $|\Gamma_{GDP}^S|$ relative to the other features. Importantly, the coefficients of all models are broadly in line with each others. For example, all models estimate a robust SSC for GDP for the UK. We observed different relative magnitudes for the importance of GDP in the US case in Fig. 7. This is reflected in the different magnitudes of SSC for all three models. Also, neither model estimates this contribution robustly, such that we should be careful with interpreting these results.

Between models, the SSC structure of the RF is often qualitatively different to that of the other two models. It tends to attribute dominant shares to a few features while attributions are more balanced for the NN and the SVM. This is again understood by the hierarchical structure of trees as seen in Section 5.1. Again, this characteristic may be desired in situations where variable selection is important. The RF may be seen as the “machine learning equivalent” of the LASSO\textsuperscript{32} in this sense.

Apart from the better test performance, machine learning models extract more significant variable contributions from the data compared to Reg. This is due to their more flexible model structure but also highlights the importance of non-linearities in the DGP. However, few SSC are robust. Thus, none of the models can be said to have learned an unbiased representation. This is likely due to the relative smallness of our datasets and shows some of the limitations of using machine learning in this context.

Overall, the Shapley regression analysis summarised in Tab. 4-6 allows statistical inference and the communication of modelling results similar to that of the linear model, while leveraging on the benefits of machine learning.

\subsection*{5.2.4 Feature dependence}

Eq. 2 requires us to compute conditional expectations of the form $E[\hat{f}_C] \equiv E_{x\setminus C}\{\hat{f}(x)|x_C\}$. Here, $S$ is the set of non-missing features in a coalition. For this to be exact, one has to assume feature independence, which can be a strong assumption for many applications. At the lowest order, variable dependencies can be linear, i.e. correlations, but they may take more complex forms. Corollary 4.1 says that error consistent models will learn such dependencies given enough training data. It is not clear, however, how to extract this information from a model without the use of Shapley values or more knowledge about the DGP. We would need a method to calculate $E[\hat{f}_C]$ which respect feature dependencies. A fast algorithm for tree-based models to do just that is given in Lundberg et al. (2018). The idea is to evaluate imputed

\textsuperscript{32}Least Absolute Shrinkage and Selection Operator.
inputs to $\mathbb{E}[\hat{f}_C]$ by following tree paths. The training set observations of the leaf node that input falls into serves as its background dataset hereby respecting observed dependencies within the data. This is conceptually similar to Wager and Athey (2018), where leaf nodes serve to match the treated and the untreated controlling for potentially complex relationship between other variables.

The comparison of Shapley values for using this exact method with values obtained using the whole training dataset as the background, i.e. assuming feature independence, allows one to quantify the error made by the independence assumption. Universal approximators learn the same feature dependencies given enough data (see Fig.2 & 7). It can therefore be assumed that differences in Shapley values observed for tree models will be comparable to models where no exact solution exists, such as NN and SVM. Thus, this comparison provides an indication for which variable contributions can be judged reliable under the independence assumption using these models.

The comparison for the RF between the exact and the approximate solution for computing $\mathbb{E}[\hat{f}_C]$ is given in Fig.8 for modelling unemployment. It shows differences $\Delta \Gamma^S$ of SSC (9) between both cases relative to the root-mean-square of estimated standard errors of the absolute value of each variable component $\mu_k = |\Gamma^S_k|$ for both cases. None of the differences for either country would test as statistically significant. This means that the assumption of feature independence is justified in the current case. The overall low cross-correlation between variables for both countries indicates this, while the above comparison accounts for more general relations.

6 Conclusion

This paper proposed Shapley regressions as a general framework for statistical inference on non-linear models, particularly those from machine learning. The underlying idea is to formulate a regression problem within the space of transformed inputs defined by the Shapley decomposition of a model. Besides the interpretability of individual model predictions, this opens machine learning problems to parametric statistics, including many techniques from econometrics, and vice versa.

There are two appealing properties of the Shapley regression framework, which provide justification for this approach. First, Shapley values have a clear interpretation derived from their game theoretic origin and desirable properties. Second, Shapley regression is identical to a conventional regression analysis for a model linear in parameters. Thus, Shapley regressions can be interpreted as an extension of parametric statistical inference into the space of non-linear and non-parametric models.

I showed that commonly used model classes in machine learning are estimation consistent for the important class of piecewise analytic functions, which covers most cases of interest. Numer-

33All results presented for the RF in this case study are based on exact calculations.
ical simulations were used to investigate the asymptotic inference properties of state-of-the-art machine learning models, such as artificial neural network, support vector machines and random forests.

Finally, I applied the Shapley regression framework to model UK and US macroeconomic time series. Machine learning models outperformed the linear benchmark most of the time. Importantly, leading feature attributions of most machine learning models are comparable. This provides further trust in the use of machine learning models as one can concentrate on technical aspect of a model amenable to a particular problem. For example, random forests tend to produce hierarchical feature attributions, making them suitable to high-dimensional problems of variable selection.

The summarising concept of Shapley share coefficient (SSC) was introduced which are close in their interpretation to the coefficients of a linear model. The main difference to statistical inference on a linear model is that results are only locally valid within the considered region. This puts more burden on careful testing of results, especially in the presence of strong non-linearities.

Despite good test performance, not all estimated and significant variable contributions turned out to be robust. This warrants caution for the application of machine learning models and suggest comprehensive model evaluation before putting such a model into practice, e.g. to inform decisions.

In summary, the Shapley regression framework provides a rigorous approach for addressing the black-box critique of machine learning models, including those voiced against modern development in artificial intelligence. In their essay “We built them, but we don’t understand them” (Kleinberg and Mullainathan (2015)) Jon Kleinberg and Sendhil Mullainathan set out the challenges and risks around designing and using algorithms to inform decisions if these algorithms are poorly understood. Shapley regressions offer a tool to partly address such concerns through well-grounded statistical inference. Thus, they may extend the scope of applicability for machine learning models, particularly for informing decisions in the presence of ever more granular data sources.
Appendix

Box 1: Statistical inference recipe for machine learning models

1. Cross-validation, training and testing of a model $\hat{f}$

2. Model decomposition
   
   (a) Shapley value decomposition $\Phi^S(\hat{f})$ [Eq. 2] on test set
   
   (b) (if any) Test of assumptions and approximations made to calculate $\Phi^S$
   
   (c) (optional) Mapping of $\Phi^S$ to parametric functional form $\hat{\Psi}(\Phi^S)$
       [see Section 4.2]

3. Model inference
   
   (a) Shapley regression [Eq. 7] with appropriate standard errors

   $y_i = \Phi^S_i \beta^S + \epsilon_i = m \sum_{k=0}^{m} \phi^S_k(\hat{f}, x_i) \hat{\beta}^S_k + \epsilon_i$

   [replace $\Phi^S$ with $\hat{\Psi}$ in case of 2(b)]

   (b) Assessment of model bias and component robustness based on $\hat{\beta}^S$
       over a region $\Omega$ of the input space:

   Robustness (components): $H^k_0 : \{\beta^S_k = 0|\Omega\}$ rejected and $H^k_1 : \{\beta^S_k = 1|\Omega\}$
       not rejected for single $k \in \{1,\ldots,m\}$

   Unbiasedness (model): $H^1_k : \{\beta^S_k = 1|\Omega\}$ not rejected $\forall k \in \{1,\ldots,m\}$

   (c) Calculate Shapley share coefficients (SSC) $\Gamma^S(\hat{f}, \Omega)$ [Eq. 9] and their standard errors
Figure 1: Error learning curves for DGP (20) and (21), root-mean-square error as fraction of DGP standard deviation. Source: Author’s calculation.
Figure 2: Parameter learning curves (black) for NN (left column), SVM (middle column) and RF (right column) for DGP $f_2$ \cite{21} with noise. Blue dashed line references true coefficient values (normalised to one). Blue and red shaded areas indicate 90\% and 10\% confidence intervals (CI) for fast and slow convergence rates $r_e \sim n^{-\xi}$, with $\xi \geq \frac{1}{2}$ and $\xi = \frac{1}{4}$, respectively. Robustness intervals $[0.9, 1.1]$ marked by green dashed lines. Source: Author's calculation.
Figure 3: Neural network (NN) out-of-sample Shapley test decomposition for modelling unemployment for the UK (upper) and the US (lower). Source: ONS, OECD and author’s calculation.
Figure 4: Support vector machine (SVM) out-of-sample Shapley test decomposition for modelling unemployment for the UK (upper) and the US (lower). Source: ONS, OECD and author’s calculation.
Figure 5: Random forest (RF) out-of-sample Shapley test decomposition for modelling unemployment for the UK (upper) and the US (lower). Source: ONS, OECD and author’s calculation.
Figure 6: Linear regression (Reg) out-of-sample Shapley test decomposition for modelling unemployment for the UK (upper) and the US (lower). Source: ONS, OECD and author’s calculation.
Figure 7: Feature dependence based on Shapley decomposition of NN (upper), SVM (middle) and RF (lower) for selected variables for modelling unemployment in the UK (LHS) and the US (RHS). Dashed lines are best-fit degree-3 polynomials. Source: BOE, ONS, BIS, OECD and author’s calculation.
Figure 8: Differences $\Delta S$ of SSC for modelling unemployment using RF for assuming feature independence versus respecting dependencies learned by the model relative to the root-mean-square of estimated standard errors of the absolute value of each variable component $\mu_k = |\Gamma_k^S|$ in both cases. Source: Author’s calculation.
Table 1: Estimated exponents of convergence rates $r_e \sim n^{-\xi}$ of machine learning models for simulated DGP (20) & (21) and coefficients of determination ($R^2$). Source: Author’s calculation.
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<th>Private debt</th>
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Table 3: UK and US model test statistics: Root mean squared error (RMSE), generalisation error (difference between test and training error as a fraction of the test error), squared model bias and variance (as fractions of the mean squared error) for different bootstrap realisations from nested cross-validation. Best/worst RMSE for each target variable is highlighted in green/red. Source: Author’s calculation.
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<td>(.009; .00; 1.52)</td>
<td>(.020; .06; 0.28)</td>
</tr>
<tr>
<td>CA</td>
<td>[0.057]*</td>
<td>0.055**</td>
<td>0.012</td>
<td>0.054</td>
<td>0.084***</td>
<td>0.107*</td>
<td>[0.073]*</td>
<td>0.095*</td>
</tr>
<tr>
<td></td>
<td>(.003; .07; 0.91)</td>
<td>(.003; .04; 1.24)</td>
<td>(.001; .16; -3.68)</td>
<td>(.003; .07; 0.11)</td>
<td>(.011; .01; 0.82)</td>
<td>(.014; .05; 0.84)</td>
<td>(.009; .09; 1.00)</td>
<td>(.012; .06; 0.17)</td>
</tr>
<tr>
<td>ERI</td>
<td>0.022**</td>
<td>[0.031]</td>
<td>0.051***</td>
<td>0.041</td>
<td>0.057</td>
<td>0.071</td>
<td>0.043</td>
<td>0.048</td>
</tr>
<tr>
<td></td>
<td>(.005; .04; 1.16)</td>
<td>(.003; .17; 1.09)</td>
<td>(.007; .00; 1.71)</td>
<td>(.005; .24; 0.10)</td>
<td>(.009; .30; 0.30)</td>
<td>(.011; .31; 0.37)</td>
<td>(.005; .29; 0.61)</td>
<td>(.007; .36; 0.08)</td>
</tr>
<tr>
<td>Unemployment 4l</td>
<td>[0.110]**</td>
<td>0.124***</td>
<td>0.139***</td>
<td>0.104**</td>
<td>0.113**</td>
<td>0.141***</td>
<td>0.202***</td>
<td>0.042</td>
</tr>
<tr>
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<td>(.015; .00; 1.09)</td>
<td>(.016; .01; 1.22)</td>
<td>(.017; .00; 0.84)</td>
<td>(.013; .04; 0.25)</td>
<td>(.004; .00; 1.34)</td>
<td>(.010; .00; 1.77)</td>
<td>(.014; .00; 1.95)</td>
<td>(.002; .66; 0.09)</td>
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</tbody>
</table>

Table 4: Shapley share coefficients $\Gamma^S_k$ for modelling UK (LHS) and US (RHS) unemployment on a one-year horizon for different models. Significance levels: * (10%), ** (5%), *** (1%). Green/red refers to positive/negative coefficients significant at the 10% level. Estimated standard errors for $\Gamma^S_k$, p-values regarding $H_0(\Omega)$ and values of the Shapley regression coefficients $\hat{\beta}^S$ are shown in parentheses. The actual regression coefficients $\hat{\beta}$ are shown for Reg. Square brackets indicate robustness of $\Gamma^S_k$ for $\hat{\beta}^S \in [0.9,1.1]$, not shown for Reg. Source: Author’s calculation.
<table>
<thead>
<tr>
<th>Country</th>
<th>INFLATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>GDP</td>
<td>NN</td>
</tr>
<tr>
<td></td>
<td>0.049</td>
</tr>
<tr>
<td></td>
<td>(.009; .47; 0.04)</td>
</tr>
<tr>
<td>Labour prod</td>
<td>0.027**</td>
</tr>
<tr>
<td></td>
<td>(.004; .22; 0.91)</td>
</tr>
<tr>
<td>Broad money</td>
<td>0.050**</td>
</tr>
<tr>
<td></td>
<td>(.003; .04; 2.26)</td>
</tr>
<tr>
<td>Private debt</td>
<td>0.041**</td>
</tr>
<tr>
<td></td>
<td>(.006; .03; 1.56)</td>
</tr>
<tr>
<td>Unemployment</td>
<td>-0.036**</td>
</tr>
<tr>
<td></td>
<td>(.008; .00; 2.96)</td>
</tr>
<tr>
<td>GDHI</td>
<td>0.316***</td>
</tr>
<tr>
<td></td>
<td>(.021; .00; 0.87)</td>
</tr>
<tr>
<td>Policy rate</td>
<td>0.053</td>
</tr>
<tr>
<td></td>
<td>(.011; .27; 0.50)</td>
</tr>
<tr>
<td>CA</td>
<td>-0.037*</td>
</tr>
<tr>
<td></td>
<td>(.005; .07; 1.73)</td>
</tr>
<tr>
<td>ERI</td>
<td>-0.036</td>
</tr>
<tr>
<td></td>
<td>(.007; .06; 0.09)</td>
</tr>
<tr>
<td>Inflation 4l</td>
<td>[0.356]***</td>
</tr>
<tr>
<td></td>
<td>(.023; .00; 1.08)</td>
</tr>
</tbody>
</table>

Table 5: Shapley share coefficients $\Gamma_{S_k}^S$ for modelling UK (LHS) and US (RHS) inflation on a one-year horizon for different models. Significance levels: * (10%), ** (5%), *** (1%). Green/red refers to positive/negative coefficients significant at the 10% level. Estimated standard errors for $\Gamma_{S_k}^S$, $p$-values regarding $H_0(\Omega)$ and values of the Shapley regression coefficients $\hat{\beta}_S$ are shown in parentheses. The actual regression coefficients $\hat{\beta}$ are shown for Reg. Square brackets indicate robustness of $\Gamma_{S_k}^S$ for $\hat{\beta}_S \in [0.9, 1.1]$, not shown for Reg. Source: Author’s calculation.
<table>
<thead>
<tr>
<th>target</th>
<th>country</th>
<th>GDP model</th>
<th>UK</th>
<th>US</th>
</tr>
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<tr>
<td></td>
<td></td>
<td>NN</td>
<td>SVM</td>
<td>RF</td>
</tr>
<tr>
<td>Labour prod</td>
<td></td>
<td>-0.026</td>
<td>-0.090</td>
<td>-0.069</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(.003; .47; 0.15)</td>
<td>(.007; .18; -1.87)</td>
<td>(.005; .47; -0.09)</td>
</tr>
<tr>
<td>Broad money</td>
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<td>0.096*</td>
<td>0.182</td>
<td>0.083**</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(.014; .10; 1.15)</td>
<td>(.021; .10; 1.71)</td>
<td>(.014; .04; 1.21)</td>
</tr>
<tr>
<td>Private debt</td>
<td></td>
<td>-0.109***</td>
<td>-0.043</td>
<td>-0.126***</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(.011; .01; 1.51)</td>
<td>(.009; .40; -1.24)</td>
<td>(.013; .00; 1.71)</td>
</tr>
<tr>
<td>Unemployment</td>
<td></td>
<td>0.125***</td>
<td>0.065***</td>
<td>0.094***</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(.010; .00; 1.65)</td>
<td>(.006; .00; 4.93)</td>
<td>(.006; .00; 3.05)</td>
</tr>
<tr>
<td>GDHI</td>
<td></td>
<td>-0.043</td>
<td>-0.075</td>
<td>-0.054</td>
</tr>
<tr>
<td></td>
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<td>(.004; .41; 0.43)</td>
<td>(.010; .50; 0.02)</td>
<td>(.007; .38; 0.32)</td>
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<tr>
<td>Inflation</td>
<td></td>
<td>-0.192***</td>
<td>-0.100*</td>
<td>-0.062***</td>
</tr>
<tr>
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<td></td>
<td>(.028; .00; 1.27)</td>
<td>(.016; .08; 2.82)</td>
<td>(.010; .00; 2.45)</td>
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<td>Policy rate</td>
<td></td>
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<td>-0.236***</td>
<td>-0.242***</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(.021; .00; 1.12)</td>
<td>(.031; .00; 1.62)</td>
<td>(.015; .00; 1.02)</td>
</tr>
<tr>
<td>CA</td>
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<td>0.030</td>
<td>0.044</td>
<td>0.020</td>
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<td>(.002; .39; 0.40)</td>
<td>(.003; .16; -2.10)</td>
<td>(.001; .04; -4.96)</td>
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<tr>
<td>ERI</td>
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<td>-0.085**</td>
<td>-0.045</td>
<td>-0.115**</td>
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<tr>
<td></td>
<td></td>
<td>(.012; .04; 1.52)</td>
<td>(.008; .12; 3.32)</td>
<td>(.009; .02; 1.41)</td>
</tr>
<tr>
<td>GDP 4l</td>
<td></td>
<td>0.095***</td>
<td>0.121***</td>
<td>0.134***</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(.007; .00; 1.69)</td>
<td>(.008; .01; 3.36)</td>
<td>(.008; .01; 1.42)</td>
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</tbody>
</table>

Table 6: Shapley share coefficients $\Gamma_k^S$ for modelling UK (LHS) and US (RHS) GDP on a one-year horizon for different models. Significance levels: * (10%), ** (5%), *** (1%). Green/red refers to positive/negative coefficients significant at the 10% level. Estimated standard errors for $\Gamma_k^S$, p-values regarding $H_0(\Omega)$ and values of the Shapley regression coefficients $\hat{\beta}^S$ are shown in parentheses. The actual regression coefficients $\hat{\beta}$ are shown for Reg. Square brackets indicate robustness of $\Gamma_k^S$ for $\hat{\beta}^S \in [0.9, 1.1]$, not shown for Reg. Source: Author’s calculation.
Proofs

Proof of Proposition 3.1

Without loss of generality, we can write \( \hat{f} \) in terms of linear and non-linear components, \( \hat{f}(x) = \hat{f}_l(x) + \hat{f}_{nl}(x) \), e.g. using a Taylor expansion. The Shapley decomposition can then be written as

\[
\Phi_S(x) = \sum_{k=0}^{m} \phi^S_k = f(x) = \hat{f}_l(x) + \hat{f}_{nl}(x) = \hat{f}_l(x) = x\hat{\beta}.
\]

(23)

The first step follows from local accuracy and the third from the assumption of linearity. Properties (1)-(5) can be easily verified. \( \square \)

Proof of Proposition 3.2

Without loss of generality, we can again write \( \hat{f} \) in terms of a linear and a non-linear component, \( \hat{f}(x) = \hat{f}_l(x) + \hat{f}_{nl}(x) \). The Shapley regression can then be written as

\[
\Phi_S(x)\hat{\beta}^S = \sum_{k=0}^{m} (\phi^S_{k,l} + \phi^S_{k,nl})\hat{\beta}^S = \hat{f}_l(x)\hat{\beta}^S = x diag(\hat{\beta})\hat{\beta}^S = x\hat{\beta}.
\]

(24)

The last two steps follows from Proposition 1 and the uniqueness of the coefficients \( \hat{\beta} \) as solution to the convex least-squared problem. This can be made explicit for the OLS estimator. By setting \( x \rightarrow x diag(\hat{\beta}) \equiv xD_{\hat{\beta}} \), one obtains

\[
\hat{\beta}^S = \frac{xD_{\hat{\beta}}y}{(xD_{\hat{\beta}}y)^T(xD_{\hat{\beta}})} = \frac{D_{\hat{\beta}}Xy}{D_{\hat{\beta}}^2 x^T x} = D_{\hat{\beta}}^{-1} \hat{\beta} = 1_{n+1}.
\]

(25)

Proof of Theorem 4.1

I provide proofs for analytic and non-analytic models, reflecting prominent model types from machine learning. Analytic models \( \hat{f}(x,\theta) \) are differentiable almost everywhere (NN and SVM in our case).

Proof. (analytic models): Let \( \hat{f}(\theta, x) \) be a function of inputs \( x \in \mathbb{R}^m \) and parameters \( \theta \in \mathbb{R}^q \), which is \( (d' + 1) \) times differentiable, where \( d' \) is the degree of the highest polynomial \( p^{d'}(x) \) of the DGP \( f(\beta, x) \), such that the Taylor expansion of \( \hat{f} \) exists. Then, there exists an open interval \( \Omega \subset \mathbb{R}^m \) where the difference between \( f \) and \( \hat{f} \) is error consistent for each \( x' \in \Omega \) around \( a \). Namely,

\[
f - \hat{f}\big|_{\Omega}(x') = \sum_{k=0}^{m} (\beta_k - \hat{\beta}_k)p^d_k(x' - a) + R(\hat{f}^{(d'+1)}(c), (x' - a)^{(n-d)})
\]

(26)
That is, the polynomial expansion of $\hat{f}$ around $a$ will be functionally identical to $f$ up to a residual $R$ with $c$ between $x'$ and $a$. By assumption, (26) vanishes with increasing sample size, from which follows $(\beta - \hat{\beta}, R) \to 0$, as $m \to \infty$. 

Second, non-analytic models are tree-based models with $\hat{f}(x) \equiv T_{x\theta}(x) = \langle x_{\theta} \rangle_{x}$, with $T_{x\theta}$ describing the set leaf nodes of the model from training. Usually, $|T| \to \infty$, as $|x_{\text{train}}| \to \infty$. Examples are classification trees, random forests or extreme trees (Geurts et al. (2006)). The main difference to analytic models is that tree-based models are not differentiable. However, many tree-based models are based on bagging (e.g. forests) which smoothens model output (Bühlmann and Yu (2002)).

Proof. (non-analytic tree-based models): Let $x' \in \Omega \subseteq D$, where $D$ is the domain of $f$ and $\Omega$ is the leaf node region of $x'$, with $|\Omega|$ being the number of $x_\theta$ in this region. The difference between $f$ and $\hat{f}$ can then be written as

$$\left| f - \hat{f} \right|_{\Omega}(x') = \sum_{k=1}^{m} \beta_k p_k^d(x') - \frac{1}{|\Omega|} \sum_{j=1}^{|\Omega|} \hat{\beta_k} p_k^d(x_j)$$

$$= \frac{1}{|\Omega|} \sum_{j=1}^{|\Omega|} \left( \sum_{k=1}^{m} (|\Omega| \beta_k p_k^d(x') - \hat{\beta_k} p_k^d(x_j)) \right)$$

$$= \sum_{k=1}^{m} p_k^d(x') (\beta_k - \hat{\beta_k}).$$

We used the model optimising condition that values $x'$ fall into leave nodes with the same expected value, i.e. $\langle x_j \rangle_{\Omega} = x'$ in the limit $m \to \infty$. The above expression can then only vanish if $\beta - \hat{\beta} \to 0$ as $n \to \infty$. 

Proof of Corollary 4.1

For each $\epsilon > 0$ there is a neighbourhood $B_\delta$ of radius $\delta > 0$ around every $x' \in \Omega$, such that $|f - \hat{f}|_{x'} < \epsilon$. For each $B_\delta$ and $\epsilon$, there will be an large enough $n'$ such that there exist $\delta' \leq \delta$ and $\epsilon'$ with $\epsilon' + \epsilon'' < \epsilon$ for which $|f - \hat{f}|_{x'} < \epsilon'$ and $|\hat{f} - \hat{f}_p|_{x'} < \epsilon''$. The conclusion follows form the assumption of error consistency.

Proof of Theorem 4.2

The second part is a consequence of error consistency. For the first part, it is enough to show that the difference between $\hat{\Psi}$ and $\Psi^*$ vanishes beyond $n_u$. Here,

$$0 = \Psi^* - \hat{\beta}^s \hat{\Psi} = \sum_{c=1}^{C} \psi^*_c - \sum_{c=1}^{C} \hat{\beta}^s_c \hat{\psi}_c = \sum_{c=1}^{C} \psi^*_c - \hat{\psi}_c.$$

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The RHS of Eq. 28 only vanishes if $\psi^*_c = \hat{\psi}_c$, $\forall c \in \{1, \ldots, C\}$. 

---

**Series expansion of RBF SVM**

A SVM regression is the weighted sum of kernel-transformed inputs of the form

$$\hat{y}_i = \hat{f}(x_i; \hat{\theta}) = \sum_{i=1}^{n} \hat{\theta}_i K(x_i, x).$$  \hspace{1cm} (29)

The sum runs over the whole training set, while only contributions from the so-called support vectors close to the target regression line have non-zero weights. $K(x, x')$ is a kernel function which returns a distance-like scalar between its two inputs. Common kernels are radial biases functions (RBF; Gaussian kernel) or polynomial kernels. SVM are known to be error consistent (see e.g. Steinwart (2002); Christmann and Steinwart (2008)). For a RBF kernel, this can be intuitively understood by looking at the Taylor expansion of the kernel,

$$K_{\text{RBF}}(x, x') = \exp\left(\gamma |x - x'|^2\right)$$

$$= \sum_{n=0}^{\infty} \frac{(-\gamma |x|^2)^n}{n!} \sum_{n=0}^{\infty} \frac{(-\gamma |x'|^2)^n}{n!} \sum_{n=0}^{\infty} \frac{(-2\gamma x \cdot x')^n}{n!}.$$  \hspace{1cm} (30)

It contains an (infinite) sum of polynomials. The magnitude of each summand is proportional to the norm of $x_i$ and its alignment with the corresponding support vector as a measure of proximity.
Shapley value mapping from Section [5.1]

The knowledge of the DGP $f(x_i)$ and the model decomposition

$$\hat{y}_i = \hat{f}(x_i) = \sum_{k=0}^{3} \phi^S_k = \Phi(x_i)$$

(31)

can be used to estimate the coefficients learned by $\hat{f}$. To do so, we solve (31) for $\hat{x}_k(\Phi^S, f(x))$, $k \in \{1, 2, 3\}$. The results are used to estimate $y = \sum_k \hat{\beta}^S_k f_k(\hat{x})$ to obtain the normalised coefficients $\hat{\beta}^u = \hat{\beta}^S / \beta$. $\hat{f}$ generalises well to $f$ if the normalised coefficients are one at a chosen confidence level.

The reconstructed feature values $\hat{x}_k(\Phi^S, f(x))$ for the two processes (20, 21) are:

- $f_{1,\gamma}$:
  $$\hat{x}_1 = \pm (\phi_1/\beta_1)^{1/3}$$
  (32)
  $$\hat{x}_2 = \phi_2/\beta_2$$
  (33)
  $$\hat{x}_3 = \phi_3/\beta_3$$
  (34)

- $f_2$:
  $$\hat{x}_1 = \frac{-(\beta_2 x_2 + \beta_3) \pm \sqrt{(\beta_2 x_2 + \beta_3)^2 - 4\beta_1(\beta_4 x_2 + \beta_0 - \phi_{12})}}{2\beta_1}$$
  (35)
  $$\hat{x}_2 = \frac{\phi_{12} - \beta_1 x_1^2 - \beta_3 x_1 - \beta_0}{\beta_2 x_1 + \beta_4}$$
  (36)
  $$\hat{x}_3 = \phi_3/\beta_5$$
  (37)
  $$\phi_{12} = \sum_{j=0}^{2} \phi_j$$

The positive/negative sign applies to positive/negative values of $x_k$. This reconstruction is not always perfect and creates outliers which can affect regression results, especially for small $n_q$. I therefore drop reconstructed values outside the 95% percentile region for each $\hat{x}_k$ or $\hat{f}(\hat{x})$. 

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