Breiman’s “Two Cultures” Revisited and Reconciled

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Abstract—In a landmark paper published in 2001, Leo Breiman described the tense standoff between two cultures of data modeling: parametric statistical and algorithmic machine learning. The cultural division between the two dominant data modeling frameworks has been growing at an alarming rate in recent years. To make real progress, we are forced to consider the question: how do these two different cultures fit together? This paper introduces a new data analysis framework, called Integrated Statistical Learning (ISL) theory, which offers solutions to blend the “two cultures” into a coherent whole by establishing a link between them. Through examples, we describe the challenges and potential gains of this new integrated statistical thinking.

Index Terms—Integrated statistical learning theory, Exploratory machine learning, Uncertainty prediction machine, ML-powered modern applied statistics, Robustness.

1 INTRODUCTION

Some time around the year 2000 a split opened up in the world of statistics.” — Brad Efron (2020)

By the dawn of the 21st century, the statistics community was clearly divided into two distinct camps—namely, the (parametric) statistics and the (algorithmic) machine learning. A vivid description of the cultural polarization was described by Leo Breiman (2001) in his iconic essay on “Statistical Modeling: The Two Cultures.” But what has triggered this split?

ML>STAT: Breiman (2001) argued that algorithmic models are far more flexible, scalable, and accurate for complex big data problems. The traditional statistical methods, by contrast, are based on a priori assumed parametric models that are mainly suitable for small datasets.

STAT>ML: This algorithmic-supremacy standpoint had been furiously debated by several eminent statisticians in the discussion of the original paper. They found Breiman’s claim problematic on several counts. First, despite convincing prediction results, the absence of an explicit data-generating model (a probabilistic base) can make algorithmic methods less useful for scientific investigation; see the commentaries by Cox (2001) and Parzen (2001). Efron (2020) articulated this admirably: “Abandoning mathematical models comes close to abandoning the historic scientific goal of understanding nature.” Second, the black-box nature of algorithmic models makes them inscrutable, opaque, and not easily interpretable. In fact, regarding machine learning models, Breiman himself agreed that “their mechanism for producing a prediction is difficult to understand...So on interpretability, they rate an F.”

Where do we stand now? To a large extent, the cultural division between these two statistical learning frameworks has been growing over the past decades, especially in the post-deep-learning era. What options are we left with? 1) Let’s get rid of black-boxes and use parametric models, at the cost of accuracy and flexibility; 2) Let’s stick to model-free black-boxes that prioritizes predictive accuracy at the cost of interpretability and robustness.

What is a realistic way forward? Perhaps unsurprisingly, neither of these two extreme positions is tenable. To make real progress in developing next-generation ML technology, we are forced to consider the question: how do these ‘two cultures’ fit together into a coherent whole? Can we build a framework of data analysis that combines the best ideas from both sides and offer some way to make two into one? In this work, we take steps towards addressing these fundamental questions by introducing a new Integrated Statistical Learning (ISL) framework. Figure 1 shows a schematic description of blueprint of our research plan, which has five major components:

(1) Uncertainty Prediction Machine: The black-box machine learning (ML) algorithms need ways to assess their own confidence in their predictions. Without attaching some degree of uncertainty, we can’t ensure fail-safe decision making with predictive models. Unfortunately, conventional ML, which is a sophisticated curve-fitting exercise, only returns point-predictions. Naturally, the question arises: Given \((x_1, y_1), \ldots, (x_N, y_N)\) with \(x_i \in \mathbb{R}^p\) and \(y_i \in \mathbb{R}\), how can we design a simple and universal ‘interface’ that can convert any ‘first generation’ ML procedure into an uncertainty distribution prediction machine—one that is capable of predicting the underlying data generating mechanism in the form of conditional distribution \(f_{Y|X=x}(y)\). Section 2 presents the theoretical underpinnings, principles, and methods that enable such a design.

(2) Interpretable Learning: We like to distil an explicit and simple statistical model out of a black-box technique for easy interpretation. Section 2 lays out such a mechanism to allow algorithmic synthesis of probabilistic models with interpretable statistical parameters.
Now, once we observe \( X \) through the observation \( \mathcal{F}_X \), the uncertainty is updated through the observation \( X = x \), if the conditional distribution \( f_{Y|X=x}(y) \) differs from the initial \( f_Y(y) \). Therefore, one can measure the additional information gain on \( Y \) after observing \( X = x \) by examining the change in the probability distributions \( f_Y(y) \) and \( f_{Y|X=x}(y) \). Information theory provides an elegant framework for quantifying this “change.”

**Definition 1.** For a continuous response variable \( Y \), the conditional information density function between \( f_Y \) and \( f_{Y|X=x} \) is defined as

\[
d(u; F_Y, F_{Y|X=x}) := d_x(u) = \frac{f_{Y|X=x}(F_Y^{-1}(u))}{f_Y(F_Y^{-1}(u))}, \quad 0 < u < 1
\]

which satisfies: \( \int_0^1 d(u; F_Y, F_{Y|X=x}) \, du = 1 \). To improve readability, \( d(u; F_Y, F_{Y|X=x}) \) will be abbreviated as \( d_x(u) \), with the understanding that this compares marginal with conditional.

**Definition 2.** The Kullback-Leibler (KL) divergence from \( f_Y(y) \) to \( f_{Y|X=x}(y) \) is defined as

\[
KL(f_{Y|X=x}; f_Y) = \int f_Y(y) \log \frac{f_{Y|X=x}(y)}{f_Y(y)} \, dy.
\]

**Theorem 2.1.** The KL-divergence between \( f_Y(y) \) and \( f_{Y|X=x}(y) \) can be rewritten as KL-divergence between \( d_x(u) \) (1) and uniform density over \([0, 1] \):

\[
KL(f_{Y|X=x}; f_Y) = KL(d_x; 1_{[0,1]}).
\]

**Proof:** First substitute \( F_Y(y) = u \) in (2), and then apply the definition (1) to deduce:

\[
KL(d_x; 1_{[0,1]}) = \int_0^1 d_x(u) \log d_x(u) \, du.
\]

**Remark 1.** Theorem 2.1 formally implies that the deviation or proximity of \( d_x(u) \) from uniformity (see Fig. 12(c) and visually compare the blue curve with the black dotted uniform line) captures the amount of (predictive) information gained on \( Y \) by virtue of a priori knowing \( X = x \). For that reason, we call \( d_x(u) \) the “conditional information density.” Consequently, it is not surprising that \( d_x(u) \) will play a fundamental role in developing our statistical theory of prediction.

For each \( x_i \) in our dataset, we get a different \( d_{x_i} \) and a different KL-divergence value (4). Can we measure the overall predictive power of \( X \) by averaging all these KL-divergence values? To satisfactorily answer that, we have to introduce the concept of entropy.

**Definition 3.** Entropy is the most natural and fundamental measure of uncertainty. For a continuous random variable with distribution \( g \), it is defined as \( H(g) = \int g \log g \). Predictability-index can be defined as the difference between the entropy of \( Y \) and \( Y|X \), which measures how much of our uncertainty about \( Y \) decreased after observing \( X \). The following theorem establishes a beautiful connection between predictability-index and our conditional information density function.

**Theorem 2.2.** The average predictability of \( Y \) with respect to \( X \) (i.e., reduction in entropy) can be interpreted as how different \( d_x(u) \) is from uniform distribution on average:

\[
H(Y) - H(Y|X) = \mathbb{E}_X\left[KL(d_x; 1_{[0,1]})\right].
\]
solely in terms of \( d_x \), which distills the discrepancy (the “gap”) between the marginal \( f_Y \) and the conditional \( f_Y|X=x \). Accordingly, conditional information density (CID) \( d_x(u) \) acts as a natural starting point for developing a statistical theory of predictive modeling.

### 2.2 \( d \)-Modulated Conditional Density Representation

**Definition 4** (\( d \)-modulation). Any arbitrary conditional density can be expressed as

\[
    f_Y|X=x(y) = f_Y(y) \cdot d_x(F_Y(y)), \tag{6}
\]

where \( f_Y(y) \) is the marginal distribution of \( Y \) and \( d_x \) is defined in Eq. (1).

A few key points about this conditional density formula:
- This new representation of conditional density is universally valid and does not impose any distributional assumptions on the outcome (e.g., dispersion, symmetry, skewness, heavy-tailedness, etc.). Most importantly, it decouples the conditional density into two components: the marginal \( f_Y \) (the ‘pivot’ density) and the conditional information density \( d_x \) (which encapsulates all the essential knowledge for predicting \( Y \) from \( X = x \)).
- We call the density perturbation formula of Eq. (6) the \( d \)-modulation of marginal \( Y \). The function \( d_x \) filters out the directions by which the initial \( f_Y(y) \) departs most significantly from the true \( f_Y|X=x \). For that reason, we call it, alternatively, the contrast density function.

We now provide another interpretation of \( d_x(u) \) by introducing the notion of conditional probability integral transform (shortly, \( \text{cPIT} \)).

**Definition 5.** For continuous \( Y \sim F_Y \) and \( Y|X \sim F_Y|X=x \), the following random variable

\[
    U_x = F_Y(Y_x) \tag{7}
\]

is defined as conditional probability integral transform. Note that when \( F_Y = F_Y|X=x \), i.e., under homogeneity, \( U_x \) follows uniform distribution.

**Theorem 2.3.** \( d_x(u) \) is the distribution of \( F_Y(Y_x) \), where, recall, \( F_Y \) is the marginal cdf of \( Y \) and \( Y_x \) denotes the conditional random variable \( Y | X = x \).

**Proof:** \( D_x(u) = \Pr(F_Y(Y_x) \leq u) = F_Y|X=x(F_Y^{-1}(u)) \). Taking derivative we get the density:

\[
    D_x'(u) = d_x(u) = \frac{f_Y|X=x(F_Y^{-1}(u))}{f_Y(F_Y^{-1}(u))}, \quad 0 < u < 1. \tag{9}
\]

**Remark 2.** Definition 4 uses marginal distribution as a natural pivot for representing conditional density of \( Y \) given \( X = x \). However, one can choose any reasonable density function (parametric, nonparametric, etc.) as a pivot, as long as its support contains the support of \( f_Y|X=x \). The generalization of the formula (6) for arbitrary pivot \( f_0(y) \) is straightforward:

\[
    f_Y|X=x(y) = f_0(y) \cdot d_x^0(F_0(y)), \tag{8}
\]

where \( d_x^0(u) \) is the contrast density between \( F_0 \) and \( F_Y|X=x \); also see Supplementary A.5.

1. Support of a distribution is the region where probability density (or mass function) is strictly positive.

### 2.3 A Robust Learning Theory

It is evident from the decomposition (6) that we need to estimate \( d_x(F_Y(y)) \) to obtain the conditional density of \( Y \) given \( X = (X_1, \ldots, X_p) \). In this section, we describe a special technique for estimating \( d_x \) that enjoys the following desirable properties:

(a1.) Robust as well as efficient: One of the unsolved dilemmas is to construct learning algorithms that are simultaneously robust against noise and leaks very little, if any, efficiency under the standard (unperturbed) situation. Here we will present a new class of statistical learning methods to balance robustness and efficiency.

(a2.) Scalable to massive high-dimensional data: An important practical goal is to have a method that works for high dimensional problems with large \((n, p)\). We achieve this goal by developing a new class of machine learning powered nonparametric estimation method.

(a3.) Interpretable: It would be nice to have a tractable (and preferably smooth) model for the contrast density \( d_x(u) \) with a few interpretable parameters that can capture increasingly complex shapes of the conditional distributions.

Its a challenging task to reliably learn the “true” relationship between the input features \( X \) and output \( Y \) from big noisy datasets. To ensure that we succeed at it, our method has to be 'doubly-robust,' and thus tackle possible contamination in both response and covariates. We will approach the estimation of \( d_x \) in two stages: first, we will make it robust with respect to the outcome variable \( Y \), and then we will discuss defence strategies to guard against corrupted input \( X \).

**Y-Robustness.** Substituting \( u = F_Y(y) \) in the definition (1), allows us to represent the ratio \( f_Y|X=x(y)/f_Y(y) \) as a function of the rank-transform (i.e., probability integral transform) \( F_Y(Y) \) through \( d_x(F_Y(y)) \). This is the first major step towards ensuring the \( Y \)-robustness of our method. As a result, one can expand \( d_x(F_Y(y)) \) in the orthonormal basis of \( F_Y(y) \). One such orthonormal system is the LP-family of rank-polynomials (Mukhopadhyay and Parzen, 2020, Mukhopadhyay and Wang, 2020, Mukhopadhyay, 2017), which we denote \( \{T_j(y; F_Y)\} \) to emphasize that they are polynomials of \( F_Y(Y) \), hence extremely robust. As the true \( F_Y \) is unknown, we instead use the empirical LP-bases \( \{T_j(y; \hat{F}_Y)\} \) for our data analysis. Supplementary A.2 describes its construction and properties.

Summing up the above discussion, we have the following LP-series representation:

\[
    d_x(F_Y(y)) = 1 + \sum_j \text{LP}_{j|x} T_j(y; F_Y). \tag{9}
\]

The problem of estimating the function \( d_x \) now boils down to estimating the orthogonal coefficients \( \text{LP}_{j|x} \). To get an expression for the conditional LP-coefficients, first recall that

2. This ‘robustness-efficiency’ dilemma (also known as ‘rank-and-size’ duality) is not new. It is at least 60 years old unsettled issue of Statistics; see Box (1953) and Tukey (1960).
Multiply both sides of the equation (9) with \( \delta \) via Theorem 2.4 allows us to interpret the LP-coefficients
\[ \text{LP} \]
where \( \delta \) is the Kronecker delta function. Consequently, we multiply both sides of the equation (9) with \( T_j(y; F_Y) \cdot \hat{f}_Y(y) \) and then integrate to produce the following identity:
\[ \text{LP}_{j|x} = \int_{-\infty}^{\infty} \delta_x(\hat{f}_Y(y)) \cdot T_j(y; F_Y) \cdot f_Y(y) \, dy, \quad \text{for } j = 1, 2, \ldots \]
This immediately yields the following important theorem.

**Theorem 2.4.** The \( j \)th conditional LP-Fourier coefficient admits the following representation
\[ \text{LP}_{j|x} = \mathbb{E}[T_j(Y; F_Y) | X = x]. \]

Theorem 2.4 allows us to interpret the LP-coefficients \( \text{LP}_{j|x} \) (which are surely function of \( x \)) as a specialized regression function, where we regress \( T_j(y; F_Y) \) on \( X \). This is extremely handy, because one can now use any general machine learning method to estimate these coefficients.

**X-Robustness.** Nevertheless, the computational theory (12) is still not fully satisfactory, as corrupted (or adversarially perturbed) \( X \) can substantially influence the coefficients. To provide further defense, we need one last piece of idea, which is a little-known but fundamental fact about ‘regression via rank-transform.’ Also see Supplementary A.5.2.

**Theorem 2.5.** For any general random variable \( X \) (whether discrete or continuous) and a measurable function \( \Psi(\cdot) \), the following result holds
\[ \mathbb{E}[\Psi(X) | X] = \mathbb{E}[\Psi(X) | F_X(X)], \quad \text{with probability 1} \]

**Proof:** We start with a basic quantile mechanics fact: for any random variable \( X \) we have
\[ Q_X(F_X(X)) = X, \quad \text{with probability 1}. \]
where \( Q_X(u) = \inf \{F_X(x) \geq u\}, \quad 0 < u < 1 \) denotes the quantile function of \( X \). Accordingly, any function of \( X \), say \( h(X) \), can be equivalently rewritten as a function of the rank-transform \( F_X(X) \), since \( h(X) = \frac{h \circ Q_X(F_X(X))}{\text{prob}} \) with probability 1.

In plain language, this result proves the “sufficiency” of rank-transform (retaining all the information) for estimating conditional mean function. Therefore, we can estimate \( \text{LP}_{j|x} \) (12) by regressing \( T_j(Y; F_Y) \) on the LP-bases of \( X \). This will allow doubly-robust estimation of \( d_x \), without sacrificing an iota of efficiency in the process. Next, we will go over some examples, to demonstrate how this theory actually translates into practice (algorithm).

**Example 1.** The Butterfly data. The butterfly data superposition of \( N = 350 \) samples, generated from \( y = x + \epsilon \) and \( y = -x + \epsilon \), with \( \epsilon \sim \mathcal{N}(0, 1) \); see Supp. Fig. 11. The scatter plots of \( T_j(y; F_Y) \) against \( u = F_X(x) \) for \( j = 1, \ldots, 4 \) are displayed in Fig. 2. We refer these graphs as LP-scatter plots. The red curves denote kkn-estimated regression curves. In the \( \mathbb{R} \) computing language, this is simply \( \text{knn}(T_j(y; F_Y)) \sim T_X \), \( j = 1, \ldots, 4 \) where \( T_k(x; F_X) \) is the \( k \)th col of LP-transformed feature matrix \( T_X \). Of course, instead of kkn method, one can choose any machine learning routine; the next example uses gradient boosting algorithm (gbm). To get the estimated conditional LP-coefficients at \( x_0 = 2 \) (indicated by the blue dots) we record the predicted values at \( u_0 = F_X(x_0) \): \( \text{LP}(x_0) = -0.20, \text{LP}(x_0) = -0.47 \), and the rest are practically zero. Substituting them into (9) yields the estimated contrast function \( \hat{d}_{x_0} \), as was shown in Fig. 12(c).

**Example 2.** BUPA liver disorders data. This is a popular dataset (available in the UCI ML repository) used by many researchers, including Breiman (2001). It consists of measurements of gamma-glutamyl transpeptidase (GGT) and alanine-aminotransferase (ALT) extracted from \( N = 345 \) male individuals’ blood samples. High levels of ALT and GGT indicate a damaged liver. Fig. 3 (a) shows the scatter plot of \( y = \log(\text{ALT}) \) versus \( x = \log(\text{GGT}) \). We seek to predict the distribution of \( \log(\text{ALT}) \) levels given \( x = 3.5 \); see the red triangle in panel (a) of Fig. 3.

Our data analysis scheme can be simply summarized as follows:
1) We start by modeling the conditional mean function using gradient-boosting algorithm. The rugged blue line denotes the estimated regression curve $\hat{\mu}(x)$. 

2) To estimate the conditional density at $x_0 = 3.5$, we choose $\mathcal{N}(\hat{\mu}(x_0), \sigma^2_y)$ as our pivot model, where $\hat{\mu}(x_0) = 3.35$ and $\sigma_y = 0.5$.

3) Following theorems 4 and 5, we estimate the conditional LP-Fourier coefficients $\hat{L}_{j|}\hat{\mu}_{j|x_0}$ and the associated contrast density function: $\hat{\sigma}_{x_0}^2(F_0(y)) \approx 1 - 0.72 T_2(y; F_0) + 0.10 T_3(y; F_0)$. This is shown in the middle panel of Fig. 3.

4) Interestingly, we can interpret the coefficients of $\hat{\sigma}_{x_0}$: the large value of $\hat{L}_{2|}\hat{\mu}_{2|x_0}$ indicates that the selected pivot needs scale-correction ("second-order" correction) to get close to the true conditional density at $x_0$. In addition, the negative sign implies the variability has to be reduced, which is vividly apparent from Fig. 3(c).

5) Similarly, the presence of third-order (positive) LP-coefficient indicates the symmetric normal pivot needs to be tilted to the right. The positive skewness of $f_{\hat{\sigma}}(y)$ is probably justified due to the presence of few large $y$-values around $x_0 = 3.5$, as indicated by squares.

6) Finally, the shape of $\hat{d}_x$ provides quick exploratory guidance for choosing appropriate families of parametric conditional distributions. For the BUPA liver-disorders data, one could select skew-normal or asymmetric logistic distribution (Friedman, 2020) as a model for $Y|X$.

### 2.4 The Basic Algorithm: ISL

We are now in a position to provide the blueprint of the algorithmic interface in Fig. 1 that converts a user-selected machine learning method into an integrated statistical learning machine (ISL) in a robust and interpretable manner. The theory in the previous section, makes it absolutely straightforward and extremely easy to implement, which essentially involves three steps. The following algorithm presents a flowchart of the core computational steps in pseudo-code, so that one can implement it using any programming language or ML-platform. We have found that the H2O open-source (see https://github.com/h2oai) is extremely efficient. But users can choose any ML environment (e.g., TensorFlow, Azure, AutoGloun, etc.) to operationalize the following workflow.

**ISL: A flowchart of the core computational steps in pseudo-code**

**Input Data** $(X_i, y_i)$ for $i = 1, \ldots, N$; user-selected machine learning method (ML); values of $m_y$ and $m_x$. And the target test-point $x_0 \in \mathbb{R}$.

**Step 1.** $T_Y \leftarrow \text{LP.basis}(y, m_y); \quad T_X \leftarrow \text{LP.basis}(X, m_x)$

for $(j = 1, \ldots, m_y)$

**Step 2.** $\text{Fitted.ML}_j \leftarrow \text{ML}(T_Y[, \ j] \sim T_X)$

**Step 3.** $\hat{L}_{j|x_0} \leftarrow \text{predict}($Fitted.ML$_j, x_0)$

**Return** $\hat{d}_{x_0} \leftarrow 1 + \sum_j \hat{L}_{j|x_0} T_Y[, \ j]$

The LP-transformation is implemented by the function $\text{LP.basis}; T_X$ is the stacked matrix $[T_{X_1}, T_{X_2}, \ldots, T_{X_6}]$, where the $j$th column of the matrix $T_{X_i}$ is simply $T_{j|x} (x; F_i)$. Step 1 makes the whole procedure robust, non-linear, and invariant under monotone transformations. Step 2 performs the training of the ML method and step 3 predicts the fitted value at the desired $X = x_0$. Based on our experience, $m_x = 4$ and $m_y = 6$ work reasonably well for most problems, including all the datasets analyzed in this paper. Finally, one can generate the uncertainty distribution $f_{Y|x_0}$ by applying the $d$-modulation formula of equation (6).

**Remark 3.** In this section we have showed: how to actualize the vision sketched in Fig. 1; how to compress-down large complicated algorithmic methods into concise parametric forms with few interpretable coefficients; how to derive a smoothness-promoting model for the response $Y$ from black-box prediction methods; and how to robustify algorithmic techniques. We have achieved each of these ends by devising a generic mechanism that holds for any machine learning method. Furthermore, Section 4 shows how these same modeling principles can unify a wide range of traditional and novel statistical methods.

### 2.5 d-Kernel Smoothing and Importance Weighting

In many applications, we are often interested in estimating quantities like $\mathbb{E}[\Psi(Y)|X = x]$ for general $\Psi(\cdot)$ function. This can be rewritten as

$$\mathbb{E}[\Psi(Y)|X = x] = \int_y \Psi(y) f_{Y|x = x}(y) \, dy$$

by applying the formula (6). This leads to the following theorem, which expresses the (local) conditional mean as a weighted average of the global samples $\{y_1, \ldots, y_N\}$.

**Theorem 2.6 (d-kernel smoothing).** Conditional expectation of $\Psi(Y)$ given $X = x$ can alternatively be expressed as:

$$\mathbb{E}[\Psi(Y)|X = x] = \mathbb{E}[\Psi(Y) \cdot d_x(F_Y(y))],$$

the rhs expectation is taken over the marginal distribution of $Y$.

**Remark 4.** The above representation implies the following $d$-weighted estimator:

$$\hat{\mathbb{E}}[\Psi(Y)|X = x] = \frac{1}{N} \sum_{i=1}^{N} w_i(x) \Psi(y_i),$$

(15)

where the empirical weighting function $w_i(x) = \hat{d}_x(F_Y(y_i))$. The formula (15) can be interpreted as a

4. This materialize the objective of “Trend 1” as proposed by Efron (2020): “Trend 1 aims to make the output of a prediction algorithm more interpretable, that is, more like the output of traditional statistical methods.”

5. Friedman (2020) advocates strongly for this.
specially-designed kernel smoothing, which could be extremely handy in some cases. Three particular examples are given below:
1) $\Psi(Y) = \mathbb{I}(Y > k)$ gives the conditional probability $\Pr(Y > k|X = x)$; This can be used to construct different risk measures based on tail events;
2) $\Psi(Y) = Y$ recovers the conditional mean $\mathbb{E}[Y|X = x]$;
3) $\Psi(Y) = (Y - \mathbb{E}(Y))^2$ yields conditional variance $\text{Var}(Y|X = x)$, whose square-root, i.e., the standard error of $Y_{|x}$, is often used as an uncertainty quantification measure for a prediction. However, we should point out that standard error is not always a prudent choice for quantifying reliability of a prediction, specially when the conditional distribution is heavy-tailed, or skewed, or multi-modal; see Section 4.6.

3 Exploratory Learning Machine

Our goal here is to develop some graphical diagnostic methods to make a transition from a black-box answer machine to an exploratory learning machine that can drill deeper and generate previously unanticipated, interesting questions to facilitate data-driven scientific discovery and understanding. This can be viewed as a pre-deployment “health check-up” for statistical learning methods that comes with a set of nonparametric exploratory tests.

3.1 Heterogeneity Component Analysis

Heterogeneity component analysis (HCA) is a technique for identifying dynamic components of $Y_{|x}$ – the aspects of the conditional distribution $f_{Y|X=x}(y)$ (such as location, scale, skewness, tail, etc.) that vary with covariates. We illustrate the main idea using the butterfly example. The procedure starts with LP-scatter plots, as was shown in Fig. 2. The first scatter plot captures how the location of $f_{Y|X=x}(y)$ (conditional mean) changes with $x$; the second one captures the change of scale (the nature of heteroscedasticity); the third one provides information on the changing skewness and so on. Thus, an obvious approach for detecting the heterogeneity components would be to:

Step 1. Compute a goodness-of-fit statistic for the $j$th LP-scatter plot by its coefficient of multiple correlation: $R_j^2 =$Proportion of variance in $T_j(y; F_Y)$ that is explained by $\{T_1(x; F_X), \ldots, T_m(x; F_X)\}$.

Step 2. Assess the significance of the $R_j^2$ by constructing F-statistic for the $j$th LP-scatter

$$F_j = \frac{R_j^2}{1 - R_j^2} \times \frac{N - m - 1}{m}$$

whose null distribution is $F_{m,N-m-1}$. Fig. 12(a) displays the significant (whose p-value < 0.05) heterogeneity components for the butterfly data.

HCA Diagnostic Plot. For real data analysis, we apply the above method on the residuals to determine what structure of the data (relationship between $y$ and $x$) remains to be described after the ML-regression. The magnitude of the heterogeneity components reveals which shape parameters of the residuals are changing with $x$. If all the components of HCA are small, then it confirms to the practitioner that errors are already homogeneous (we call this scenario: residual “flatland”) with respect to the covariates and nothing left to be modeled.

Example 3. Bone mineral density data (Bachrach et al., 1999). It contains measurements on the relative change in spinal bone density (over one year period) of $N = 485$ North American adolescents, as a function of age. This dataset was previously analyzed by Hastie et al. (2009, p. 152). Fig. 4(a) shows the data along with the estimated regression smoother. We like to understand whether there is any “excess” heterogeneity beyond the 1st-order regression modeling. To check the homogeneity of the residuals with respect to the covariate $x$, we apply our HCA procedure on the residuals $y_i - \hat{y}_i$. The result is shown in the top row of Fig. 4, which says that the regression model has not taken into account the heteroscedasticity of the data (furthermore, there is some evidence of varying skewness in the conditional distribution). These unmodelled (dynamic) heterogeneity components should be modeled to improve the prediction. The heterogeneity is partly due to the presence of both male and female youngsters’ in the dataset.

Example 4. Online news popularity data (Fernandes et al., 2015). The objective of this study is to predict the popularity of online news. The data consists of $N = 39,644$ articles,

6. The article in row # 31,038 seems to be an outlier (in fact it is an erroneous data: check its 4th and 5th feature values–rate of unique and non-stop words–which can’t be more than 1). Due to the doubly-robust property, our method can handle this easily. But standard machine learning methods might get unduly influenced by this one data point (article); see Fig. 17 in the Supplementary.
and for each article we have $y$ (log$_{10}$ of number of shares in social networks, a measure of popularity) and $p = 59$ extracted features (such as number of words in the title, number of links, number of images, etc.) $x$. This dataset was recently analyzed by Friedman (2020).

We start by fitting gradient boosting regression to the data. At this point, it is a natural question to ask whether the gbm captured most of the important information in the data. If not, what is missing? To spot the unmodelled dynamic components, we start from the residuals, as shown in Fig. 4(c). The HCA diagnostic plot of panel (d) is constructed based on the Lasso-selected features, which clearly indicates the presence of a strong heteroscedasticity. However, it is also true that the conditional distribution will be asymmetric and long-tailed, but our analysis shows that they are not swiftly changing with covariates; they are more or less static in nature, not dynamically evolving shape-parameters. This further corroborates the findings of Friedman (2020, Sec. 6.2).

3.2 Pivot Uncertainty Modeling

The parameters LP$_{ij|x}$ of the contrast density $d_x$ can be viewed as “generalized coordinates” that describe the shape of $f_{Y|x=x}(y)$ relative to some reference distribution (pivot) $f_0(y)$.

In fact, these shape-coefficients LP$_{ij|x}$ capture the whole dynamics of how the conditional distribution $f_{Y|x=x}(y)$ evolves with $x$. Naturally, a large value of qPivot($x$) $= \sum_j |LP_{ij|x}|^2$ (16) indicates higher uncertainty of the pivot $f_0$ for modeling the conditional $f_{Y|x=x}$. The ‘q’ in qPivot stands for quantification of lack-of-fit. The above formula can also be interpreted as measuring the deviation of $d_x(u)$ from uniformity (follows from Parseval’s identity):

$$\int_0^1 (d_x(u) - 1)^2 du = \sum_j |LP_{ij|x}|^2$$ (17)

Example 5. Movie box-office revenue data (Voudouris et al., 2012). The goal of this study is to build a forecasting model for film revenues. The data contain $N = 4031$ pairs of observations $(x_i, y_i)$ on the 1990s film data, where $x_i$ is the log of opening box-office revenues and $y_i$ is the log of box office revenues after the first week. We are interested in predicting the distribution of the outcome $y$ for $x = 12$ and 14-marked with the blue and green dotted line in Supp. Fig. 18 (a). Our starting pivot $f_0 = N(14, \sigma_y = 3.25)$ is shown in the panel (b). To check whether the conditional density at $x$ follows this presumed parametric law, we estimate the respective contrast densities; see the middle panel of Supp. Fig. 18. The bimodality of $d_{x=12}$ indicates that the Gaussian pivot failed to capture the presence of films with two kinds of box-office earnings. This is also reflected in the pattern of the scatter plot, which is divided into two branches. On the other hand, the inverted ‘U’ shape (quadratic) of $d_{x=14}$ implies that the variability of the Gaussian pivot needs to be adjusted (2nd-order correction) to go from $f_0(y) \rightarrow f_{Y|x=14}(y)$. The final refined conditional density models at $x = 12$ and 14 are shown in the bottom row of Supp. Fig. 18. They are estimated using the $d$-modulation technique, described in Sec. 2.2.

3.3 Goodness-of-fit Diagnostics

The flexibility of our approach (as depicted in Fig. 1) enables us to choose any machine learning method to design the integrated learning machine. Nonetheless, a user may want to know which ML-powered distribution prediction algorithm(s) better explains the pattern in the data. To answer that here we introduce a graphical exploratory procedure to evaluate the overall goodness-of-fit of the method. Deploying statistical models without knowing how well it actually fits the data, is a statistical sin. We illustrate our idea using the butterfly data.

ML-driven ISL system. To construct an ISL for the butterfly data, we start with three base learners: k-nearest neighbors (knn), random forest (RF), and gradient-boosting algorithm (GBM). We train each model using the generic ISL-procedure described in section 2.4. To check whether these models are congruent with the observed data, we proceed as follows:

Step 1. Compute the generalized quantile-residuals for all three models on a hold-out validation dataset (separate from the training samples) of size $n_v$.

$$U_{\ell i} = \hat{F}_{Y|x=x}(y_i), \quad \text{for } i = 1, \ldots, n_v; \quad \text{and } \ell = 1, 2, 3,$$ (18)

Note that if the model accurately captures the pattern in the validation data, then we expect the distribution of the $U_{\ell i}$’s to be close to uniform.

Step 2. Display the histograms and QQ-plot of $\{U_{\ell 1}, \ldots, U_{n_v}\}$. This is done in Fig. 19 for the butterfly data, which shows the knn model satisfactorily captures the observed reality.

Step 3. To quantify the model-fit we propose the following test statistic for the $\ell$-th model:

$$qDIV_\ell = \sum_{j=1}^m \left| n_v^{-1} \sum_{i=1}^n \text{Log}_j (U_{\ell i}) \right|^2,$$ (19)

where Log$_j$ denotes the $j$th orthonormalized shifted Legendre polynomial on unit interval. The qDIV statistic measures the distance between Uniform[0,1] and the sample distribution of $U$. We have used $m = 6$ in all our examples. One can also compute pvalues using the $\chi^2$ null distribution of qDIV. For more details see Mukhopadhyay and Wang (2020).

Step 4. It is also interesting to note that one can directly compute these generalized quantile-residuals from the contrast distribution function, since (by substituting $F_0(t) = u$):

$$F_{Y|x=x}(y) = \int_{-\infty}^y f_0(t) \, d_x(F_0(t)) \, dt = \int_0^y d_x(u) \, du = D_x(F_0(y)).$$
For the butterfly data qDIV values (with p-values in the parentheses) for knn, random forest, and gbm respectively are 0.0466 (0.82), 1.12 (1.4 × 10^{-10}), and 0.386 (0.0026). These statistic values can be used as a general measure of “lack-of-fit” for comparing (ranking or even tuning hyperparameters) the performances of different ML-engine based ISL models.

## 4 Some Connections and Unification

The theoretical constructs of our integrated learning framework are not based on an isolated trick, but are fundamental to statistics. Our formalism and principles of data modeling unify many traditional and modern statistical methods. Its significance on the practice and pedagogy of modern applied statistics is described in Supplementary A.6. In this section, we will report some surprising findings to highlight this fact.

### 4.1 K-sample Comparison Problems

Given random samples from \( k \) distributions \( F_i(y), i = 1, \ldots, k \) we seek to test the following null hypothesis:

\[
H_0 : F_1(y) = \cdots = F_k(y) \quad \text{for all } y,
\]  

with the alternative hypothesis that at least two distributions are different. Before attacking this general problem of equality of distributions, we start with a simpler problem of testing the equality of means. The Kruskal–Wallis statistic provides a nonparametric test for it.

**Definition and notation.** We have \( k \) mutually independent random samples with sizes \( n_1, \ldots, n_k \) with the combined sample size \( N \); \( R_i \) denotes the rank of \( Y_i \) in the pooled samples; \( G_\ell \) is the set of indices for the \( \ell \)-th group. The Kruskal–Wallis test statistic is defined as:

\[
KW = \frac{12}{N(N+1)} \sum_{\ell=1}^{k} n_\ell \left( \frac{1}{n_\ell} \sum_{i \in G_\ell} R_i - \frac{N+1}{2} \right)^2. \tag{20}
\]

One may go one step further and ask how to test the equality of scale parameters (variability over different groups), which can be tested using Mood statistic, defined as

\[
\text{Mood} = \frac{180}{N(N+1)(N^2-4)} \sum_{\ell=1}^{k} n_\ell \left( \frac{1}{n_\ell} \sum_{i \in G_\ell} \left( R_i - \frac{N+1}{2} \right)^2 - \frac{N^2-1}{12} \right)^2. \tag{22}
\]

For testing high-order differences like skewness (not to mention kurtosis, which will take almost two full lines to write), the formula of the test statistic gets even more complicated.

\[
\text{SKEW} = \frac{7}{N(N+1)(N^2-4)(N^2-9)} \sum_{\ell=1}^{k} n_\ell \left( \frac{20}{n_\ell} \sum_{i \in G_\ell} \left( R_i - \frac{N+1}{2} \right)^3 - \frac{3N^2-7}{n_\ell} \sum_{i \in G_\ell} \left( R_i - \frac{N+1}{2} \right) \right)^2 \tag{23}
\]

A practitioner at this point might wonder how to make sense of these monstrous formulae? Is any intuitive or systematic derivation possible? The answer is yes. We just have to look at the problem from a different perspective. A modern introduction to the \( k \)-sample comparison problem is given below.

Step 1. View it as a \( (X, Y) \) problem: Define \( X \) to be the group-index variable taking values \( \{1, \ldots, k\} \). We now have \( \{(x_i, y_i)\}_{i=1}^{N} \) data, which can be displayed as a scatter plot.

Step 2. Reformulate \( H_0 \): The \( k \)-sample null-hypothesis of equality of distributions can now be viewed as a test of the homogeneity problem (see Section 3.1) \( f_{Y|X=\ell}(y) = f_Y(y) \) for \( \ell = 1, \ldots, k \). Now since

\[
f_{Y|X=\ell}(y) = f_Y(y) \left\{ 1 + \sum_{j} \text{LP}_{j|\ell} T_{j}(y; F_Y) \right\}, \tag{24}
\]

the original \( k \)-sample hypothesis (20) can alternatively be rewritten as

\[
H_0' : \text{LP}_{j|\ell} = 0, \quad \text{for all } j.
\]

Hence, intuitively, it makes sense to apply the HCA diagnostic test of section 3.1 on the \( (x, y) \) data. Doing so we get almost all of the known results of \( k \)-sample modeling in one-shot. To justify the statement, we start with the following theorem.

**Theorem 4.1.** The Kruskal–Wallis statistics can be expressed as

\[
KW = \sum_{\ell=1}^{k} n_\ell \left| \text{LP}_{1|\ell} \right|^2 = NR_1^2. \quad \text{change in location}
\]  

where the first-order LP-multiple correlation \( R_1^2 \) is defined in the step 1 of Section 3.1.

The proof is in Supplementary A.3. The beauty of this result is that it seamlessly generalizes to high-order cases (proofs are tedious but not difficult and are left to the reader):

\[
\text{MOOD} = \sum_{\ell=1}^{k} n_\ell \left| \text{LP}_{2|\ell} \right|^2 = NR_2^2; \quad \text{change in scale}
\]  

\[
\text{SKEW} = \sum_{\ell=1}^{k} n_\ell \left| \text{LP}_{3|\ell} \right|^2 = NR_3^2; \quad \text{change in skewness}
\]  

\[
\text{KURT} = \sum_{\ell=1}^{k} n_\ell \left| \text{LP}_{4|\ell} \right|^2 = NR_4^2, \quad \text{change in tail}
\]

and so on. The practical benefit of these results is that they provide a systematic and comprehensive \( k \)-sample analysis program that is radically simple to implement, since LP-multiple correlations \( R_j^2 \) can be computed in a single line R-code: `summary(lm(T_j(y) ~ T_X)$r.squared).

**Example 6.** LDL cholesterol of Quail. This is a randomized experiment investigating LDL (low-density lipoprotein) cholesterol in quails; \( N = 39 \) quails were randomly assigned to \( k = 4 \) diets for a specific period of time, where each diet mixed with a different drug compound. The boxplot is shown in Fig. 5. The scientific question is: whether or not different drug compounds change the LDL

```r
# Sample code for LDL cholesterol analysis
library(supplementary)

# Data preparation
quail_data <- read.csv("quail_data.csv")

# Groupings for different diets
diet_groups <- factor(quail_data$diet)

# Calculating sample statistics
sample_stats <- summary(diet_groups)

# For each diet group:
for (diet in levels(diet_groups)) {  
  # Compute sample mean
  mean_LDL <- mean(quail_data$LDL[diet_groups == diet])
  # Compute sample variance
  var_LDL <- var(quail_data$LDL[diet_groups == diet])
  # Output results
  cat("Diabetes Diet", diet, "Mean LDL: ", mean_LDL, " Variance: ", var_LDL, "\n")
}
```

The practical benefit of these results is that they provide a systematic and comprehensive \( k \)-sample analysis program that is radically simple to implement, since LP-multiple correlations \( R_j^2 \) can be computed in a single line R-code: `summary(lm(T_j(y) ~ T_X)$r.squared).`
cholesterol levels. The right panel of Fig. 5 shows the re-scaled HCA-plot, where we multiply the $j$th LP-multiplet correlations $R_j^2$ by the sample size $N$. It shows that the distribution of LDL cholesterol levels is changing in scale and location.

**Remark 5.** For this data, the classical anova F-test yields $p$-value 0.345, thus fails to detect any significant location differences—clearly contradicting the boxplots. Why has the F-test failed? There are two primary reasons for that: (i) the highly non-Gaussian nature of the distribution of $Y$ along with a few outlying values were enough to kill the parametric F-test; and (ii) the assumption of equal variance is also seems inaccurate. The takeaway: we need both robustness and efficiency to reliably detect high-order distributional differences.

### 4.2 Probabilistic Index Model

Do baseball players gain weight as they get older? Can a larger dose of an antipsychotic drug reduce the severity of depression (after adjusting for the confounders)? All of these problems are concerned with modeling the *comparison probability regression:* $CPR(x) = Pr(Y \leq Y[x])$, as a function of $x$, instead of as a usual conditional mean regression. $CPR(x)$ encodes the probability that a randomly selected subject from the population with $X = x$ has a higher response value than a randomly selected subject from the full data. This concept was pioneered by Olivier Thas under the name of probabilistic index model; see Thas et al. (2012).

To understand how $CPR(x)$ is related to our framework, we start with the following integral representation:

$$Pr(Y \leq Y[x]) = \int F_Y(y) \, dF_Y|X=x(y). \quad (26)$$

This leads to the following important result.

**Theorem 4.2.** $Pr(Y \leq Y[x])$ can be expressed as a regression of probability integral transform $F_Y(Y)$ on $X$:

$$Pr(Y \leq Y[x]) = E[F_Y(Y)|X = x] = \frac{1}{\sqrt{12}} LP_{1|x} + 1/2. \quad (27)$$

The first equality follows from (26), and the second equality follows from supplementary equation (E5.4). The representation (27) justifies the name comparison probability regression, which is simply a standardized 1st-order LP-regression function. As a consequence of Theorem 4.2, the whole procedure can be implemented in one line of R-code: $lm(T_1(y) \sim T_X)$.

**Example 7.** *Baseball data* (Matloff, 2017). Fig. 6 shows the age versus weight scatter plot of $N = 1015$ major-league baseball players. We would like to investigate whether baseball players gain weight as they age. In other words, our task is to check whether $Pr(Y \leq Y_{[x]}$) increases as a function of age or not. We proceed as follows:

Step 1. Compute the LP-basis functions: $T_1(y; \tilde{F}_Y)$ for the response and $T_X = \{T_1(x; \tilde{F}_X), \ldots, T_m(x; \tilde{F}_X)\}$ for the predictor variable. Here we have picked $m = 4$.

Step 2. (penalized) LP-regression: Perform the multiple linear regression $T_1(y; \tilde{F}_Y)$ on $T_X$. Return the Bayesian information criterion (BIC) selected model. If no variable is selected then it supports the null hypothesis $H_0 : Pr(Y \leq Y_{[x]} = 0.5$. For the baseball data we get:

$$\hat{LP}_{1|x} = E[T_1(y; \tilde{F}_Y)|X = x] = 0.17 T_1(x; \tilde{F}_X). \quad (28)$$

There is no intercept as $E[T_1(y; \tilde{F}_Y)] = 0$, by construction. Although athletes strive to keep physically fit, eq. (28) seems to suggest that the baseball players do gain some weight (in a slow linear rate) over time. Fig. 6(b) makes this somewhat clear to understand how.

### 4.3 Shape Predictors: Generalized Feature Selection

When dealing with a large number of covariates, it is often important to understand which are the most *predictive* for the response variable $Y$. Currently, the most popular feature selection method is $L_1$-penalized lasso regression (Hastie et al., 2009, Ch. 3.4), which minimizes the residual sum of squares $\|y-X\beta\|^2_2$ subject to $\|eta\|_1 \leq \lambda$. This will be denoted algorithmically as lasso($y \sim X$). It is important to note that, when we perform lasso-regression of $y$ on $X$, we only get ‘first-order’ informative features—the variables that influence the conditional mean $E[Y|X = x]$. Naturally, the question is: can we go beyond mean, and find the most relevant attributes to describe the *shape* of the conditional distribution $f_{Y|X=x}(y)$? Can we better utilize the lasso-technique to find those “shape predictors”? The following
is an attempt to construct such a generalized feature selection program, which is extremely easy to implement. The algorithmic logic proceeds as follows:

Step 1. Let’s start by recalling our basic model: $f_Y|X=x(y) = f_Y(y) \cdot d_x(F_Y(y))$. The covariates $x = (x_1, \ldots, x_p)$ can influence the distribution of the response $Y$ only through $d_x \circ F_Y(y) = 1 + \sum \text{LP}_j |x \cdot T_j(y; F_Y)$, which is governed by the following system of equations:

$$
\text{LP}_1[x] = \mathbb{E}[T_1(Y; F_Y)|X = x] \\
\text{LP}_2[x] = \mathbb{E}[T_2(Y; F_Y)|X = x] \\
\vdots
$$

The first equation captures the dynamics of conditional mean as a function of $x$, while the second one describes how the variability (scale) changes with $x$, and so on.

Step 2. Thus, it is quite obvious that one can extract the $j$th order features, denoted by $O_j = \{ \ell : \beta^{(j)}_\ell \neq 0 \}$, where $\beta^{(j)}$ is the output of lasso($T_j(y) \sim X$), which efficiently computes the solution of

$$
\hat{\beta}^{(j)} = \arg \min_{\beta^{(j)} \in \mathbb{R}} \left\{ \| T_j(y) - X \beta^{(j)} \|^2_2 + \lambda \| \beta^{(j)} \|_1 \right\}. \tag{29}
$$

Accordingly, $O_1$ is the collection of location-informative variables, $O_2$ contains all the scale-informative ones, $O_3$ variables are responsible for predicting change in skewness of $Y$, etc.

Step 3. To make the analysis doubly-robust and to allow for non-linearity of the features, simply substitute $Y$, accordingly, we use $k = 1$ and $j = 3$ in the formula (30). The top 10 features are displayed in Fig. 7, which shows the nature of contributions of different variables. For example, the variable ‘is_weekend’ (whether the news article was published on the weekend or not) plays a dual role of being both a location and scale invariant. Consequently, our multi-layered robust methodology provides a refined understanding of the “which and how” aspects of feature selection.

Finally, one can perform targeted feature screening. An investigator can specifically query which variables are mostly responsible for heteroscedasticity, or change in symmetry, etc. Table 1 shows the top five features in each category: location, scale, and skewness. These insights will ultimately help the applied researchers to better understand the nature of the association between response $Y$ and $X = (X_1, \ldots, X_p)$.

### 4.4 The XYZ Problem: Distributional Impact Analysis

In application fields such as healthcare, economics, and social sciences, data often arise in $XYZ$ format, where $Z$ is the binary treatment variable, $Y$ is the response variable, and $X$ is the (possibly large) collection of covariates. One such problem is discussed below.

**Example 9.** Rosner’s FEV data (Rosner, 1995). Table 2 (in Supplementary) describes the data. The main interest lies in understanding the impact of smoking on the distribution of FEV as a function of age. This is important for individualized custom-tailored decision-making, where a treatment might be locally effective for a sub-population (characterized by certain values of $x$) without being globally effective for the whole heterogeneous population.

---

8. see Supplementary A.5.4. for a related discussion on ‘robust lasso.’
most impacted by the treatment and how they are impacted. For example, at age 14 variability is the dominant mode of difference, whereas at age 15, location is the main source of difference. Overall, the DIF barplot does a modest job of capturing the heterogeneous impact of teenage smoking on lung function across different age groups; see Remark A.5.5 in the Supplementary.

4.5 Quantile Regression

Quantile regression has become a pervasive tool in a host of real-world application areas. Despite the great progress made in the last 40 years (Koenker, 2017), some practical challenges still remain: how to develop a nonparametric quantile regression method that can scale to high-dimensional problems? How to efficiently incorporate non-linearity? How to ensure that the estimated conditional quantile curves will not cross each other? Our statistical learning theory can offer some realistic solutions to these frontier problems of quantile regression. But before getting into that, let us start with a slightly broader context. Our integrated statistical learning architecture (as depicted in Fig. 1) has two notable consequences for the practice of nonparametric data modeling:

- Our technology opens up brand-new ways of building “ML-powered” statistical models that are simultaneously flexible and scalable for large \((n, p)\) problems, where classical nonparametric methods are not a viable option.
- Our ISL learning architecture is designed using a high-level universal language that is agnostic to the specific ML method, to ensure the portability and durability of the technology.

To perform ISL-based quantile regression, simply extract the appropriate percentiles from the conditional density estimate \(f_{Y|X=x}(y)\). The beauty of our method is that we don’t have to develop theories specialized to a certain kind of machine learning method like random forest (Meinshausen, 2006) or neural network (Cannon, 2011)—it’s all done using a single generic procedure; see Supplementary A.5.3 and Fig. 16 for more details.

Example 10. Dutch Boys data (Fredricks et al., 2000). This dataset is a part of the Fourth Dutch Growth Study, which comprised of observations on age and BMI of \(N = 7294\) Dutch boys. The goal is to estimate age-specific reference growth curves based on conditional quantile function \(Q_{Y|X=x}(u)\). Supp. Fig 20 displays our result where we have used lasso (as implemented in the glmnet R-package) as the base learner inside ISL engine. Following the current World Health Organisation (WHO) recommendation, we have used \(u = (.03, .15, .50, .85, .97)\) for constructing the cross-sectional reference growth charts; see Remark A.5.6 in the Supplementary.

4.6 Prediction Interval Estimator

One way to summarize the conditional density is through prediction interval (PI), which provides a concise view of the most likely values of the response variable\(^{10}\). For a specified

\[\text{ML-powered}\]

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- Our ISL learning architecture is designed using a high-level universal language that is agnostic to the specific ML method, to ensure the portability and durability of the technology.

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10. Knowing the range of values of \(Y\) as opposed to a single-point estimate, helps decision-makers choose proper action by carefully evaluating the degree of uncertainty (length of the PIs).
level \( \alpha \), the goal is to construct an interval that covers no less than \( (1 - \alpha) \) of the probability mass of \( f_{Y|X=x}(y) \). How to construct such PIs? We discuss three different procedures. The real challenge is to make it as narrow as possible, while maintaining the desired coverage. • Quantile-based PI. As noted by Meinshausen (2006), one can use quantile regression to build PIs. A 100(1-\( \alpha \))% prediction interval for \( Y \) given \( X = x \) can be estimated as

\[
\text{qPI}_{1-\alpha}(x) = \left[ Q_{Y|X=x}(\alpha/2), Q_{Y|X=x}(1-\alpha/2) \right].
\]

Naturally, the width (precision) of the quantile-PIs vary over the covariate space.

• Standard-error-based PI. If we are brave enough to assume some parametric form of the conditional density, then we can greatly simplify the expression for the PIs. For example, under Gaussianity assumption, one can express (34) in the following compact form, just involving conditional mean \( \mu(x) \) and standard-deviation:

\[
\text{gPI}_{1-\alpha}(x) = \left[ \mu(x) \pm z^{(\alpha/2)} \sigma_{\|x} \right]
\]

where \( \Phi^{-1}(\alpha) = z^{(\alpha)} \), \( \text{Var}(Y|X = x) = \sigma_{\|x}^2 \), and ‘g’ denotes the Gaussian-PI.

• Highest-density PI. This is the shortest possible interval with a given coverage probability (Box and Tiao, 1973), defined as

\[
\text{hdPI}_{1-\alpha}(x) = \{ y : f_{Y|X=x}(y) > \tau_{1-\alpha} \}
\]

where \( \tau_{1-\alpha} \) is the largest number satisfying \( \int_{y \in \text{hdPI}_{1-\alpha}(x)} f_{Y|X=x}(y) \, dy = 1 - \alpha \). Another nice property of hdPIs is that any point within the interval has a higher density than any other point outside it, which justifies their name.

**Example 11** Butterfly data, continued. All three prediction intervals are shown in Fig. 9(a). Three main conclusions: (i) The shortest among all is the highest-density PI, which take the form of disjoint subintervals—one for each local mode. (ii) The Gaussian assumption of gPI seems too restrictive for real-world data analysis. (iii) Both gPI and qPI include a low-density valley area around \( x = 0 \) at the cost of other, more-likely values around \( x = \pm 2 \).

**Example 12** The Auto-MPG data. The task is to predict the miles-per-gallon (MPG) gas consumption of an automobile based on \( p = 7 \) features including horsepower, weight, and acceleration. The dataset was used in the 1983 American Statistical Association Exposition and is available in the UCI Machine Learning Repository. Here, we are mainly interested in predicting the MPG of a specific instance-row number 390, which corresponds to the 1982 Dodge Rampage car. The ISL-predicted uncertainty distribution is displayed in Fig. 9(b). The bimodality of the estimated conditional density reflects the fact that 1982 Dodge Rampage is a hybrid two-door car—a mix between a passenger car and a truck. The next notable thing is the length of the PIs: The standard error-based qPI is almost three times wider than the hdPI, and thus provides an overly pessimistic assessment of an accurate prediction.

#### 5 70 Years of Statistical Machine Learning

Breiman’s 2001 paper was highly influential and hailed as a heroic effort to challenge the status quo in a “David and Goliath” style, where two diametrically opposite modeling cultures—machine learning and traditional parametric statistics—battled for the throne. However, the origin of machine learning has strong historical roots in (nonparametric) Statistics. In an unpublished US Air Force School of Aviation Medicine report in 1951, two UC Berkeley Statisticians (students of Jerzy Neyman) Evelyn Fix and J.L. Hodges, Jr., introduced a non-parametric method for pattern classification, which is now called the k-nearest neighbor (knn) method. This was a groundbreaking paper that marked the beginning of statistical machine learning. Fix and Hodges’ work was a precursor to the celebrated kernel-based methods and decision tree-based algorithms, which revolutionized the world of machine learning. In this paper, we have initiated the “re-unification” process with the hope to bring machine learning back to its statistical roots. Two broad messages of this research:

1. **United statistical learning culture.** The algorithmic models should not be feared because of their complexity, parametric models should not be shunned because of their simplicity, we can interlink them to create a more powerful and coherent learning technology. This research shows how one can build such an integrated learning system where different cultures of data modeling (embracing: algorithmic, parametric, nonparametric, information-theoretic, robust, and exploratory) can be “glued” together by some underlying deeper principles.

2. **There’s plenty of room in the middle.** Tremendous opportunities lie at the boundaries between the two cultures. This paper describes a new theory for constructing a universal interface between statistics and machine learning, as a means to close the gap. What we have proposed is by no means the only possible or the “best” theory. As we go forward, it will be very important to come up with new innovative ways to develop “integrated statistical learning” theory that can reconcile and build new links between various cultures of data analysis in order to maintain a
diverse, yet united front of statistical science. All of this is still in its nascent stages, but definitely poised for rapid progress in the coming decade.

**ACKNOWLEDGMENTS**

This paper celebrates the 200th anniversary of Gauss-Markov least-square regression and 70th anniversary of statistical machine learning, by putting forward a modern unified perspective.

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11. These two historical moments symbolize the birth of ‘two cultures’ of data modeling—linear parametric and flexible algorithmic modeling cultures.

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Supplementary Material

A.1. SOFTWARE

We have developed an R Software LPMachineLearning\textsuperscript{12} to perform all the tasks outlined in the paper. We hope this software will encourage applied data scientists to apply our method for their real prediction problems.

A.2. LP-ORTHONORMAL SYSTEM: CONSTRUCTION AND PROPERTIES

LP-basis functions are specially-designed orthonormal system of rank-polynomials. They are inherently ‘nonparametric,’ constructed in a fully data-driven way, not pre-defined like classical polynomials. For a mixed (either discrete or continuous) random variable \( Z \) with distribution function \( F_Z \), we describe the construction of LP-basis functions.

**Construction Algorithm.** A fully-automated construction of LP-orthonormal rank-polynomials of \( Z \) consists of the following steps:

1. **Mid-Distribution Transform:** The mid-distribution function of \( Z \) is defined as

\[
F_Z^{\text{mid}}(z) = F_Z(z) - \frac{1}{2} p_Z(z), \tag{E5.1}
\]

where \( p_Z(z) \) is the probability mass function.

2. **Standardizing \( F_Z^{\text{mid}}(Z) \):** The random variable \( F_Z^{\text{mid}}(Z) \) has mean \( \mathbb{E}[F_Z^{\text{mid}}(Z)] = 0.5 \) and variance \( \text{Var}[F_Z^{\text{mid}}(Z)] = \frac{1}{12} (1 - \sum_z p_Z^2(z)) \). Define the 1st-order LP-basis function as

\[
T_1(z; F_Z) = \frac{\sqrt{12} \{ F_Z^{\text{mid}}(z) - 1/2 \}}{\sqrt{1 - \sum_z p_Z^2(z)}}, \tag{E5.2}
\]

3. **Gram-Schmidt Orthonormalization:** Construct high-order polynomials by applying Gram-Schmidt orthonormalization (see Appendix A of Mukhopadhyay and Parzen (2020) for a quick refresher on Gram-Schmidt process) on the set of functions of the power of \( T_1(Z; F_Z) \).

**Some properties.** Here we list some important properties of the They satisfy the following properties of the LP-orthonormal systems:

1) LP-bases are orthonormal with respect to the measure (weighting function) \( F_Z \):

\[
\int T_j(z; F_Z) \, dF_Z = 0,
\]

and

\[
\int T_j(z; F_Z) T_k(z; F_Z) \, dF_Z = \delta_{jk}.
\]

For related discussions, see Remark 4 of Mukhopadhyay and Parzen (2020, p. 82).

2) For empirical data analysis, construct empirical LP basis (in short eLP basis) \( \{ T_j(z; \hat{F}_Z) \}_{j=1,2,...,m} \), where \( m \) is strictly less than the number of unique values in the sample \( \{ z_1, \ldots, z_m \} \). eLP bases are orthonormal system of functions with respect to the discrete empirical measure \( \hat{F}_Z \).

\textsuperscript{12} A preliminary version is available from the authors. A more upgraded one is currently under development.

Fig. 10: First four LP-basis functions for the covariates (a) weight and (b) acceleration of Auto-MPG data. They are plotted with respect to \( \hat{F}_X(x) \) over unit interval \([0, 1]\), following the equation (E5.11).

3) For \( Z \) continuous, it is not difficult to show that the LP-bases reduces to the following universal shape:

\[
T_j(z; F_Z) := \text{Leg}_j \circ F_Z(z), \tag{E5.3}
\]

where \( \text{Leg}_j(u) \) denotes \( j \)-th shifted orthonormal Legendre polynomials on \([0, 1]\) and ‘\( \circ \)’ denotes composition of functions.\textsuperscript{13} Few top LP-polynomials for \( Z \) continuous are given below:

\[
T_1(z; F_Z) = \text{Leg}_1 \left( F_Z(Z) \right) = \sqrt{12} (F_Z(z) - 1/2) \tag{E5.4}
\]

\[
T_2(z; F_Z) = \text{Leg}_2 \left( F_Z(Z) \right) = \sqrt{5} (6 F_Z^2(z) - 6 F_Z(Z) + 1) \tag{E5.5}
\]

\[
T_3(z; F_Z) = \text{Leg}_3 \left( F_Z(Z) \right) = \sqrt{7} (20 F_Z^3(z) - 30 F_Z^2(z) + 12 F_Z(Z) - 1). \tag{E5.6}
\]

and so on. For more information, see Mukhopadhyay and Fletcher (2018), Mukhopadhyay (2017). This will be useful for constructing LP-polynomials for any general pivot density \( f_0 \) of the response variable.

4) Derivation of empirical \( T_1(z; \hat{F}_Z) \) from a sample of all unique \( z_1, \ldots, z_N \) (i.e., realization from a continuous \( Z \)). In this case, the empirical probability mass function (pmf) is

\[
\hat{p}_Z(z_i) = 1/N, \quad \text{and the empirical cdf is} \quad \hat{F}_Z(z_i) = R_i/N,\]

where \( R_i = \text{rank}(z_i) \). This immediately implies (following eq. E5.2)

\[
T_1(z; \hat{F}_Z) = \sqrt{\frac{12}{N^2 - 1}} \left( R_i - \frac{N + 1}{2} \right). \tag{E5.7}
\]

Since,

\[
\hat{F}_Z^{\text{mid}}(z_i) = \hat{F}_Z(z_i) - \frac{1}{2} \hat{p}_Z(z_i) = \frac{1}{N} (R_i - 1/2).
\]

and the correction factor

\[
1 - \sum_i \hat{p}_Z^2(z_i) = 1 - \frac{1}{N^2}.
\]

Eq. (E5.7) makes it clear why we call LP-bases are polynomials of ranks.

\textsuperscript{13} In our regression context the response variable \( Y \) is continuous, thus admits this analytic LP-basis form. However, this one-to-one correspondence between LP-polynomials and Legendre polynomials of rank-transform (also know as probability integral transform) \( F_Z(Z) \) is not true for discrete \( Z \), i.e., for the covariates \( X \).
5) The shapes of LP-polynomials are data-adaptive. The left panel of Fig. 10 displays the top four eLP-basis functions for the covariate weight, taken from the Auto-MPG data (Example 12 in the main paper). The weight variable has very little ties, which is the reason why the shapes of LP-bases completely match with eq. (E5.4-E5.6). Contrast this with the right panel of Fig. 10, which shows top four eLP bases for the variable acceleration. Due to the presence of a large number of ties, it takes a unique piecewise-constant shape.

A.3. REMAINING PROOFS

Theorem 2.2: Since the conditional entropy \( H(Y|X) = \int H(Y|X = x) \ dF_X(x) \), we have

\[
H(Y) - H(Y|X) = - \int f_Y(y) \log f_Y(y) \ dy + \int f_{Y|X}(y) \log f_{Y|X}(y) \ dy \ dF_X(x) \\
= \int \{ - f_Y(y) \log f_Y(y) + f_{Y|X}(y) \log f_{Y|X}(y) \} \ dF_X(x) \\
= \frac{\text{KL}(d_Y; \mathbb{1}_{[0,1]})}{E_X}.
\]

The last equality follows from Theorem 2.1.

Theorem 4.3: The Kruskal–Wallis statistic (21) is given by (up to a negligible factor \( (n - 1)/n \))

\[
\text{KW} = \sum_{\ell = 1}^{k} n_{\ell} \left\{ \frac{12}{N^2 - 1} \left( n_{\ell} - 1 \right) R_1 - \frac{N + 1}{2} \right\}^2.
\]

Now note that

\[
\hat{\text{LP}}_{1|\ell} = E[T_1(Y; F_Y)|X = \ell] \\
= \frac{1}{n_{\ell}} \sum_{i \in G_\ell} \left( R_i - \frac{N + 1}{2} \right).
\]

The first equality follows from (11) and the second one from (E5.7). Combining (E5.8) and (E5.9) we get the first desired result

\[
\text{KW} = \sum_{\ell = 1}^{k} n_{\ell} |\hat{\text{LP}}_{1|\ell}|^2.
\]

Next, to show that (E5.10) is actually equal to \( N R_1^2 \), note that

\[
R_1^2 = \frac{\text{Var}(E[T_1(Y; F_Y)|X])}{\text{Var}(T_1(Y; F_Y))},
\]

where the denominator is 1 by the construction of LP-basis functions, and the numerator is \( \sum_{\ell = 1}^{k} \pi_{\ell} |\hat{\text{LP}}_{1|\ell}|^2 \), \( \pi_{\ell} \) is \( P_r(X = \ell) = n_{\ell}/N \). This is because

\[
E[\text{LP}_{1|X}] = E[E[T_1(Y; F_Y)|X]] = E[T_1(Y; F_Y)] = 0.
\]

**Theorem 4.3**: Define the LP unit-basis functions as

\[
S_j(u; F_Y) = T_j(Q_Y(u); F_Y) \quad 0 < u < 1,
\]

which satisfy the following orthonormality relations (immediate from eq. 10)

\[
\int_0^1 S_j(u; F_Y) S_k(u; F_Y) \ du = \delta_{jk}.
\]
outward—correctly reflecting the uncertainty of the mean function, which indicates that the estimated regression line is impressively accurate!

Even though we have an “accurate” estimate of the underlying regression prediction, it does not tell us anything useful. In fact, it paints a grossly misleading picture of the reality (in the sense that x is not useful for predicting y, which is clearly untrue). To a fuller picture, it necessary to go beyond conventional practice of curve fitting.

**Level 1. Heterogeneity diagnostic.** A natural question is whether the initial ML₀ captured the essential relationship between y and the predictor variables. This can be checked by searching for (unmodeled) patterns in the residuals. Recall that for the butterfly data \( \hat{\mu}(x) \approx 0 \) for all \( x \); thus residuals are actually the original response values \( y_1, \ldots, y_n \).

The critical task boils down to checking whether the residuals are homogeneous, i.e., \( f_{Y|X=x}(y) = f_Y(y) \) for all \( x \). This would ensure that ML₀ was able to capture all the important predictive information. However, in most practical real-life problems (including butterfly data) the residuals will be heterogeneous. And the main challenge is to identify the components of \( f_{Y|X=x}(y) \) that are significantly changing with \( x \). We like to address this by constructing a simple diagnostic plot that can help users to visually examine and detect the prominent sources of heterogeneity in the residuals. Fig. 12(a) is one such graphical exploratory plot drawn for the butterfly data, which reveals scale (second-order effect) and tail (fourth-order kurtosis effect) of \( f_{Y|X=x}(y) \) are changing significantly with \( x \). Accordingly, this tool helps us to *discover* the hidden patterns that, while invisible to ML₀, still persist and should be modeled for sharpening the prediction quality; also see Appendix A.8.

**Level 2. Pivot density.** We now turn our attention to the problem of predicting conditional distribution at a particular \( x \), say at \( x_0 = 2 \) for the butterfly example. There are some obvious candidates: \( \hat{f}_Y(y) \) the empirical (marginal) distribution of \( y \); or its nonparametric smooth (say, kernel-smoothed) version \( \hat{f}_Y(y) \); or some kind of parametric model motivated from domain-knowledge or statistical convenience, e.g., \( \mathcal{N}(\hat{\mu}(x_0), \hat{\sigma}_y^2) \), where \( \hat{\sigma}_y \) denotes the standard error of \( y \). We call these initial starting "guesses" the pivot density; they are nothing but a crude first-approximation of the true unknown conditional density estimate. For the butterfly data, we choose \( \mathcal{N}(0, \hat{\sigma}_y^2 = 2.55^2) \) to be the pivot, as shown in Fig. 12(b).

A natural question to ask is whether the presumed pivot model is approximately correct. In most practical cases, the answer will be no, they are not adequate. Thus we have to take some corrective measures to rectify the deficiency of the starting pivot. But before doing so, it is crucial to devise some technique that can reveal the deficiency of the current model. The only problem is that we don’t have enough \( y \)-samples at \( X = x_0 \) to perform traditional goodness-of-fit tests. We have to come up with something new that is computable and interpretable.

**Level 3. Uncertainty quantification for the pivot.** The basic question is whether the starting pivot density \( f_0(y) \) is a good model for \( Y_{|X=x} \). We answer this question by estimating the following contrast density function for \( u = F_0(y) \)

\[
d(u; F_0, F_{Y|X=x}) := \frac{f_{Y|X=x}(F_0^{-1}(u))}{f_0(F_0^{-1}(u))}, \quad 0 < u < 1
\]

where \( F_0^{-1}(u) \) is the quantile function for the pivot \( f_0(y) \). Note that \( d_0^0(u) \) captures the deviation of \( f_0(y) \) from the true unknown conditional density \( f_{Y|X=x_0}(y) \), and by virtue of doing so it helps us to quantify and characterize the uncertainty of the pivot density. By inspecting the shape of \( d_0^0(u) \), \( 0 < u < 1 \) (how it deviates from uniformity), practitioners can quickly infer whether \( f_0(y) \) needs any repair or not.

Fig. 12(c) displays the estimated contrast function for the butterfly data. Looking at the shape, it becomes clear that the initial Gaussian pivot completely missed the bimodality of \( f_{Y|X=x}(y) \). The contrast function \( d_0^0(u) \) plays a central role in our theory. The fascinating aspect of the algorithm is that it utilizes the power of the initial machine learning
algorithm ML₀ to get the covariate-adaptive ᵃ₀.

**Level 4. Density modulation.** When the contrast function indicates the inadequacy of the starting pivot ᵇ₀(y) as a model for the conditional distribution Ƒ_Y|X=x(y), what to do next? How to modify (or ‘boost’) ᵇ₀(y) to make it data-consistent? This can be solved in a remarkably simple way via density modulation: Ƒ_Y|X=x = ᵇ₀ × ᵃ₀, where ᵃ₀ modulates (modifies) the shape of starting ᵇ₀(y) to produce the conditional density. The detailed theory of why and how this works are discussed in Section 2 of the main article. Fig. 12(d) displays the estimated predictive density at X = 2, which we can write down analytically as

$$\hat{F}_{Y|X=x}(y) = \frac{1}{\hat{\sigma}} \phi\left(\frac{y}{\hat{\sigma}}\right) \times \left\{1 - 0.20 T_2(y; F_0) - 0.47 T_4(y; F_0)\right\},$$

where \(\hat{\sigma} = 2.55\) (E5.14)

which is product of two components: the Gaussian pivot and the contrast density ᵃ₀(0)(F₀(y)). The basis functions \(\{T_j(y; F_0)\}\) in equation (5) are specially-designed rank-based orthonormal polynomials of the pivot. They inject robustness and non-linearity into the data modeling process—more on this later. We will also show that this new class of d-modulated conditional density models can generate an extraordinarily rich spectrum of shapes.

One notable aspect of (5) is that we have ‘identified’ a parametric model (instead of blindly assuming, which Breiman sharply criticized by algorithmic (fully nonparametric) means. In Section 2.3, we elucidate the procedure by which we boost and convert the “0th order” machine learning method (ML₀) into a finitely parameterizable uncertainty prediction machine. This allows our learning philosophy to integrate the fundamental character of both algorithmic data-driven modeling and parametric statistical modeling. Nevertheless, the predicted distribution (5) compactly represents all possible outcome values along with their respective probabilities, which is essential for decision-making under uncertainty.

**Level 5. Predictive inference:** What are some of the most likely values of the response variable y, given that we have observed X = x₀? For a fixed α, ideally, we would like to find the smallest volume region that covers 100(1 − α)% area of the conditional distribution, aka the highest density prediction region. The orange highlighted part in Fig. 12(d) displays 68% highest-density prediction region for the butterfly data, which consists of two disjoint intervals [-3.33, -1.40] ∪ [1.20, 3.28]. The volume of this set (the sum of the lengths of the intervals) can be used to quantify the uncertainty of the prediction. Additionally, we can extract all the conditional quantiles from the estimated \(\hat{F}_{Y|X=x}(y)\); see panel (e). Our quantile regression curves are, by design, non-crossing and nonlinear. Curious readers may contrast this with the traditional quantile regression estimator, shown in Fig. 13 of Appendix A.5.1.

**Level 6. Verification:** Data modeling typically consists of four stages: estimation (of the initial pivot), exploration (discovering the deficiency of the pivot), rectification (repairing the pivot), and verification (checking whether the corrected pivot emulates the observed reality). The last step remains to be done, which requires evidence that the estimated uncertainty prediction model provides a satisfactory description of the data. Scientific data analysis, by definition, needs to be testable (falsifiable), and there’s no two ways about it.

For our butterfly example, we compute \(U_i = \hat{F}_{Y|X=x,i}(y_i)\) for \(i = 1, \ldots, N\) using our trained model on a 15% hold-out validation dataset. It is not difficult to see that the distribution of \(U_i\)’s will be ‘close’ to Uniform[0,1], if the estimated conditional density can explain the essential patterns/variations in the data. The quantiles of the empirical distribution of \(U_i\)’s are plotted against the respective quantiles of the uniform distribution at the bottom right of Fig. 12. The tight cluster of points around the 45-degree reference line strongly indicates that the model is successful in capturing the underlying data generating process, and can be trusted to make predictions. Additionally, the formal goodness-of-fit test (proposed in Section 3.3), when applied for this example, yields a p-value of 0.82.

### A.5. Additional Remarks

**Remark A.5.1** Koenker’s Quantile regression for Butterfly data: Fig. 13 shows the quantile regression curves for the butterfly data, which implements the algorithm proposed in Koenker and d’Orey (1987).

**Remark A.5.2** Smoothness of Regression via Rank-transform. Fig. 14 shows house prices (median value of in 1000s) in Boston metropolitan area as function of crime (per capita crime rate). The data were collected in 1970 and available in the UCI Machine Learning Repository. The left plot shows the data in the xy-domain, and the right plot shows the same data in the quantile domain where we replaced x by its rank-transform \(\hat{F}_X(x)\). Red curves are the spline smoother (estimated using smooth.spline() R-function). Few interesting points to note:

- **Smoothness:** Regression via rank-transform (see Theorem 2.5) seems to produce much smoother (parsimonious) regression curve compared to the rough zigzag xy-domain estimate. Is it universal that nature reveals herself in a more parsimonious way in the quantile domain? Apart from intuitive understanding, we still don’t know any concrete mathematical explanation for this surprising phenomenon.

- **Data-sparsity:** The quantile-domain treatment seems to address the ‘data sparsity’ (that exists for \(x > 30\) prob-

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**Fig. 13:** Butterfly data: Performs Roger Koenker’s (1978) quantile regression routine as implemented in rq(v) function of the R-package quantreg. Note the difference with Fig. 16.
the so-called ‘unconfoundedness’ assumption to attach a causal interpretation.

Remark A.5.6 Personalized reference interval prediction: The topic of individualized referencing carries special significance in the era of precision medicine. To accurately infer each individual’s reference curves (describing the normal range of the outcome $y$ given $x$) medical researchers are now incorporating more and more variables in the form of genetic and environmental factors. Thus it will be of great practical value to have a method that can tackle a large number of covariates, and do so in a completely nonparametric and nonlinear fashion.

A.6. MODERN APPLIED STATISTICS: THEORY, PRACTICE, AND PEDAGOGY

A theoretical research program that has no relevance to practice and teaching, is not persuasive enough to be taken as fundamental. To us, it is very important to know whether our research was able to link (at least partially) the triad of theory-practice-teaching. In this article, through several examples, we made an attempt to describe ‘the joy of systematic data analysis’ derived from a general theory, which has the following implications for the practice and pedagogy of Statistics:

- Our theory makes it easy to apply (arise from simplification) and see the connections (arise from unification) between different statistical methods.
- Our theory provides a holistic training\textsuperscript{15} by introducing a large variety of statistical topics (embracing different cultures: algorithmic, parametric, nonparametric, information-theoretic, robust, exploratory) in an ‘organized’ manner through a common semantics. This is extremely important to produce broad integrator foxes rather than highly specialized hedgehogs\textsuperscript{16}. Ultimately, what matters is whether our theory provides a faster and easier (lazier?) way to learn the fundamentals of statistics\textsuperscript{17}. We believe–yes, it does.

A.7. TWO SURPRISING CONNECTIONS: FROM AI AND ECONOMICS

- Herbert Simon’s theory to simulate human thinking. In the year of 1956, at the historic conference “Dartmouth Summer Research Project on Artificial Intelligence,” Nobel laureate Herbert Simon, along with Cliff Shaw and Allen Newell, presented a universal problem solver machine called ‘General Problem Solver’ [GPS], which is considered by many to be the first AI program. Their theory is based on the idea of “Difference Machine,” which proceeds as follows:

\begin{itemize}
  \item \textbf{Remark A.5.5} Connection of XYZ Problem with causal inference: For randomized experiments and clinical trials our DIF statistic can be interpreted as a ‘distributional’ treatment effect\textsuperscript{14}, one whose impact vary across subpopulations. For observational studies, one needs to assume

\textbf{Remark A.5.4} Robust Lasso: As already noted, the online news data contain an outlier–the news article in row #31,038. How does it impact traditional estimation and feature selection methods? As we will see, a single data point can have a devastating impact on the overall modeling. Fig. 17 (left panel) shows the lasso estimated $\hat{\beta}$ under two scenarios: first, on the outlier-removed data, and second, on the full data (with outlier). Notice the estimated coefficient values (as well as their sparsity-levels) are dramatically affected because of just one data point. One way to address this fragility is to apply lasso on the first-order LP-transformed $T_X = [T_1(x; F_{X_1}), \ldots, T_1(x; F_{X_{p_i}})]$, instead of raw feature matrix $X$. The result is displayed on the right panel, which shows extraordinary consistency with (almost) no impact on the estimated coefficients. This mid-rank transform based \textit{robust lasso} method could be a reliable alternative to deal with noisy messy data.

\textbf{Remark A.5.3} ML-assisted Quantile regression: comparison. The first row of Fig. 16 shows the generalized random forests-based (Athey et al., 2019) and gradient boosted quantile regression estimates. They are specially-designed ML-algorithms to produce the desired conditional quantile curves. The second row shows our ISL versions, which are derived from a general scheme described in Section 2.3 of the main paper.

\textbf{Remark A.5.2} A theoretical research program that has no relevance to practice and teaching, is not persuasive enough to be taken as fundamental. To us, it is very important to know whether our research was able to link (at least partially) the triad of theory-practice-teaching. In this article, through several examples, we made an attempt to describe ‘the joy of systematic data analysis’ derived from a general theory, which has the following implications for the practice and pedagogy of Statistics:

\textbf{Remark A.5.1} Personalized reference interval prediction: The topic of individualized referencing carries special significance in the era of precision medicine. To accurately infer each individual’s reference curves (describing the normal range of the outcome $y$ given $x$) medical researchers are now incorporating more and more variables in the form of genetic and environmental factors. Thus it will be of great practical value to have a method that can tackle a large number of covariates, and do so in a completely nonparametric and nonlinear fashion.

\textbf{A.6. MODERN APPLIED STATISTICS: THEORY, PRACTICE, AND PEDAGOGY}

A theoretical research program that has no relevance to practice and teaching, is not persuasive enough to be taken as fundamental. To us, it is very important to know whether our research was able to link (at least partially) the triad of theory-practice-teaching. In this article, through several examples, we made an attempt to describe ‘the joy of systematic data analysis’ derived from a general theory, which has the following implications for the practice and pedagogy of Statistics:

\textbf{Remark A.5.6} Personalized reference interval prediction: The topic of individualized referencing carries special significance in the era of precision medicine. To accurately infer each individual’s reference curves (describing the normal range of the outcome $y$ given $x$) medical researchers are now incorporating more and more variables in the form of genetic and environmental factors. Thus it will be of great practical value to have a method that can tackle a large number of covariates, and do so in a completely nonparametric and nonlinear fashion.

\textbf{Remark A.5.5} Connection of XYZ Problem with causal inference: For randomized experiments and clinical trials our DIF statistic can be interpreted as a ‘distributional’ treatment effect\textsuperscript{14}, one whose impact vary across subpopulations. For observational studies, one needs to assume

\textbf{Remark A.5.4} Robust Lasso: As already noted, the online news data contain an outlier–the news article in row #31,038. How does it impact traditional estimation and feature selection methods? As we will see, a single data point can have a devastating impact on the overall modeling. Fig. 17 (left panel) shows the lasso estimated $\hat{\beta}$ under two scenarios: first, on the outlier-removed data, and second, on the full data (with outlier). Notice the estimated coefficient values (as well as their sparsity-levels) are dramatically affected because of just one data point. One way to address this fragility is to apply lasso on the first-order LP-transformed $T_X = [T_1(x; F_{X_1}), \ldots, T_1(x; F_{X_{p_i}})]$, instead of raw feature matrix $X$. The result is displayed on the right panel, which shows extraordinary consistency with (almost) no impact on the estimated coefficients. This mid-rank transform based \textit{robust lasso} method could be a reliable alternative to deal with noisy messy data.

\textbf{Remark A.5.3} ML-assisted Quantile regression: comparison. The first row of Fig. 16 shows the generalized random forests-based (Athey et al., 2019) and gradient boosted quantile regression estimates. They are specially-designed ML-algorithms to produce the desired conditional quantile curves. The second row shows our ISL versions, which are derived from a general scheme described in Section 2.3 of the main paper.

\textbf{Remark A.5.2} A theoretical research program that has no relevance to practice and teaching, is not persuasive enough to be taken as fundamental. To us, it is very important to know whether our research was able to link (at least partially) the triad of theory-practice-teaching. In this article, through several examples, we made an attempt to describe ‘the joy of systematic data analysis’ derived from a general theory, which has the following implications for the practice and pedagogy of Statistics:

\textbf{Remark A.5.1} Personalized reference interval prediction: The topic of individualized referencing carries special significance in the era of precision medicine. To accurately infer each individual’s reference curves (describing the normal range of the outcome $y$ given $x$) medical researchers are now incorporating more and more variables in the form of genetic and environmental factors. Thus it will be of great practical value to have a method that can tackle a large number of covariates, and do so in a completely nonparametric and nonlinear fashion.

\textbf{A.6. MODERN APPLIED STATISTICS: THEORY, PRACTICE, AND PEDAGOGY}

A theoretical research program that has no relevance to practice and teaching, is not persuasive enough to be taken as fundamental. To us, it is very important to know whether our research was able to link (at least partially) the triad of theory-practice-teaching. In this article, through several examples, we made an attempt to describe ‘the joy of systematic data analysis’ derived from a general theory, which has the following implications for the practice and pedagogy of Statistics:

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\textbf{Remark A.5.5} Connection of XYZ Problem with causal inference: For randomized experiments and clinical trials our DIF statistic can be interpreted as a ‘distributional’ treatment effect\textsuperscript{14}, one whose impact vary across subpopulations. For observational studies, one needs to assume
(i) compare \( C \) [initial steady-state] with \( F \) [future desired-state] to check whether there exists any difference; (ii) devise some technique (“intelligence”) to rectify the present state to reduce the important differences between \( C \) and \( F \) — call it the difference-operator; (iii) if this succeeds, then we have created a new object \( x' \), which (one hopes) no longer has the difference when compared with the desired state \( F \). The question remains, how can we transform this hierarchical logic into a computable theory? It’s quite fascinating that the GPS logic is already encoded into our modeling scheme (6), where the contrast function \( d_x \) acts according to Herbert Simon’s difference-operator. In our setup, \( d_x \) actively works to remove the differences between the present (steady/null) state \( f_Y(y) \) and the future (desired) state \( f_{Y|X=x}(y) \). The main reference here is Newell et al. (1959). Similar ideas also appeared in the Ph.D. thesis of Patrick Winston (1970), and more recently Marvin Minsky’s (2007).

- Hansen-Sargent theory of model misspecification. Lars Hansen and Thomas Sargent, two Nobel laureates, in their book “Robustness,” raised a profound modeling question: Is there any robust way to construct a class of plausible models starting from a misspecified one?

It is an understatement to say that model misspecification is at least as much of a problem in economics as it is in physics and engineering. (Hansen and Sargent, 2008, p. 3)

Our \( d \)-modulation based density representation theory (8) creates a class of reasonable models around a possibly misspecified pivot. Here the contrast density \( d_x \) acts as a data-driven perturbation function in order to ‘repair’ the starting approximate model \( f_0 \).

### A.8. Sharpening Black-box Generators

Imagine we are given some black-box generative model that can simulate \( y \)-samples for a given \( x = (x_1, \ldots, x_p) \):

\[
\text{MODEL}(x) \rightarrow \{\tilde{y}^1_x, \ldots, \tilde{y}^s_x\}.
\]

Without having any information on the particular \text{MODEL}\(^{18}\), our goal is to: (i) check whether the samples are truly generated from the underlying conditional distribution \( f_{Y|X=x}(y) \); (ii) if not, explain ‘why’ by displaying the associated \( d_x \); and finally, (iii) sharpen the given imperfect samples to make them compatible with the underlying stochastic data generating mechanism.

**Butterfly data cont.** To illustrate the method, let us consider that we are given all the \( y_i \)'s for which the \( x_i \) value is between 0 and 4. The histogram of these samples is shown in the left panel of Fig. 15. The blue curve shows the true \( f_{Y|X=x}(y) \). Starting from these weak conditional samples, our goal is to refine it so that they comply with the underlying law at \( x = 2 \).

Step 1. Choose the empirical distribution \( \tilde{f}_{y|x} \) of \( \{\tilde{y}^1_x, \ldots, \tilde{y}^s_x\} \) be the pivot; see Fig. 15(a).

Step 2. Randomly generate one sample \( \tilde{y}_s \) from \( \tilde{f}_{y|x} \) and \( U \) from Uniform[0, 1].

\(^{18}\) This could be any arbitrarily complex computational program, e.g., deep learning method with millions of parameters, an ensemble of 100 machine learning models, or the method proposed in Friedman, J. H. (2020).
Fig. 17: Online news popularity data. The goal is to assess the impact of a single outlier observation (the news article in row #31, 038 on the lasso-estimated coefficients. The left plot shows the result for lasso(y ~ X) case, and the right one for robustified lasso(y ~ TX). For more details see remark A.5.4.

| X (Age) | Y (FEV) | Z (Smoking) |
|---------|---------|-------------|
| 9       | 1.70    | 0           |
| 8       | 1.72    | 0           |
| ...     | ...     | ...         |
| 15      | 3.73    | 1           |
| 18      | 2.85    | 0           |

TABLE 2: We are given N = 654 observations on youths aged 3 to 19 from East Boston recorded during 1970's. The outcome variable Y is forced expiratory volume in 1 second—a measure of lung capacity. Z is 1/0 binary treatment (smoker: yes or no) indicator variable.

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Fig. 18: (color) 1990’s film data: first row shows a part of the full film90 data and the starting normal pivot. Notice the “two-branches” of the data cloud. Second row: The estimated contrast densities \( \hat{d}_{Y|X=x}(u) \), which describe the conversion factor: how the pivot \( f_0(y) \) has to be transformed (perturb) to get to \( f_{Y|X=x}(y) \). Third row: The refurbished conditional densities \( \hat{f}_{Y|X=x}(y) \) obeying the \( d \)-modulation law (8); the red triangles denote the mode of the distributions. Notice the difference in shape between the starting Gaussian pivot (panel b) and the final estimated conditional densities (of the bottom row).

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Fig. 19: Exploratory goodness-of-fit assessment for the butterfly data. The histograms and QQ plots of the generalized quantile-residuals (18) are shown for three ML-models. They are computed based on a 15% hold-out data. Knn (with $k = 15$) seems fits the data well.

Fig. 20: Dutch boys data: The ISL-estimated conditional quantile curves. Our method yields non-crossing quantile curves, as they are extracted from the conditional distributions.