

¹ This is a draft version of work in progress, content will be revisited in subsequent versions.

² Robust Local Polynomial Regression with Similarity Kernels

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⁵ Abstract

Local Polynomial Regression (LPR) is a widely used nonparametric method for modeling complex relationships due to its flexibility and simplicity. It estimates a regression function by fitting low-degree polynomials to localized subsets of the data, weighted by proximity. However, traditional LPR is sensitive to outliers and high-leverage points, which can significantly affect estimation accuracy. This paper revisits the kernel function used to compute regression weights and proposes a novel framework that incorporates both predictor and response variables in the weighting mechanism. By introducing two positive definite kernels, the proposed method robustly estimates weights, mitigating the influence of outliers through localized density estimation. The method is implemented in Python and is publicly available at <https://github.com/yaniv-shulman/rsklpr>, demonstrating competitive performance in synthetic benchmark experiments. Compared to standard LPR, the proposed approach consistently improves robustness and accuracy, especially in heteroscedastic and noisy environments, without requiring multiple iterations. This advancement provides a promising extension to traditional LPR, opening new possibilities for robust regression applications.

6 **1. Introduction**

7 Local polynomial regression (LPR) is a powerful and flexible statistical technique that has
8 gained increasing popularity in recent years due to its ability to model complex relationships be-
9 tween variables. Local polynomial regression generalizes the polynomial regression and moving
10 average methods by fitting a low-degree polynomial to a nearest neighbors subset of the data at
11 the location. The polynomial is fitted using weighted ordinary least squares, giving more weight
12 to nearby points and less weight to points further away. The value of the regression function for
13 the point is then obtained by evaluating the fitted local polynomial using the predictor variable
14 value for that data point. LPR has good accuracy near the boundary and performs better than all
15 other linear smoothers in a minimax sense [2]. The biggest advantage of this class of methods
16 is not requiring a prior specification of a function i.e. a parametrized model. Instead only a
17 small number of hyperparameters need to be specified such as the type of kernel, a smoothing
18 parameter and the degree of the local polynomial. The method is therefore suitable for modeling
19 complex processes such as non-linear relationships, or complex dependencies for which no the-
20 oretical models exist. These two advantages, combined with the simplicity of the method, makes
21 it one of the most attractive of the modern regression methods for applications that fit the general
22 framework of least squares regression but have a complex deterministic structure.

23 Local polynomial regression incorporates the notion of proximity in two ways. The first is
24 that a smooth function can be reasonably approximated in a local neighborhood by a simple
25 function such as a linear or low order polynomial. The second is the assumption that nearby
26 points carry more importance in the calculation of a simple local approximation or alternatively
27 that closer points are more likely to interact in simpler ways than far away points. This is achieved
28 by a kernel which produces values that diminish as the distance between the explanatory variables
29 increase to model stronger relationship between closer points.

30 Methods in the LPR family include the Nadaraya-Watson estimator [10, 18] and the estimator
31 proposed by Gasser and Müller [7] which both perform kernel based local constant fit. These
32 were improved on in terms of asymptotic bias by the proposal of the local linear and more general
33 local polynomial estimators [16, 3, 9, 4, 5]. For a review of LPR methods the interested reader
34 is referred to [2].

35 LPR is however susceptible to outliers, high leverage points and functions with discontinu-
36 ities in their derivative which often cause an adverse impact on the regression due to its use
37 of least squares based optimization [17]. The use of unbounded loss functions may result in
38 anomalous observations severely affecting the local estimate. Substantial work has been done to
39 develop algorithms to apply LPR to difficult data. To alleviate the issue [15] employs variable
40 bandwidth to exclude observations for which residuals from the resulting estimator are large. In
41 [3] an iterated weighted fitting procedure is proposed that assigns in each consecutive iteration
42 smaller weights to points that are farther then the fitted values at the previous iteration. The pro-
43 cess repeats for a number of iterations and the final values are considered the robust parameters
44 and fitted values. An alternative common approach is to replace the squared prediction loss by
45 one that is more robust to the presence of large residuals by increasing more slowly or a loss that
46 has an upper bound such as the Tukey or Huber loss. These methods however require specifying
47 a threshold parameter for the loss to indicate atypical observations or standardizing the errors
48 using robust estimators of scale [8]. For a recent review of robust LPR and other nonparametric
49 methods see [17, 11]

50 The main contribution of this paper is to revisit the kernel used to produce regression weights.
51 The simple yet effective idea is to generalize the kernel such that both the predictor and the re-

52 response are used to calculate weights. Within this framework, two positive definite kernels are
 53 proposed that assign robust weights to mitigate the adverse effect of outliers in the local neighborhood by estimating the density of the response at the local locations. Note the proposed
 54 framework does not preclude the use of robust loss functions, robust bandwidth selectors and
 55 standardization techniques. In addition the method is implemented in the Python programming
 56 language and is made publicly available. Experimental results on synthetic benchmarks demon-
 57 strate that the proposed method achieves competitive results and generally performs better than
 58 LOESS/LOWESS using only a single training iteration.

60 The remainder of the paper is organized as follows: In section 2, a brief overview of the
 61 mathematical formulation of local polynomial regression is provided. In section 3, a framework
 62 for robust weights as well as specific robust positive definite kernels are proposed. Section 4
 63 provides an analysis of the estimator and a discussion of its properties. In section 5, implemen-
 64 tation notes and experimental results are provided. Finally, in section 6, the paper concludes with
 65 directions for future research.

66 2. Local polynomial regression

67 This section provides a brief overview of local polynomial regression and establishes the no-
 68 tation subsequently used. Let (X, Y) be a random pair and $\mathcal{D}_T = \{(X_i, Y_i)\}_{i=1}^T \subseteq \mathcal{D}$ be a training
 69 set comprising a sample of T data pairs. Suppose that $(X, Y) \sim f_{XY}$ a continuous density and
 70 $X \sim f_X$ the marginal distribution of X . Let $Y \in \mathbb{R}$ be a continuous response and assume a model
 71 of the form $Y_i = m(X_i) + \epsilon_i$, $i \in 1, \dots, T$ where $m(\cdot) : \mathbb{R}^d \rightarrow \mathbb{R}$ is an unknown function and ϵ_i
 72 are independently distributed error terms having zero mean representing random variability not
 73 included in X_i such that $\mathbb{E}[Y|X=x] = m(x)$. There are no global assumptions about the function
 74 $m(\cdot)$ other than that it is smooth and that locally it can be well approximated by a low degree
 75 polynomial as per Taylor's theorem. Local polynomial regression is a class of nonparametric re-
 76 gression methods that estimate the unknown regression function $m(\cdot)$ by combining the classical
 77 least squares method with the versatility of non-linear regression. The local p -th order Taylor
 78 expansion for $x \in \mathbb{R}$ near a point X_i yields:

$$m(X_i) \approx \sum_{j=0}^p \frac{m^{(j)}(x)}{j!} (x - X_i)^j := \sum_{j=0}^p \gamma_j(x)(x - X_i)^j \quad (1)$$

79 To find an estimate $\hat{m}(x)$ of $m(x)$ the low-degree polynomial (1) is fitted to the N nearest neighbors
 80 using weighted least squares such to minimize the empirical loss $\mathcal{L}_{lpr}(\cdot; \mathcal{D}_N, h)$:

$$\mathcal{L}_{lpr}(x; \mathcal{D}_N, h) := \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \gamma_j(x)(x - X_i)^j \right)^2 K_h(x - X_i) \quad (2)$$

$$\hat{\gamma}(x) := \min_{\gamma(x)} \mathcal{L}_{lpr}(x; \mathcal{D}_N, h) \quad (3)$$

81 Where $\gamma, \hat{\gamma} \in \mathbb{R}^{p+1}$; $K_h(\cdot)$ is a scaled kernel, $h \in \mathbb{R}_{>0}$ is the bandwidth parameter and $\mathcal{D}_N \subseteq \mathcal{D}_T$
 82 is the subset of N nearest neighbors of x in the training set where the distance is measured on

83 the predictors only. Having computed $\hat{\gamma}(x)$ the estimate of $\hat{m}(x)$ is taken as $\hat{\gamma}(x)_1$. Note the
 84 term kernel carries here the meaning typically used in the context of nonparametric regression
 85 i.e. a non-negative real-valued weighting function that is typically symmetric, unimodal at zero,
 86 integrable with a unit integral and whose value is non-increasing for the increasing distance
 87 between the X_i and x . Higher degree polynomials and smaller N generally increase the variance
 88 and decrease the bias of the estimator and vice versa [2]. For derivation of the local constant and
 89 local linear estimators for the multidimensional case see [6].

90 3. Robust weights with similarity kernels

91 The main idea presented is to generalize the kernel function used in equation (2) to produce
 92 robust weights. This is achieved by using a similarity kernel function defined on the data domain
 93 $K_{\mathcal{D}} : \mathcal{D}^2 \rightarrow \mathbb{R}_+$ that enables weighting each point and incorporating information on the data in
 94 the local neighborhood in relation to the local regression target.

$$\mathcal{L}_{rsk}(x, y; \mathcal{D}_N, H) := \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x, y)(x - X_i)^j \right)^2 K_{\mathcal{D}}((x, y), (X_i, Y_i); H) \quad (4)$$

$$\hat{\beta}(x, y; \mathcal{D}_N, H) := \min_{\beta(x, y)} \mathcal{L}_{rsk}(x, y; \mathcal{D}_N, H) \quad (5)$$

95 Where H is the set of bandwidth parameters. There are many possible choices for such a
 96 similarity kernel to be defined within this general framework. However, used as a local weighting
 97 function, such a kernel should have the following attributes:

- 98 1. Non-negative, $K_{\mathcal{D}}((x, y), (x', y')) \geq 0$.
- 99 2. Symmetry in the inputs, $K_{\mathcal{D}}((x, y), (x', y')) = K_{\mathcal{D}}((x', y'), (x, y))$.
- 100 3. Tending toward decreasing as the distance in the predictors increases. That is, given a
 101 similarity function on the response $s(\cdot, \cdot) : \mathbb{R}^2 \rightarrow \mathbb{R}_+$, if $s(y, y')$ indicates high similarity
 102 the weight should decrease as the distance between the predictors grows, $s(y, y') > \alpha \implies$
 103 $K_{\mathcal{D}}((x, y), (x + u, y')) \geq K_{\mathcal{D}}((x, y), (x + v, y')) \quad \forall \|u\| \leq \|v\|$ and some $\alpha \in \mathbb{R}_+$.

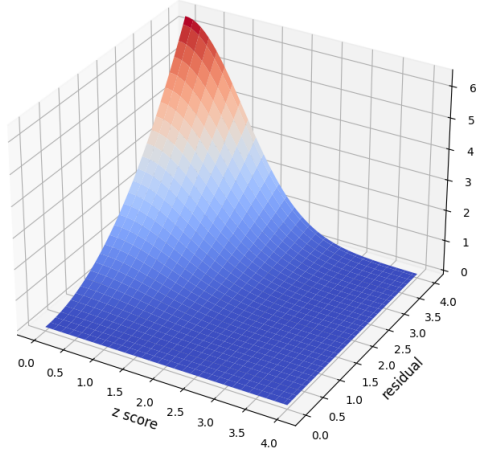
104 In this work two such useful positive definite kernels are proposed. Similarly to the usual
 105 kernels used in (2), these tend to diminish as the distance between the explanatory variables
 106 increases to model stronger relationship between closer points. In addition, the weights produced
 107 by the kernels also model the "importance" of the pair (x, y) . This is useful for example to down-
 108 weight outliers to mitigate their adverse effect on the ordinary least square based regression.
 109 Formally let $K_{\mathcal{D}}$ be defined as:

$$K_{\mathcal{D}}((x, y), (x', y'); H_1, H_2) = K_1(x, x'; H_1)K_2((x, y), (x', y'); H_2) \quad (6)$$

110 Where $K_1 : \mathbb{R}^d \times \mathbb{R}^d \rightarrow \mathbb{R}_+$ and $K_2 : \mathcal{D}^2 \rightarrow \mathbb{R}_+$ are positive definite kernels and H_1, H_2 are
 111 the sets of bandwidth parameters. The purpose of K_1 is to account for the distance between a
 112 neighbor to the local regression target and therefore may be chosen as any of the kernel functions
 113 that are typically used in equation (2). The role of K_2 is described now in more detail as this

114 is the main idea proposed in this work. Using K_2 , the method performs robust regression by
 115 detecting local outliers in an unsupervised manner and assigns them with lower weights. There
 116 are many methods that could be employed to estimate the extent to which a data point is a local
 117 outlier however in this work it is estimated in one of the following two ways.

Figure 1: Loss function, assuming a standard quadratic function of the residual, a standard normal density for K_2 and excluding the K_1 distance kernel scaling.



118 *Conditional density*

The first proposed method for K_2 is proportional to the estimated localized conditional marginal distribution of the response variable at the location:

$$K_2((x, y), (x', y'); H_2) = \hat{f}(y | x; H_2)\hat{f}(y' | x'; H_2) \quad (7)$$

The nonparametric conditional density estimation is performed using the Parzen–Rosenblatt window (kernel density estimator):

$$\hat{f}(y | x; H_2) = \hat{f}(x, y; H_2)/\hat{f}(x; H_2) \quad (8)$$

$$= \hat{f}(v; \mathbf{H}_v)/\hat{f}(x; \mathbf{H}_x) \quad (9)$$

$$= \frac{|\mathbf{H}_x|^{1/2} \sum_{i=1}^N K_v(\mathbf{H}_v^{-1/2}(v - V_i))}{|\mathbf{H}_v|^{1/2} \sum_{i=1}^N K_x(\mathbf{H}_x^{-1/2}(x - X_i))} \quad (10)$$

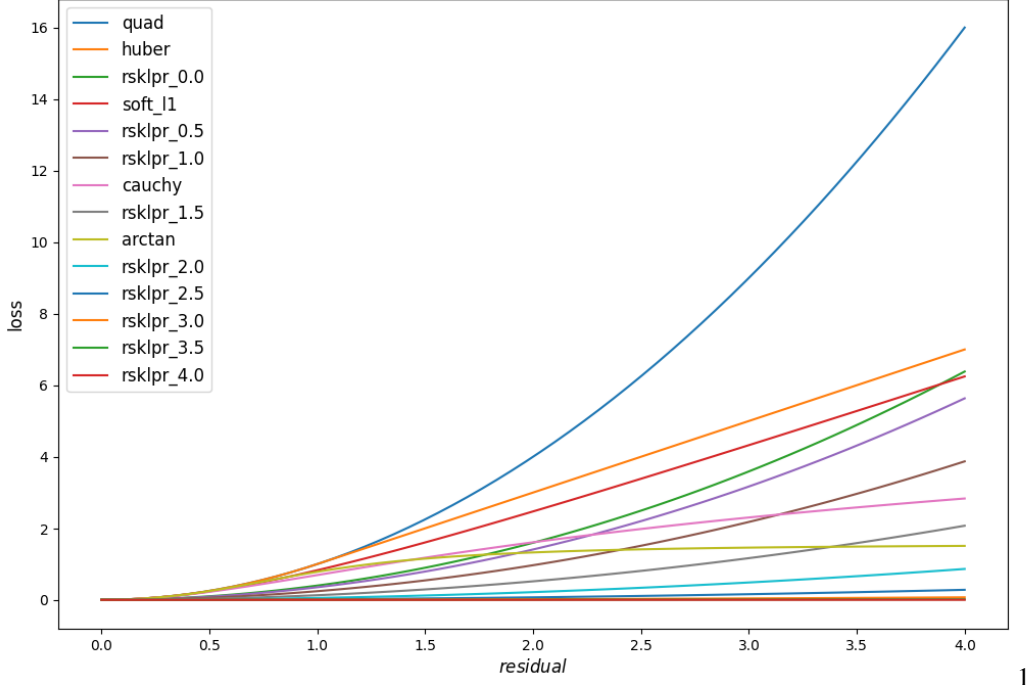
119 Where $v = [x, y] \in \mathbb{R}^{d+1}$ is the concatenated vector of the predictors and the response; and $\mathbf{H}_v, \mathbf{H}_x$
 120 are bandwidth matrices.

121 *Joint density*

The second proposed kernel is proportional to the joint distribution of the random pair, this could be useful for example to also down-weight high leverage points:

$$K_2((x, y), (x', y'); H_2) = \hat{f}(x, y; H_2)\hat{f}(x', y'; H_2) \quad (11)$$

Figure 2: The plot illustrates the proposed loss function, a number of common robust losses and the standard quadratic residual loss for comparison. It is assumed that that K_2 is equivalent to the standard normal density and the K_1 distance kernel scaling is excluded. The numbers appended to "rsklpr" indicate how many standard deviations away from the mean the density is calculated. It is evident that the loss is heavily attenuated in regions of low density.



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122 Where the joint density can be estimated using the same aforementioned approach.

123

124 Regardless of the choice of kernel, the hyperparameters of this model are similar in essence
 125 to the standard local polynomial regression and comprise the span of included points, the kernels
 126 and their associated bandwidths. Note that this estimator can be replaced with other robust
 127 density estimators and better results are anticipated by doing so however exploring this option is
 128 left for future work.

129 **4. Properties**

130 This section discusses some properties of the estimator. Note the notation in this section is
 131 simplified by excluding explicit mentions of D_N and H , however the analysis is conditional on
 132 the nearest neighbors in the sample, D_N .

133 *4.1. Invariance to y at the regression location and simplification of the objective*

134 The objective (5) is invariant to the value of y at the location (x, y) for the proposed similarity
 135 kernels.

136 *Proof:* The optimization is invariant to the scale of the objective function. Therefore:

$$\hat{\beta}(x, y) := \min_{\beta(x,y)} \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x, y)(x - X_i)^j \right)^2 K_{H_1}(x - X_i) \hat{f}(x, y) \hat{f}(X_i, Y_i) \quad (12)$$

$$= \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x, y)(x - X_i)^j \right)^2 K_{H_1}(x - X_i) \hat{f}(X_i, Y_i) \quad (13)$$

¹³⁷ The equality holds because $\hat{f}(x, y)$ is a constant scalar that uniformly scales the weights.
¹³⁸ Since the objective is now independent of y , it follows that:

$$\hat{\beta}(x, y) := \min_{\beta(x)} \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x)(x - X_i)^j \right)^2 K_{H_1}(x - X_i) \hat{f}(X_i, Y_i) \quad (14)$$

$$:= \hat{\beta}(x) \quad \forall y \quad (15)$$

¹³⁹ This simplification enables more efficient calculations of the estimator because the dependence on y is removed from the objective function. Note that $\hat{f}(X_i, Y_i)$ can also be replaced with
¹⁴⁰ $\hat{f}(Y_i | X_i)$ with similar results.
¹⁴¹

¹⁴² 4.2. Weighted arithmetic mean of the standard LPR

¹⁴³ The proposed estimator is equivalent to the weighted arithmetic mean of the terms in the
¹⁴⁴ standard LPR loss (2), with weights $w_i = \hat{f}(X_i, Y_i)$.

¹⁴⁵ *Proof:* Since the optimization is invariant to scaling, we have:

$$\hat{\beta}(x) := \min_{\beta(x)} \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x)(x - X_i)^j \right)^2 K_{H_1}(x - X_i) \hat{f}(X_i, Y_i) \quad (16)$$

$$= \min_{\beta(x)} \left(\sum_{i=1}^N \hat{f}(X_i, Y_i) \right)^{-1} \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x)(x - X_i)^j \right)^2 K_{H_1}(x - X_i) \hat{f}(X_i, Y_i) \quad (17)$$

$$= \min_{\beta(x)} \left(\sum_{i=1}^N w_i \right)^{-1} \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x)(x - X_i)^j \right)^2 K_{H_1}(x - X_i) w_i \quad (18)$$

¹⁴⁶ The normalization by $\sum_{i=1}^N w_i$ shows the equivalence to the weighted arithmetic mean, with
¹⁴⁷ the weights $w_i = \hat{f}(X_i, Y_i)$.

¹⁴⁸ 4.3. Asymptotic degeneration of the estimator to the standard LPR

¹⁴⁹ Asymptotically, the proposed estimator degenerates to the standard LPR when the weights
¹⁵⁰ w_i are uncorrelated with the standard LPR terms. Formally, as $N \rightarrow \infty$, $\hat{\beta}(x) \rightarrow \hat{\gamma}(x)$, where
¹⁵¹ $\hat{\gamma}(x)$ is the standard LPR estimator, and the condition that $\left(Y - \sum_{j=0}^p \beta_j(x)(x - X)^j \right)^2 K_{H_1}(x - X)$

¹⁵² and $\hat{f}(X, Y)$ are uncorrelated holds. It is assumed that (X_i, Y_i) are independent and identically
¹⁵³ distributed (i.i.d.) random variables and that $\hat{f}(X, Y) > 0$ almost everywhere.

¹⁵⁴ *Proof:* Define

$$g(X, Y) := \left(Y - \sum_{j=0}^p \beta_j(x)(x - X)^j \right)^2 K_{H_1}(x - X),$$

it follows that:

$$\hat{\beta}(x) := \min_{\beta(x)} \left(\sum_{i=1}^N \hat{f}(X_i, Y_i) \right)^{-1} \sum_{i=1}^N g(X_i, Y_i) \hat{f}(X_i, Y_i) \quad (19)$$

$$= \min_{\beta(x)} \left(\frac{1}{N} \sum_{i=1}^N \hat{f}(X_i, Y_i) \right)^{-1} \left(\frac{1}{N} \sum_{i=1}^N g(X_i, Y_i) \hat{f}(X_i, Y_i) \right) \quad (20)$$

As $N \rightarrow \infty$, by the law of large numbers, we obtain:

$$\left(\frac{1}{N} \sum_{i=1}^N \hat{f}(X_i, Y_i) \right)^{-1} \xrightarrow{a.s.} \frac{1}{\mathbb{E}[\hat{f}(X, Y)]} \quad (21)$$

$$\frac{1}{N} \sum_{i=1}^N g(X_i, Y_i) \hat{f}(X_i, Y_i) \xrightarrow{a.s.} \mathbb{E}[g(X, Y) \hat{f}(X, Y)] \quad (22)$$

Assuming $\mathbb{E}[\hat{f}(X, Y)] \neq 0$, it follows that:

$$\hat{\beta}(x) \xrightarrow{a.s.} \min_{\beta(x)} \frac{\mathbb{E}[g(X, Y) \hat{f}(X, Y)]}{\mathbb{E}[\hat{f}(X, Y)]} \quad (23)$$

If $g(X, Y)$ and $\hat{f}(X, Y)$ are uncorrelated, then:

$$\mathbb{E}[g(X, Y) \hat{f}(X, Y)] = \mathbb{E}[g(X, Y)] \mathbb{E}[\hat{f}(X, Y)] \quad (24)$$

$$\hat{\beta}(x) \xrightarrow{a.s.} \min_{\beta(x)} \mathbb{E}[g(X, Y)] \quad (25)$$

Therefore, as $N \rightarrow \infty$:

$$\hat{\beta}(x) \xrightarrow{a.s.} \min_{\beta(x)} \mathbb{E} \left[\left(Y - \sum_{j=0}^p \beta_j(x)(x - X)^j \right)^2 K_{H_1}(x - X) \right] \quad (26)$$

¹⁵⁵ This is the same objective minimized by the standard LPR estimator in the asymptotic sense.
¹⁵⁶ Thus, the proposed estimator degenerates to the standard LPR as $N \rightarrow \infty$, provided that $g(X, Y)$
¹⁵⁷ and $\hat{f}(X, Y)$ are uncorrelated.

158 4.4. Relationship to Kernel Methods and RKHS

159 In this subsection, the relationship of the proposed method to kernel methods and Reproducing Kernel Hilbert Spaces (RKHS) is explored. The use of positive definite kernels in defining
 160 the weights $K_{\mathcal{D}}$ allows the proposed estimator to be interpreted within the RKHS framework,
 161 providing deeper insights into its properties and connections to existing kernel-based methods.
 162

163 Recall that in the proposed method, the weights in the loss function (4) are defined using a
 164 compound positive definite kernel $K_{\mathcal{D}}$ on the data domain \mathcal{D} :

$$\mathcal{L}_{\text{rsk}}(x, y; \mathcal{D}_N, H) := \sum_{i=1}^N \left(Y_i - \sum_{j=0}^p \beta_j(x, y)(x - X_i)^j \right)^2 K_{\mathcal{D}}((x, y), (X_i, Y_i); H). \quad (27)$$

165 As per equation (6), the kernel $K_{\mathcal{D}}$ is defined as a product of two positive definite kernels:

$$K_{\mathcal{D}}((x, y), (x', y'); H_1, H_2) = K_1(x, x'; H_1) \cdot K_2((x, y), (x', y'); H_2), \quad (28)$$

166 where K_1 is a kernel function depending only on the predictors x and x' , typically chosen
 167 as the traditional distance-based kernel used in local polynomial regression, and K_2 is a kernel
 168 function that incorporates both predictors and responses.

169 Since $K_{\mathcal{D}}$ is a product of positive definite kernels, it is itself a positive definite kernel. Therefore,
 170 there exists a feature mapping $\phi : \mathcal{D} \rightarrow \mathcal{H}$, where \mathcal{H} is a Hilbert space, such that:

$$K_{\mathcal{D}}((x, y), (x', y')) = \langle \phi(x, y), \phi(x', y') \rangle_{\mathcal{H}}. \quad (29)$$

171 Thus, the weights $K_{\mathcal{D}}((x, y), (X_i, Y_i))$ can be interpreted as inner products in the feature space
 172 \mathcal{H} . Consequently, the loss function (27) can be viewed as a weighted least squares problem
 173 where the weights are determined by the similarity between the feature representations of the
 174 data points and the point of interest.

175 Furthermore, consider the role of the Kernel Density Estimator (KDE) in the proposed method.
 176 The KDE at a point (x, y) is given by:

$$\hat{f}(x, y) = \frac{1}{N} \sum_{i=1}^N K_2((x, y), (X_i, Y_i); H_2). \quad (30)$$

177 Note that the KDE uses K_2 , not the full kernel $K_{\mathcal{D}}$, since K_2 is the kernel used in the density
 178 estimation of the joint (or conditional) distribution involving both x and y .

179 Since K_2 is a positive definite kernel, there exists a feature mapping $\psi : \mathcal{D} \rightarrow \mathcal{G}$, where \mathcal{G} is
 180 another Hilbert space, such that:

$$K_2((x, y), (x', y')) = \langle \psi(x, y), \psi(x', y') \rangle_{\mathcal{G}}. \quad (31)$$

181 Therefore, the KDE at (x, y) can be expressed in terms of inner products in the feature space
 182 \mathcal{G} :

$$\hat{f}(x, y) = \frac{1}{N} \sum_{i=1}^N \langle \psi(x, y), \psi(X_i, Y_i) \rangle_{\mathcal{G}}. \quad (32)$$

183 This expression shows that the KDE at (x, y) is proportional to the inner product between the
 184 feature mapping $\psi(x, y)$ and the mean of the feature mappings of the data:

$$\hat{v}_{\psi} = \frac{1}{N} \sum_{i=1}^N \psi(X_i, Y_i), \quad (33)$$

185 so that:

$$\hat{f}(x, y) = \langle \psi(x, y), \hat{v}_{\psi} \rangle_{\mathcal{G}}. \quad (34)$$

186 This interpretation shows that the KDE measures how closely the feature representation
 187 $\psi(x, y)$ of a point (x, y) aligns with the average feature representation \hat{v}_{ψ} of the data in the space
 188 induced by K_2 . In the proposed method, this alignment influences the weights in the regression,
 189 as the density estimates $\hat{f}(x, y)$ or $\hat{f}(Y_i | X_i)$ derived from K_2 directly affect the overall weights
 190 $K_{\mathcal{D}}((x, y), (X_i, Y_i))$. This interplay underpins the robustness and adaptability of the proposed
 191 method.

192 By leveraging positive definite kernels for defining $K_{\mathcal{D}}$, the method inherently operates within
 193 the RKHS framework, where weights represent similarities in feature space. This perspective
 194 highlights the connection between the kernel-based weighting and the feature mappings, offering
 195 insights into the estimator's flexibility and robustness.

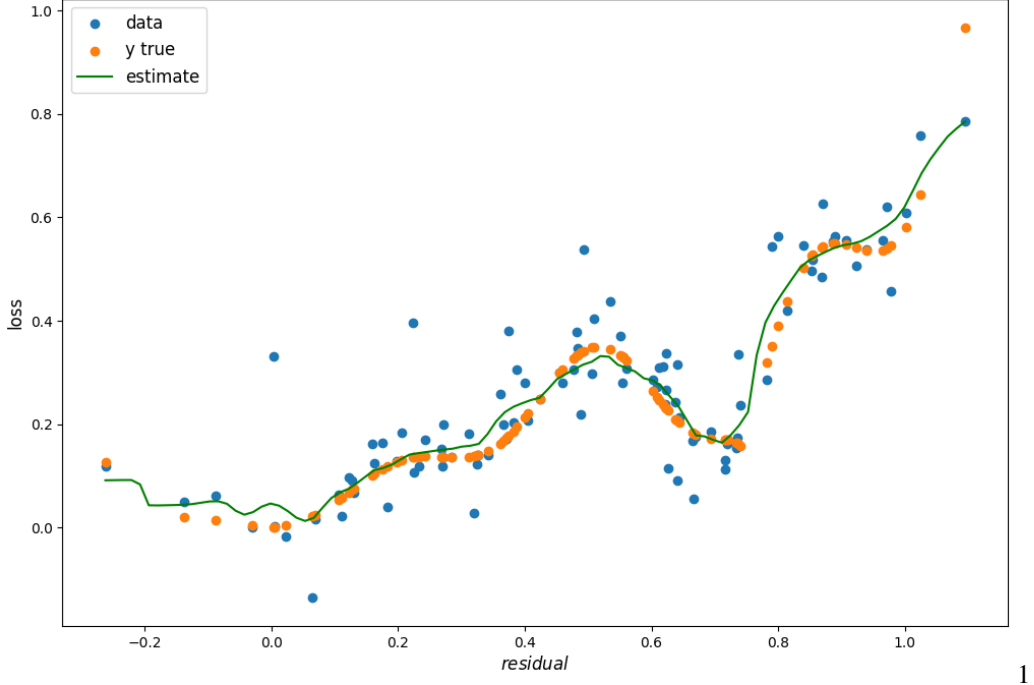
196 5. Experiments and Implementation Notes

197 The proposed method was implemented in Python. Following [3], distances between pairs
 198 in each neighborhood are normalized to the range $[0, 1]$. For the kernel function $K_1(x, x'; H_1)$, a
 199 simple Laplacian kernel $e^{-\|x-x'\|}$ was used, as it demonstrated more efficient and consistent em-
 200 pirical performance than the tricube kernel suggested in [3]. For density estimation, a factorized
 201 multidimensional KDE with scaled Gaussian kernels was applied. Five bandwidth estimation
 202 methods were tested: Scott's rule [12], the Normal Reference rule, global Least Squares Cross-
 203 Validation (LSCV), local LSCV, and local Modified Least Squares Cross-Validation (MLCV). In
 204 some experiments, the bandwidth for the predictor kernel was empirically adjusted as a simple
 205 function of the window size.

206 For computational efficiency, certain calculations were omitted because the local regression
 207 in equation (5) is invariant to the scale of the weights. This includes excluding scaling constants
 208 fixed within a neighborhood for a specific local regression target, such as those in computing
 209 $\hat{f}(y | x)$ and $\hat{f}(x, y)$ in equations (7) and (11), respectively.

210 Experiments were conducted using a variety of synthetic benchmarks to evaluate the perfor-
 211 mance of the method's linear and quadratic variants against other local polynomial regression
 212 methods. These include LOWESS and iterative robust LOWESS [13], local linear and local con-
 213 stant kernel regression [13], local quadratic regression [14], and radial basis function networks
 214 [14].

Figure 3: Regression example of synthetically generated 1D data with heteroscedastic noise. Additional experimental results and demonstrations including multivariate settings and bootstrap based confidence intervals are available at <https://nbviewer.org/github/yaniv-shulman/rsklpr/tree/main/src/experiments/> as interactive Jupyter notebooks [1]



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215 The experimental setups included several non-linear synthetic curves and planes with added
 216 noise, representing dense and sparse data, homoscedastic and heteroscedastic noise characteris-
 217 tics, and different neighborhood sizes. The results indicate that no single method universally
 218 outperforms the others; performance varies by setting. However, the proposed method ex-
 219 hibited competitive performance overall, delivering the best results across numerous settings,
 220 particularly in heteroscedastic environments. It also generally outperformed its direct counter-
 221 parts, LOESS/LOWESS and quadratic LPR, with just a single iteration. Moreover, the proposed
 222 method showed lower sensitivity to neighborhood size, resulting in reduced variance. This sta-
 223 bility makes it an attractive option, especially when choosing hyperparameters without ground
 224 truth data.

225 The complete experimental results are available as interactive Jupyter notebooks at <https://nbviewer.org/github/yaniv-shulman/rsklpr/tree/main/src/experiments/> [1].

227 6. Future Work and Research Directions

228 This work introduces a new robust variant of Local Polynomial Regression (LPR), opening
 229 several avenues for further exploration and refinement. Since the proposed method generalizes
 230 the traditional LPR, there are opportunities to replace certain standard components in equation
 231 (5) with more robust alternatives. These could include approaches such as robust methods for

232 bandwidth selection or substituting the conventional quadratic residual function with alternatives
233 better suited for handling outliers.

234 Additionally, further development of this framework may involve exploring different kernel
235 functions K_D and assessing how robust density estimators influence overall performance. Ex-
236 tending the method within the RKHS framework presents another valuable direction. This could
237 allow for the introduction of a regularization term in the loss function, enhancing control over
238 estimator smoothness and mitigating the risk of overfitting. Through these future directions, the
239 robustness and adaptability of the proposed method could be substantially advanced.

240 **References**

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