

The Economics of European Regions: Theory, Empirics, and Policy

Dipartimento di Economia e Management

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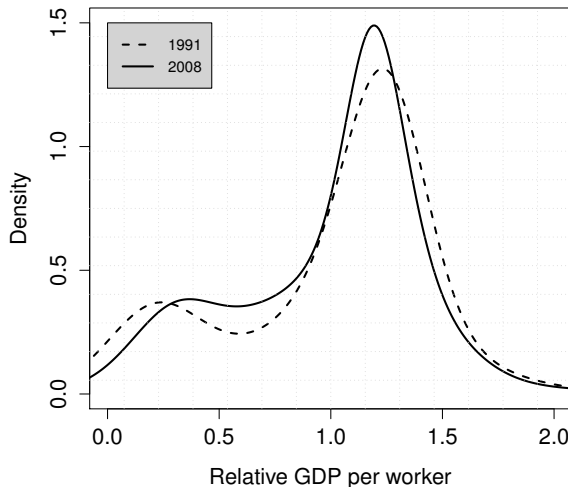
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Distribution of Regional GDP per Worker

	1991	2008
Gini	0.25	0.23
BIPOL	0.83	0.78



Estimate of The Density Function

Let be x a continuous random variable and f its probability density function (pdf).

The pdf characterizes the distribution of the random variable x since it tells “how x is distributed”.

Moreover, from pdf it is possible to calculate the mean and the variance (it they exists) of x and the probability that x takes on values in a given interval.

Histogram

Histograms are nonparametric estimates of an *unknown density function*, $f(x)$, **without assuming any well-known functional form**. In order to build an histogram, you have to:

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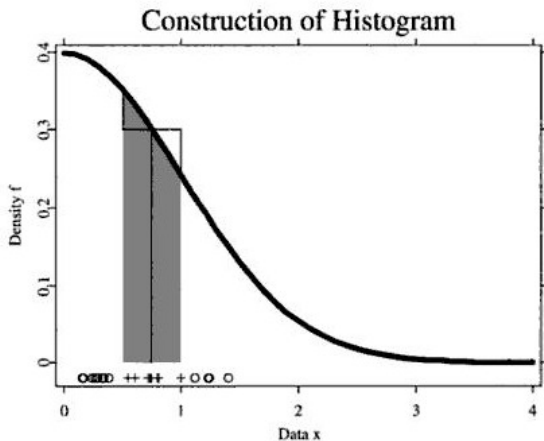
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- 2 count how many observations fall into each bin (n_j for each bin j);
- 3 for each bin divide the frequency by the sample size n and the binwidth h , to get the relative frequencies $f_j = \frac{n_j}{nh}$

Histogram: Cont.



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→ we need to find an “optimal” binwidth, which represents an optimal compromise.

Histogram: Cont.

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- 1 each observation x in $[m_j - \frac{h}{2}, m_j + \frac{h}{2})$ is estimated by the same value, $\hat{f}_h(m_j)$, where m_j is the center of the bin;
- 2 $f(x)$ is estimated using the observations that fall in the interval containing x , and that receive the same weight in the estimation. That is, for $x \in B_j$,

$$\hat{f}_h(m_j) = \frac{1}{nh} \sum_{i=1}^n I(X_i \in B_j),$$

where I is the indicator function.

Nonparametric density estimation

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- Density estimation is a generalization of the histogram.
- It is based on **Kernel functions**: estimate $f(x)$ using the observations that fall into an interval around x , which (typically) receive decreasing weight the further they are from x .

Kernel functions

Consider the *uniform* kernel function, which assigns *the same weight to all observations in an interval* of length $2h$ around observation x , $[x - h, x + h)$:

$$\hat{f}_h(x) = \frac{1}{2nh} \#\{X_i \in [x - h, x + h)\}$$

can be obtained by means of a kernel function $K(u)$ such that:

$$K(u) = \frac{1}{2} I(|u| \leq 1)$$

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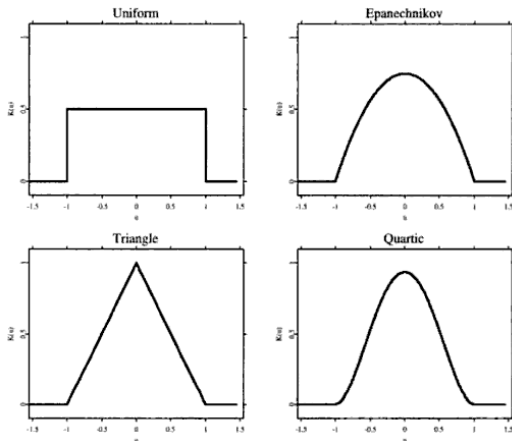
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- For each observation that falls into the interval $[x - h, x + h)$ the indicator function takes on value 1
- Each contribution to the function is weighted equally no matter how close the observation X_i is to x

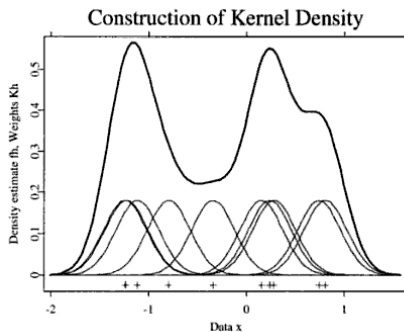
Kernel functions: Cont.

A Kernel function in general (e.g. Epanechnikov, Gaussian, etc), assigns higher weights to observations in $[x - h, x + h]$ closer to x .



Kernel density

A kernel density estimation appears as a sum of bumps: at a given x , the value of $\hat{f}_h(x)$ is found by vertically summing over the “bumps”:



$$\hat{f}_h(x) = \sum_{i=1}^n \frac{1}{nh} K\left(\frac{x - X_i}{h}\right) = \sum_{i=1}^n \frac{1}{n} \underbrace{K_h(x - X_i)}_{\text{"rescaled kernel function"}}$$

Properties of Kernel density estimator

Same problems found for the histogram, that is the bias and the variance depending on h , also hold for the Kernel:

$$Bias\{\hat{f}_h(x)\} = E\{\hat{f}_h(x)\} - f(x);$$

that positively depends on h ;

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So, how do we choose h given the trade-off between bias and variance?

Choosing the bandwidth h

- (a) Define MSE (mean squared error)

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- (c) Define AMISE (an approximation of MISE) → still h_{opt} depends on the unknown $f(x)$, in particular on its second derivative $f''(x)$.
- (d) One possibility is a plug-in method suggested by Silverman, and consists in assuming that the unknown function is a Gaussian density function (whose variance is estimated by the sample variance). In this case h_{opt} has a simple formulation, and can be defined as a rule-of-thumb bandwidth.

Adaptive Kernel

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- But we can get a better estimate by allowing the window width of the kernels to vary from one point to another.
- In particular, a natural way to deal with long-tailed densities is to use a broader kernel in regions of low density.
- Thus an observation in the tail would have its mass smudged out over a wider range than one in the main part of the distribution.

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- The adaptive kernel approach copes with this problem by means of a two-stage procedure:
 - ① get an initial estimate to have a rough idea of the density
 - ② use the former density to get a pattern of bandwidths corresponding to various observations to be used in a second estimate

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Step 1 Find a *pilot estimate* $\tilde{f}(x)$ that satisfies $\tilde{f}(x_i) > 0 \forall i$

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where g is the geometric mean of the $\tilde{f}(x_i)$, $\log g = n^{-1} \sum \log \tilde{f}(x_i)$;
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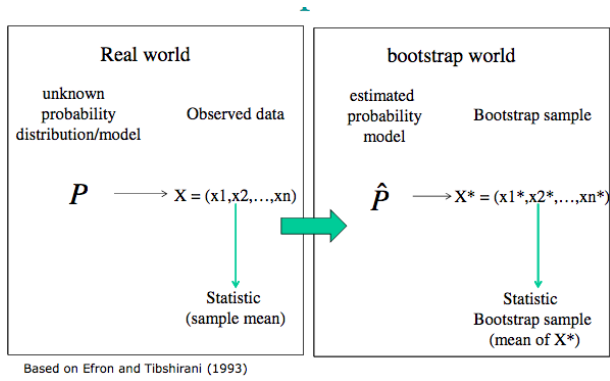
$$\hat{f}(x) = nh^{-1} \sum \lambda_i^{-1} K\{h^{-1}\lambda_i^{-1}(x - X_i)\} \quad (2)$$

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- Justification:
 - 1 If the sample is representative for the population, the sample distribution (empirical distribution) approaches the population (theoretical) distribution if n increases;
 - 2 If the number of resamples (B) from the original sample increases, the bootstrap distribution approaches the sample distribution.

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The distribution of $\hat{f}^*$ about $\hat{f}$ can therefore be used to mimic the distribution of $\hat{f}$ about f , that is it can be used to calculate the confidence intervals for estimates.

References

Histogram and Density Estimation

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- Estimate: Chapter 1
- Inference (confidence bands): Chapter 2

Adaptive Density Estimation

Silverman, B.W. (1986). Density estimation for statistics and data analysis. *CRC press*.

- Estimate: Chapter 5.3