REGRESSION PROBLEMS

In a regression problem, we have a set of evidence variables X[I], ..., X[D] and a response variable Y that we want to predict.

For instance, let's say we want to predict chalesterol level given age and weight:

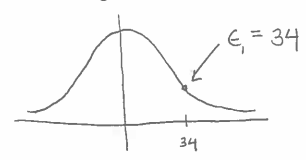
We have N training examples, each consisting of a vector X: and a scalar Y:, to learn from.

- 2) We assume the response variable is generated using the following steps:
 - we start with a vector of weights (one weight per evidence variable), e.g.

$$W = \begin{bmatrix} W[1] \\ W[2] \end{bmatrix} = \begin{bmatrix} -50 \\ 2 \\ W[3] \end{bmatrix}$$

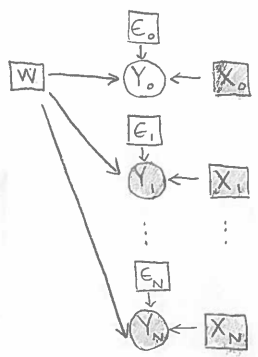
- we compute the weighted linear combination of the evidence variables (this will result in a single number):

we sample a random number \in from a distribution φ e.g. \in , \propto Normal $(0, \sigma^2)$



- we compute the response variable as a function p of of W^TX and E, e.g.

3) Given this setup, we want to predict the value of an Unobserved response variable Yo given its observed evidence variables Xo and our training data. In its simplest formulation, the causal diagram looks as follows:



Legend:

[]: background

O: endogenous

shaded: observed

We assume a probability distribution P over the background variables such that all background variables are marginally independent, i.e.

 $P(w, \epsilon_0, ..., \epsilon_n, \chi_0, ..., \chi_N) = P_w(w)P_{\epsilon}(\epsilon_0)...P_{\epsilon}(\epsilon_n).P_{\epsilon}(\chi_0)...P_{\epsilon}(\chi_n)$ Moreover we assume that all variables ϵ_n are drawn from the same distribution P_{ϵ} and all variables χ are

drawn from the same distribution Px.

4) What's the deal with the E's?

We'll call these stochastic terms. The idea is to allow some softness around the deterministic point wixi. In other words, just because w x: = 148 for age = 24 and weight = 150, that doesn't mean we want the model to claim that EVERY person whose age is 24 and whose weight is 150 must have a chalesterol level of EXACTLY 148.

Instead, we want their cholesterol levels to be disposed around 148.

5) Suppose we choose Pe ~ Normal (0, 02) for some fixed variance of, and suppose we choose $P(z, \epsilon) = z + \epsilon$. This is what we did in (2), which gave us a chalesterol level of 148+34=182 for a 24-year-old, 1501b subject.

DETERMINISTIC TERM WTX = [-50 2 1][17

= 148

STOCHASTIC TERM

REGRESSION PROBLEMS

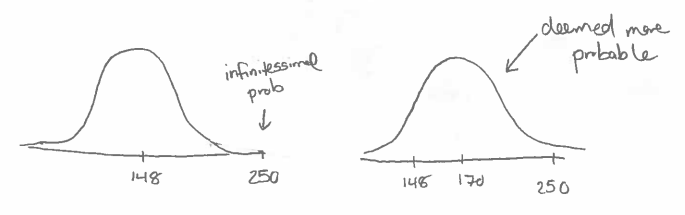
6) These choices:

$$\begin{cases}
P_{\epsilon} \sim \text{Normal}(0, \sigma^{2}) \\
P(Z, \epsilon) = Z + \epsilon
\end{cases}$$

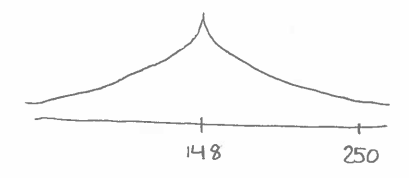
give us the ordinary linear regression model

F) But it may not be the best choice. One possible downside of the normal distribution is that it has rapidly diminishing tails, so a normal distribution centered at 148 may give a nearly infinitessimal probability to 250.

In practice, this means that using the normal distribution makes regression very sensitive to cuttiers and noise, because even you have a thousand 24-year-olds with cholesterol 148, just one 24-year-old with cholesterol 250 can cause the learned mean to shift significantly upwards, because this will be deemed more probable than even one parson with cholesterol 250:



(8) If this is a problem for you, you can use a "heavy-tailed" distribution (i.e. they diminish in probability much more slowly). One example is the Laplace distribution.



Often, however, such distributions are not as computationally convenient.

9) This chaice:

\[\begin{aligned} P_{\infty} & \taplace(0, b) \\ P(\pi, \infty) = \pi + \infty \]

gives us the "robust" linear regression model

REGRESSION PROBLEMS

10) Another common choice for Pc and p comes into play when the response variable is Boolean-valued, e.g.:

$$X$$
 (evidence vars)
 $X[1]$ $X[2]$ $X[3]$
(offset) (age) (weight)
 $X_1 = \begin{bmatrix} 1 & 24 & 150 \end{bmatrix}$
 $X_2 = \begin{bmatrix} 1 & 50 & 164 \end{bmatrix}$

$$X_{N} = \begin{bmatrix} 1 & 22 & 205 \end{bmatrix}$$

Y (response var)

(has high cholesterol)

O = Y₁

1 = Y₂

;

(1) We could use ordinary linear regression, but then we end up predicting values in the range (-00,00), rather than restricting ausselves to the set 20, 13.

Another chaice:

$$P_{\epsilon} \sim \text{Constant}(0,1)$$

$$P(z,\epsilon) = 1$$

$$(1+e^{-z})^{-1} < \epsilon (z)$$

gives us the famous logistic regression model

REGRESSION FROBLEMS
12 Let's unpack this: "we generate a random number & between O and 1 from Constant(0,1) 1
$\boxed{\mathbb{V}} \longrightarrow \boxed{\mathbb{V}}_{0} \longleftarrow \boxed{\mathbb{X}}_{0}$
(ii) We compute a threshold
$r = \frac{1}{1 + e^{-\omega^T x_n}}$ (note: $0 < \gamma < 1$)
(iii) $Y_n = \begin{cases} 1 & \text{if } \epsilon < \tau \\ 0 & \text{otherwise} \end{cases}$
In other words, the probability that $Y_n = 1$ is equal to $\frac{1}{1+e^{-w^T \times n}}$
(13) This function, $f(z) = (1 + e^{-z})^{-1}$, looks like this: when z is negative, $f(z) < \frac{1}{2}$ when z is positive, $f(z) > \frac{1}{2}$
is called the louistic function (1)

or le signoid).

(H) No matter what regression model we're using, we typically want to predict the value of the unobserved response variable Yo given its observed evidence variables Xo (recall Xo is a vector) and our other observations (i.e. Xn, Yn for $n \ge 1$). In other words, we want argmax $P(y_0 \mid x_0, x_1, ..., x_N, y_1, ..., y_N)$.

15) The exact computation would be:

argmax P(y. 1x., x, x, y, y, y, y)

= argmax $\int P(y_0, w | x_0, x_1, ..., x_N, y_1, ..., y_N) dw$ [Law of Total Probability]

But integrals are painful to work with, so let's not go there.

(6) Instead, we can do a point estimate approach. (a) compute the most probable value of û given the observations: $\hat{\omega} = \operatorname{argmax} P(\omega | x_0, x_1, ..., x_N, y_1, ..., y_N)$ = argmax $P(x_0, x_1, ..., x_N, y_1, ..., y_N | w) P(w)$ $P(x_0, x_1, ..., x_N, y_1, ..., y_N) \quad [Bayes Rule]$ = $argmax P(x_0, x_1, ..., x_N, y_1, ..., y_N | \omega) P(\omega)$ [remove constant factors] = argmax P(x0 | w)P(x, | x0, w) ... P(yn | x0, ..., xn, y1, ..., yn-1, w)P(w) [Chain Rule of Prob] = argmax $P(x_0) P(x_1) \cdots P(x_N) P(y_1 | w_1 x_1) \cdots P(y_N | w_1 x_N) P(w)$ [d-separation] = argmax $P(y, |w, x_1) \cdot \cdots \cdot P(y_N | w, x_N) P(w)$ = argmax P(w) TP (yn | w, xn) [remove constant factors] (b) compute the most probable value of \hat{y}_0 given $\hat{\omega}$: $\hat{y}_0 = \underset{y_0}{\operatorname{argmax}} P(y_0 | \hat{\omega}, x_0, x_1, ..., x_N, y_1, ..., y_N)$

= argmax P(yolw, xo) [d-separation]

REGLESSION PROBLEMS

- 17 In short:
 - (a) compute $\hat{w} = \operatorname{argmax} P(w) \prod_{n=1}^{N} P(y_n | w, x_n)$
 - (b) compute $\hat{y}_o = \operatorname{argmax} P(y_o | \hat{w}, x_o)$

This is called the MAP (maximum a posteriori) estimate.

(B) A special case of the MAP estimate assumes that P(w) is the same for every possible w. Since it then becomes a constant factor, we can drop how comit from the argmax:

(a) compute $\hat{w} = \operatorname{argmax} \prod_{n=1}^{N} P(y_n | w_n x_n)$

(b) compute j. = argmax P(y, 1û, x.)

This is called the MLE (maximum likelihood estimate).

how can this
be, if there's
an infinite space
of w-vectors?
don't warry
about it