Applied Machine Learning

Generalization

Reihaneh Rabbany



Learning objectives

- generalization
- bias and variance trade-off
- validation and cross-validation
- evaluation

Generalization

$$x$$
 input features $\xrightarrow{\text{ML algorithm} \atop \text{features}} \xrightarrow{\text{ML algorithm} \atop \text{with parameters } w} \xrightarrow{\text{ML algorithm} \atop \text{labels}} y$

training: parameter estimation

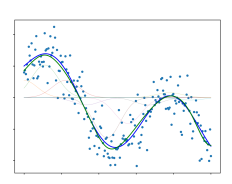
$$\mathcal{D} = \{(x^{(n)}, y^{(n)})\}_{n=1}^N$$

$$\mathcal{D} = \{(x^{(n)},y^{(n)})\}_{n=1}^N egin{array}{c} w^* = rg \min_w J(w) \ J(w) = rac{1}{N}\sum_{n=1}^N l(y^{(n)},f(x^{(n)};w)) \end{array}$$

Example: Linear regression

model:
$$\hat{y} = f_w(x) = w^ op x : \mathbb{R}^D o \mathbb{R}$$

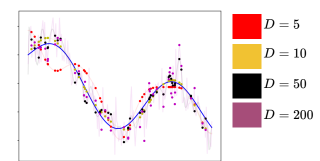
cost function:
$$J_w = rac{1}{N} \sum_n rac{1}{2} (y^{(n)} - \hat{y}^{(n)})^2$$



who the trained model generalizes to unseen data? how accurate is the model in general?

inference: $\hat{y} = f(x; w^*)$

on unseen data for which we haven't seen the label

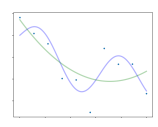


Generalization and model complexity

simple models cannot fit the data

bias

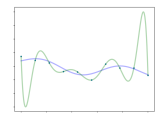
large training error due to underfitting



expressive models can overfit the data

- small training error
- large test error due to overfitting

variance



regularization can help us trade-off between bias and variance

we want to see how these two terms contribute to the generalization error

Generalization and model complexity

example

columns: a different type of model g(x)

rows: different datasets

datasets are from the same distribution

$$x^{(n)},y^{(n)}\sim p(x,y)$$

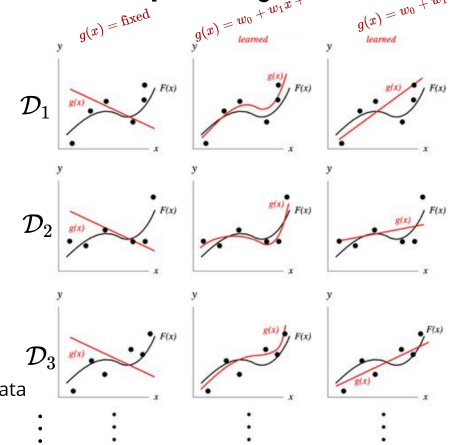
F(x) the best possible model

higher variance

the complex model varies more with the dataset it may not generalize well for this reason

higher bias

the simple model is biased to a particular type of data it underfits, but it has a low variance



Bias-variance decomposition: Setup

decompose the generalization error to see the effect of bias and variance (for L2 loss) assume a true distribution $\ p(x,y)$

best prediction given L2 loss $f(x) = \mathbb{E}_p[y|x]$

assume that a dataset $\mathcal{D} = \{(x^{(n)}, y^{(n)})\}_n$ is sampled from p(x,y)

let $\hat{f}_{\mathcal{D}}$ be our model based on the dataset

what we care about is the generalization error (aka expected loss, expected risk)

$$\mathbb{E}[(\hat{f}_{\mathcal{D}}(x)-y)^2]$$

all blue items are random variables

Bias-variance decomposition

what we care about is the generalization error

above simplifies to the following (the remaining terms are going to be zero)

$$=\mathbb{E}[(\hat{f}_{\mathcal{D}}(x)-\mathbb{E}_{\mathcal{D}}[\hat{f}_{\mathcal{D}}(x)])^2]+\mathbb{E}[(f(x)-\mathbb{E}_{\mathcal{D}}[\hat{f}_{\mathcal{D}}(x)])^2] \ +\mathbb{E}[\epsilon^2]$$
 variance bias^2 unavoidable noise error

Bias-variance decomposition

the expected loss is decomposed to:

$$\mathcal{E}[(\hat{f}_{\mathcal{D}}(x) - \mathbb{E}_{\mathcal{D}}[\hat{f}_{\mathcal{D}}(x)])^2] + \mathbb{E}[(f(x) - \mathbb{E}_{\mathcal{D}}[\hat{f}_{\mathcal{D}}(x)])^2] + \mathbb{E}[\epsilon^2]$$

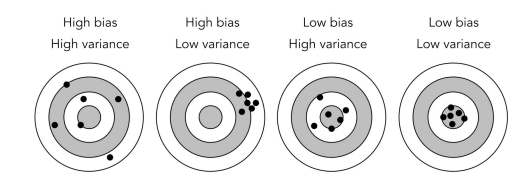
variance: how change of dataset affects the prediction

bias: how average over all datasets **noise error:** the error differs from the regression function even if we used the

true model f(x)

different models vary in their trade off between error due to bias and variance

- simple models: often more biased
- complex models: often have more variance

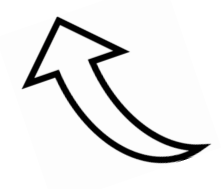


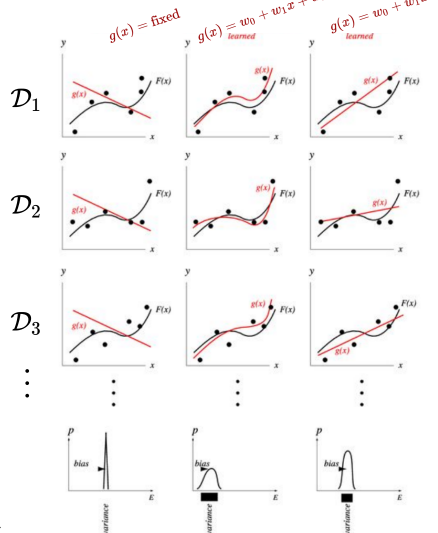
Bias vs. variance

example

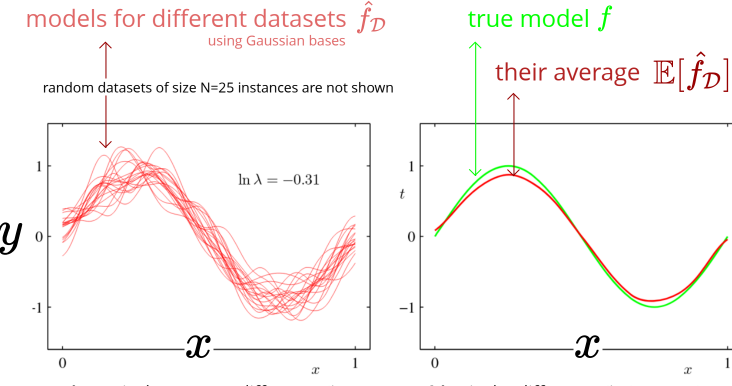
distribution of error (cost) due to randomness of dataset

we care about the expected error bias causes a high error for all choices of dataset higher variance also increases the expected error





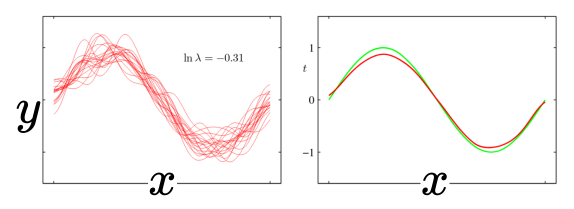
Example: bias vs. variance



variance is the average difference (in squared L2 norm) between these curves and their average

bias is the difference (in L2 norm) between two curves

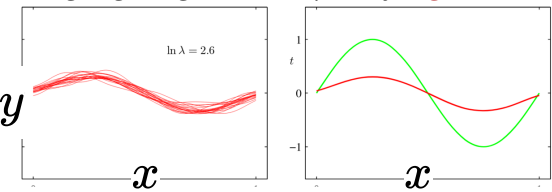
Example: bias vs. variance



side note

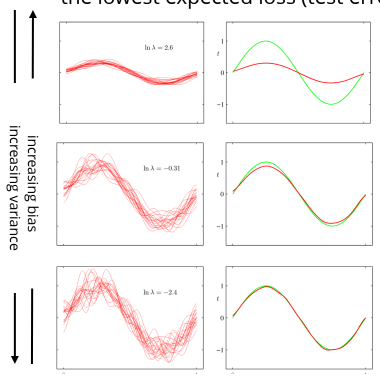
the average fit is very good, despite high variance **model averaging:** uses "average" prediction of expressive models to prevent overfitting

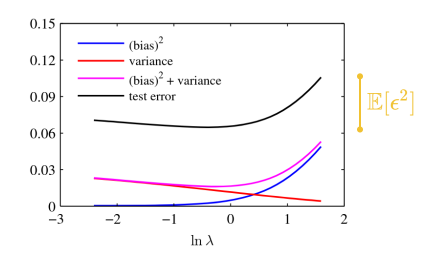
using larger regularization penalty: higher bias - lower variance



Example: bias vs. variance

the lowest expected loss (test error) is somewhere between the two extremes

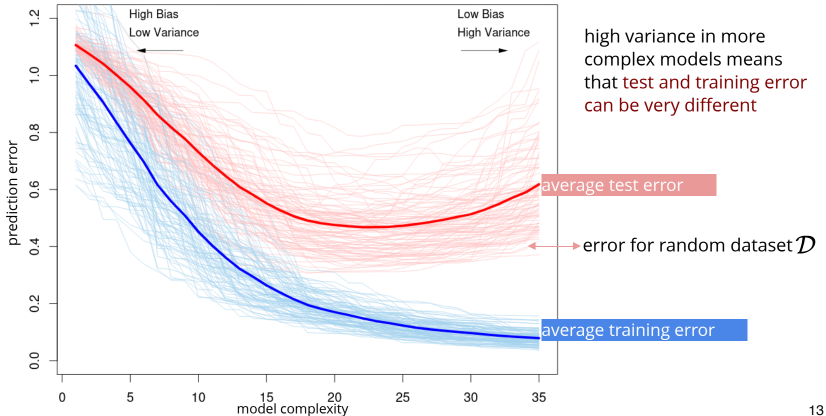




in practice, how to decide which model to use?

Effect on training and test error

high bias in simplistic models means that training error can be high



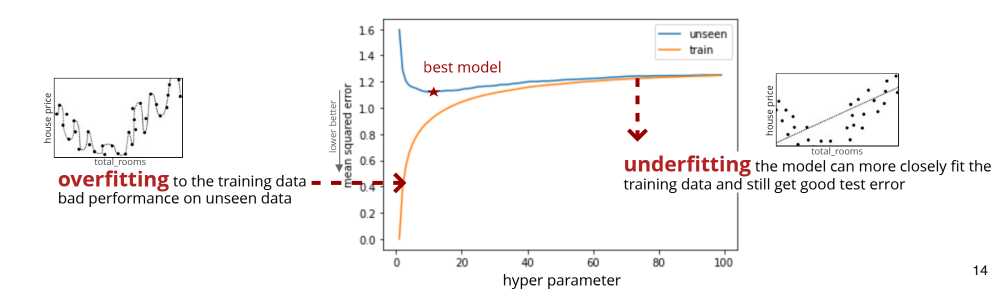
Model selection

many ML algorithms have hyper-parameters (e.g., the number of nonlinear basis to use)

how should we select the best hyper-parameter?

example

performance of a regression model on California Housing Dataset



Model selection

what if unseen data is completely different from training data? no point in learning!

assumption: training data points are samples from an unknown distribution independent identically distributed (**IID**)

$$x^{(n)},y^{(n)}\sim p(x,y)$$

unseen data comes from the same distribution.

		unseen
00000000000000000000000000000000000000	0 0 0 0 0 0 0 0 1 1 1 1 1 1 1 1 2 2 2 2	00000000000000000000000000000000000000
9999999999999999999999999999999999999		7

Loss, cost and generalization

assume we have a **model** $f: x \mapsto y$ for example $f: |\mathbf{3}| \mapsto 3$

and we have a **loss function** that measures the error in our prediction $~\ell:y,\hat{y} o\mathbb{R}$

for example
$$\ell(y,\hat{y})=(y-\hat{y})^2$$
 for regression $\ell(y,\hat{y})=\mathbb{I}(y
eq\hat{y})$ for classification

we train our models to minimize the cost function:

$$J = rac{1}{|\mathcal{D}_{ ext{train}}|} \sum_{x,y \in \mathcal{D}_{ ext{train}}} \ell(y,f(x))$$

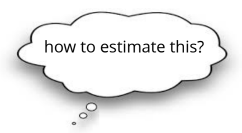
how to estimate this?

We can drop this, why?

what we really care about is the <code>generalization</code> error: $\mathbb{E}_{x,y\sim p}\;\ell(y,f(x))$

we can not measure this, why?

we can set aside part of the given data and use it to estimate generalization error

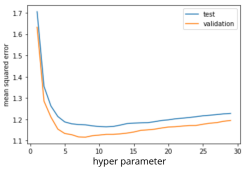


Validation set

what we really care about is the **generalization error**: $\mathbb{E}_{x,y\sim p}\;\ell(y,f(x))$ we can set aside part of the training data and use it to **estimate** the generalization error

pick a hyper-parameter that gives us the best **validation error** at the very end, we report the error on **test set**

validation and test error could be different because they use limited amount of data

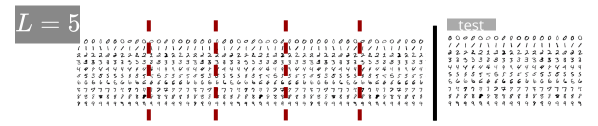


how to get a better estimate of generalization error? increase the size of the validation set? *this reduces the training set*

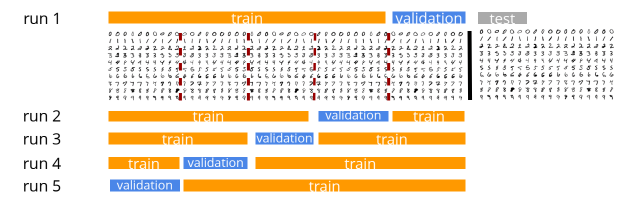


Cross-validation helps us in getting better estimates + uncertainty measure

- divide the (training + validation) data into L parts
- use one part for validation and L-1 for training



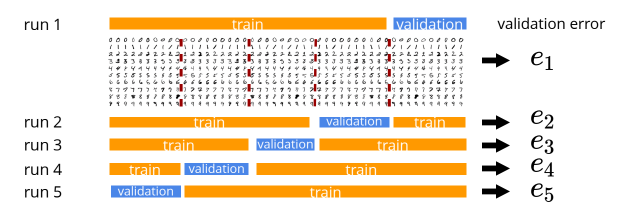
- divide the (training + validation) data into L parts
- use one part for validation and L-1 for training



- use the **average** validation error and its variance (uncertainty) to pick the best model
- report the test error for the final model

this is called **L-fold** cross-validation in **leave-one-out** cross-validation L=N (only one instance is used for validation)

- divide the (training + validation) data into L parts
- use one part for validation and L-1 for training



 use the average validation error and its variance (uncertainty) to pick the best model

$$ar{e}=rac{1}{5}\sum_{i=1}^{5}e_{i}$$

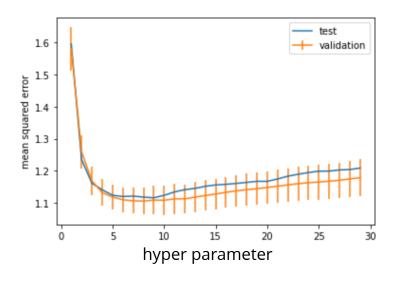
report the test error for the final model

tr.

this is called **L-fold** cross-validation

in **leave-one-out** cross-validation L=N (only one instance is used for validation)

example the plot of the mean and standard deviation in 10 fold cross-validation



test error is plotted only to show its agreement with the validation error; in practice we don't look at the test set for hyper-parameter tunning

a rule of thumb: pick the simplest model within one std of the model with lowest validation error

Performance metrics for classification

Not all errors are the same In particular in classification, we have different types of mistakes

false positive (type I) and false negative (type II)

example:

patient does not have disease but received positive diagnostic (Type I error) patient has disease but it was not detected (Type II error)

a message that is not spam is assigned to the spam folder (Type I error) a message that is spam appears in the regular folder (Type II error)

Performance metrics for classification

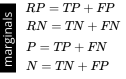
binary classification results:

FP false positive (type |)
FN false negative (type | |)

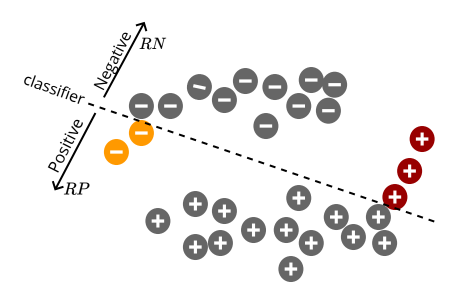
TP true positive

TN true negative

confusio	n matrix	Truth		\sum
	Result	TP	FP	RP
		FN	TN	RN
_	Σ	Р	N	



$$TN + TP + FN + FP = ?$$



example:	Truth		Σ
Result	14	2	16
	3	11	14
\sum	17	13	

Performance metrics for classification

confusi	on matrix	Truth		\sum
Result	TP	FP	RP	
	nesun	FN	TN	RN
	Σ	Р	N	

(example:	Truth		Σ
Result	14	2	16	
	3	11	14	
	Σ	17	13	

$$Precision = \frac{TP}{RP} = \frac{14}{16}$$
 $Recall = \frac{TP}{R} = \frac{14}{17}$

$$\frac{1}{F_{\beta}score} = \frac{1}{1+\beta^2} \frac{1}{Precision} + \frac{\beta^2}{1+\beta^2} \frac{1}{Recall}$$

$$egin{align*} Accuracy &= rac{TP+TN}{P+N} \ Precision &= rac{TP}{RP} \ Recall &= rac{TP}{P} \ F_1score &= 2rac{Precision imes Recall}{Precision+Recall} \ F_{eta}score &= (1+eta^2)rac{Precision imes Recall}{eta^2 Precision+Recall} \ \end{array}$$
 (Harmonic mean)

recall is β times more important compared to precision

$$egin{aligned} Miss\ rate &= rac{FN}{P} \ Fallout &= rac{FP}{N} \end{aligned}$$

false positive rate

$$False\ discovery\ rate = rac{FP}{RP}$$

$$Selectivity = rac{TN}{N}$$

specificity

$$Fallout = rac{FP}{N}$$
 $False\ discovery\ rate = rac{FP}{RP}$
 $Selectivity = rac{TN}{N}$
 $False\ omission\ rate = rac{FN}{RN}$

$$False\ omission\ rate = rac{FN}{RN} \ Negative\ predictive\ value = rac{TN}{RN}$$

Trade-off between precision and recall

How many false positives do we tolerate? How important are false negatives?

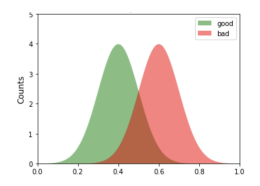
e.g. spam in inbox v.s. negative test for cancer test

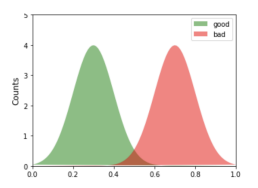
We can often control the trade-off between type I & type II error

e.g. by changing the threshold of p(y = 1|x) if we produce class score (probability)

Most ML algorithm produces class score or probability p(y=1|x) $1 \qquad 0$ no false positive also no true positive also no true negative also no true negative

goal: evaluate class scores/probabilities (independent of choice of threshold)





Trade-off between precision and recall

How many false positives do we tolerate?

How important are false negatives?

e.g. spam in inbox v.s. negative test for cancer test

We can often control the trade-off between type I & type II error

e.g. by changing the threshold of p(y = 1|x) if we produce class score (probability)

goal: evaluate class scores/probabilities (independent of choice of threshold)

Receiver Operating Characteristic ROC curve, a function of threshold t

TPR(t) = TP(t)/P (**recall**, sensitivity at t, hit rate) **FPR(t)** = FP(t)/N (**fallout**, false alarm at t, type I error rate)

Area Under the Curve (**AUC**) is used as a threshold independent measure of quality of the classifier

$$AUC = \sum_t TPR(t)(FPR(t) - FPR(t-1))$$
 , box-rule approximation

Most ML algorithm produces class score or probability threshold p(y = 1|x)no false negative no false positive also no true positive also no true negative ROC CURVE PERFECT CLASSIFIER

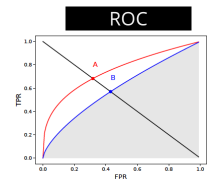
FALSE POSITIVE RATE

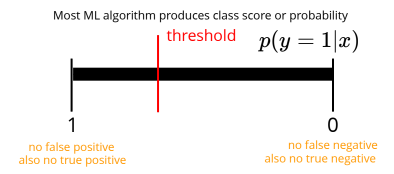
Precision-recall curves

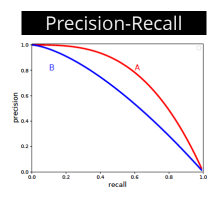
Similar to ROC curve but more helpful in some situation

- when size of negative set is also a model parameter, e.g. in information retrieval
- when there is class imbalance, e.g. in fraud detection when $N\gg P$), since ROC curves are insensitive to class imbalance

Instead we curve Precision vs Recall for different thresholds

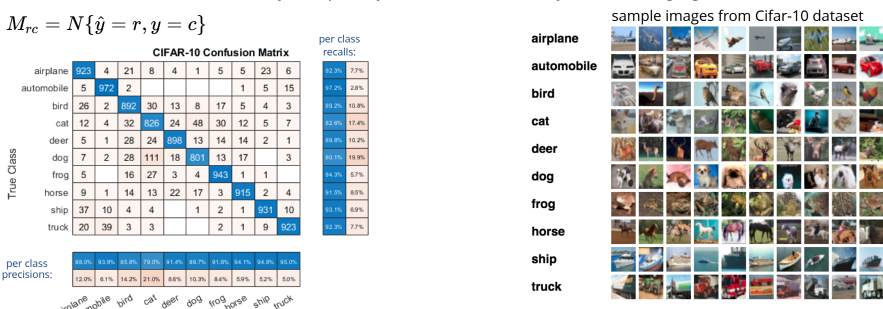






Confusion Matrix for multiclass classification

A CxC table that shows how many samples of each class are classified as belonging to another class



classifier's accuracy is the sum of diagonal divided by the sum-total of the matrix, you can also report the average of the F_1 scores per class (macro), or weight the average by class sizes

when evaluating a classifier it is useful to look at the confusion matrix

Bias and Fairness Challenge

The model learns from the distribution of the input data {train, validation, test are still sampled based on some process}

the demographic and phenotypic composition of training and benchmark datasets are important

Growing use, growing concerns

- Amazon's hiring algorithm decides not to invite women to interview, read it here
- Google's online ad algorithm decides to show high-income jobs to men much more often than to women, read about it here
- A machine learning algorithm denies you credit based on race or gender, read it here
- Health care algorithm offers less care to black patients, read it here, and here
- Florida risk score algorithm used in courts assign higher risk to black defendants, read it here

How can we factor these in the evaluation of models?

Many recent works, for example see this book on fairness & ML, here, read this article on bias detectives, or this course on data ethics

Face-recognition software is perfect – if you're a white man





Inductive bias

learning algorithms make implicit assumptions learning or inductive bias e.g., we are often biased towards **simplest explanations** of our data

Occam's razor between two models (explanations) we should prefer the simpler one

example

both of the following models perfectly fit the data

$$egin{array}{c|cccc} rac{ extbf{x1} & extbf{x2} & extbf{x3} & extbf{f}}{1 & 0 & 1 & 0} & \hat{f}(x) = x_2 & ext{this one is simpler} \ rac{-1}{1} & -1 & 0 & 1 \ 1 & 1 & 1 & 1 & 1 \end{array}$$





OCCAM'S RAZOR



why does is make sense for learning algorithms to be biased?

- the world is not random
- there are regularities, and induction is possible (why do you think the sun will rise in the east tomorrow morning?

what are some of the inductive biases in using linear regression?

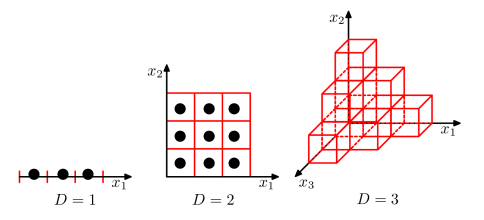


Curse of dimensionality

learning in **high dimensions** can be difficult since the volume of space grows exponentially fast with the dimension

example:

suppose our data is uniformly distributed in some range, say $x \in [0,3]^D$ predict the label by counting labels in the same unit of the grid to have at least one example per unit, we need 3^D training examples for D=180 we need more training examples than the number of particles in the universe

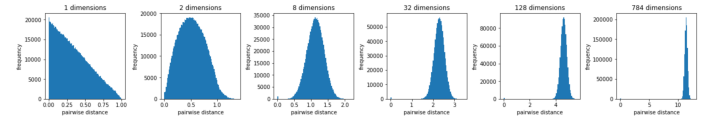




Curse of dimensionality

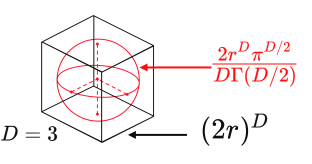
in high dimensions most points have similar distances!

histogram of pairwise distance of 1000 points with random features of D dimensions



as we increase dimension, distances become "similar"!

- Q. why are most distances similar?
- **A.** in high dimensions most of the volume is close to the corners!



$$\lim_{D o\infty}rac{\mathrm{volum}(\circ)}{\mathrm{volum}(\square)}=0$$

a "conceptual" visualization of the same idea # corners and the mass in the corners grow quickly with D

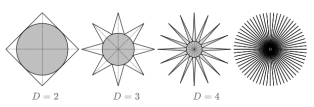
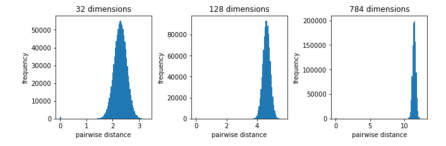


image: Zaki's book on Data Mining and Analysis

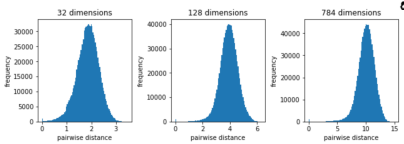
Real-word vs. randomly generated data

how come ML methods work for image data (D=number of pixels)? pairwise distance for random data



in fact KNN works well for image classification

pairwise distance for D pixels of MNIST digits



112345678

the statistics do not match that of random high-dimensional data!

Test Error Ra	te (%)
Linear classifier (1-layer NN)	12.0
K-nearest-neighbors, Euclidean	5.0
K-nearest-neighbors, Euclidean, deskewed	2.4
K-NN, Tangent Distance, 16x16	1.1
K-NN, shape context matching	0.67
1000 RBF + linear classifier	3.6
SVM deg 4 polynomial	1.1
2-layer NN, 300 hidden units	4.7
2-layer NN, 300 HU, [deskewing]	1.6
LeNet-5, [distortions]	0.8
Boosted LeNet-4, [distortions]	0.7

Manifold hypothesis

real-world data is often far from uniformly random

manifold hypothesis: real data lies close to the surface of a manifold

example

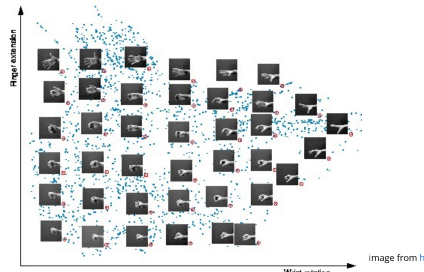
data dimension: D=3

manifold dimension: $\hat{D}=2$

example

data dimension: D = number of pixels (64x64)

manifold dimension: $\hat{D}=2$



No free lunch

consider the binary classification task:



read more here

there are $2^4 = 16$ binary functions that perfectly fit our dataset

our **learning algorithm** can produce one of these as our classifier $\hat{f}: \{0,1\}^3 \to \{0,1\}$

the same algorithm cannot perform well for all possible class of problems (f)

no free lunch

each ML algorithm is biased to perform well on some class of problems

there is no single algorithm that performs well on all class of problems

Summary

- complex models can have very different training and test error (generalization gap)
- regularization bounds this gap by penalizing model complexity
- bias-variance trade off:
 - formalizes the relation between
 - training error (bias)
 - o complexity (variance) and
 - and the test error (bias + variance)
 - not so elegant beyond L2 loss
- what we care about is the generalization of ML algorithms
 - overfitting: good performance on the training set doesn't mean the same for the test set
 - underfitting: we don't even have a good performance on the training set
- estimated using a validation set or better, we could use cross-validation