



**Heiko Paulheim** 

# Why Hyperparameter Tuning?

- What we have seen so far
  - many learning algorithms for classification, regression, ...
- Many of those have hyperparameters
  - k and distance function for k nearest neighbors
  - splitting and pruning options in decision tree learning
  - hidden layers in neural networks
  - C, gamma, and kernel function for SVMs
  - **–** ...
- But what is their effect?
  - hard to tell in general
  - rules of thumb are rare

#### Parameters vs. Hyperparameters

- Parameters
  - ...are learned during training
  - Typical examples:
    - Coefficients in (linear) regression
    - Weights in neural networks
    - •
  - Training:
    - Find set of parameters so that objective function is minimized/maximized
      - (on a holdout set)



#### Parameters vs. Hyperparameters

- Hyperparameters
  - ...are fixed before training
  - Typical examples:
    - Network layout and learning rate in neural networks
    - k in kNN
    - ...
  - Training:
    - Find set of parameters so that objective function is minimized/maximized
      - (on a holdout set)
      - given a previously fixed set of hyperparameters

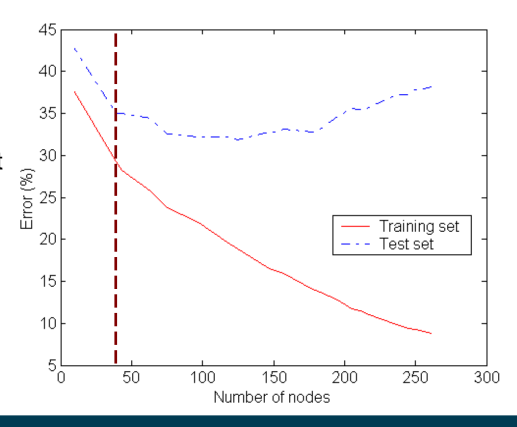


# Hyperparameter Tuning – a Naive Approach

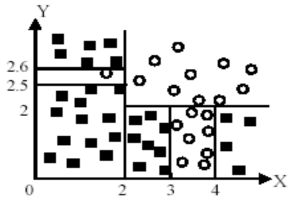
- You probably know that approach from the exercises
  - 1. run classification/regression algorithm
  - 2. look at the results (e.g., accuracy, RMSE, ...)
  - 3. choose different parameter settings, go to 1
- Questions:
  - when to stop?
  - how to select the next parameter setting to test?

#### Recap overfitting:

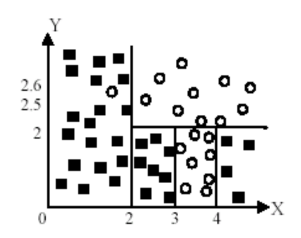
- classifiers may overadapt to training data
- the same holds for parameter settings
- Possible danger:
  - finding parameters that work well on the training set
  - but not on the test set
- Remedy:
  - train / test / validation split

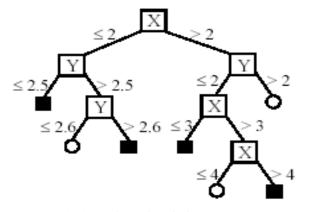


Parameter option: pruning (yes/no)

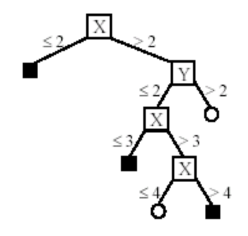


(A) A partition of the data space



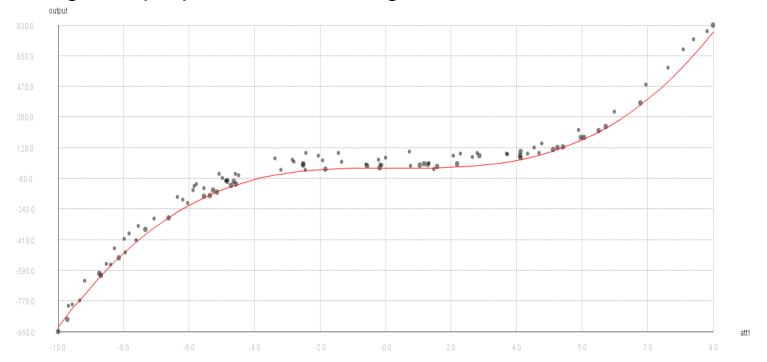


(B). The decision tree

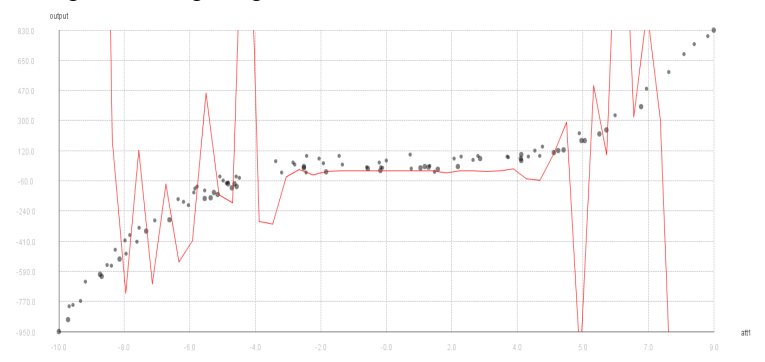


4/22/24

- Real example: train a local polynomial regression model
  - Parameter to tune: find the optimal maximum degree of the polynomial
- Tuning with proper validation: degree = 3



- Real example: train a local polynomial regression model
  - Parameter to tune: find the optimal maximum degree of the polynomial
- Tuning overfitting: degree = 9



# Hyperparameter Tuning: Brute Force

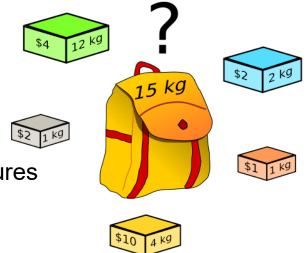
- Try all parameter combinations that exist
- Consider, e.g., a k-NN classifier
  - try 30 different distance measures
  - try all k from 1 to 1,000
  - use weighting or not
    - $\rightarrow$  60,000 runs of k-NN

→ we need a better strategy than brute force!

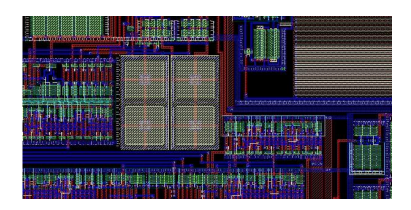
- Hyperparameter tuning is an optimization problem
- Finding optimal values for N variables
- Properties of the problem:
  - the underlying model is unknown
    - i.e., we do not know changing a variable will influence the results
  - we can tell how good a solution is when we see it
    - i.e., by running a classifier with the given parameter set
  - but looking at each solution is costly
    - e.g., 60,000 runs of k-NN
- Such problems occur quite frequently

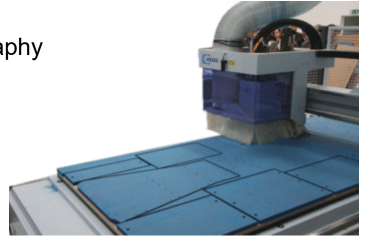
- Related problem:
  - feature subset selection
  - cf. Data Mining 2, first lecture
- Given n features, brute force requires 2<sup>n</sup> evaluations
  - for 20 features, that is already one million
    - → ten million with cross validation

- Knapsack problem
  - given a maximum weight you can carry
  - and a set of items with different weight and monetary value
  - pack those items that maximize the monetary value
- Problem is NP hard
  - i.e., deterministic algorithms
     require an exponential amount of time
  - Note: feature subset selection for N features requires 2 evaluations

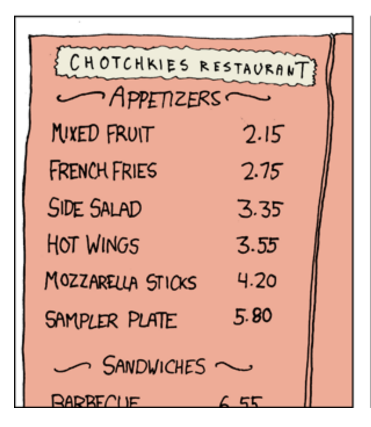


- Many optimization problems are NP hard
  - Routing problems (Traveling Salesman Problem)
  - Integer factorization
     hard enough to be used for cryptography
  - Resource use optimization
    - e.g., minimizing cutoff waste
  - Chip design
    - minimizing chip sizes





MY HOBBY: EMBEDDING NP-COMPLETE PROBLEMS IN RESTAURANT ORDERS





http://xkcd.com/287/

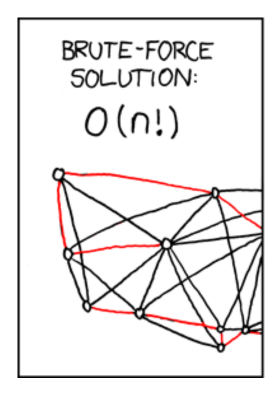
#### Hyperparameter Tuning: Brute Force

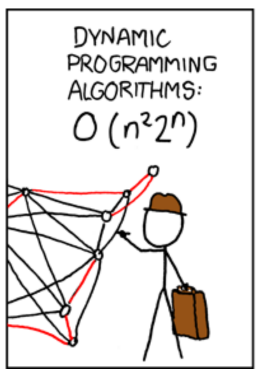
- Properties of Brute Force search
  - guaranteed to find the best parameter setting
  - too slow in most practical cases
- Grid Search
  - performs a brute force search
  - with equal-width steps on non-discrete numerical attributes (e.g., 10,20,30,..,100)
- Hyperparameter with a wide range (e.g., 0.0001 to 1,000,000)
  - with ten equal-width steps, the first step would be 1,000
  - but what if the optimum is around 0.1?
  - logarithmic steps may perform better for some parameters

# **Hyperparameter Tuning: Heuristics**

- Properties of Brute Force search
  - guaranteed to find the best parameter setting
  - too slow in most practical cases
- Needed:
  - solutions that take less time/computation
  - and often find the best parameter setting
  - or find a *near-optimal* parameter setting

#### **Beyond Brute Force**







https://xkcd.com/399/

# Hyperparameter Tuning: One After Another

- Given n parameters with m degrees of freedom
  - brute force takes m<sup>n</sup> runs of the base classifier
- Simple tweak:
  - 1. start with default settings
  - 2. try all options for the first parameter2a. fix best setting for first parameter
  - 3. try all options for the second parameter3a. fix best setting for second parameter
  - 4. ...
- This reduces the runtime to n\*m
  - i.e., no longer exponential!
  - but we may miss the best solution

# **Hyperparameter Tuning: Interaction Effects**

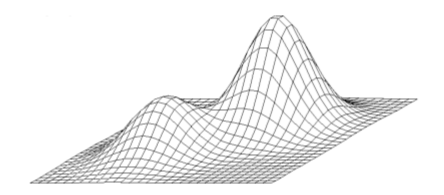
- Interaction effects make parameter tuning hard
  - i.e., changing one parameter may change the optimal settings for another one
- Example: two parameters p and q, each with values 0,1, and 2
  - the table depicts classification accuracy

	p=0	p=1	p=2
q=0	0.5	0.4	0.1
q=1	0.4	0.3	0.2
q=2	0.1	0.2	0.7

# **Hyperparameter Tuning: Interaction Effects**

- If we try to optimize one parameter by another (first p, then q)
  - we end at p=0,q=0 in six out of nine cases
  - on average, we investigate 2.3 solutions

	p=0	p=1	p=2
q=0	0,5	0.4	0.1
q=1	0.4	0.3	0.2
q=2	0.1	0.2	0.7



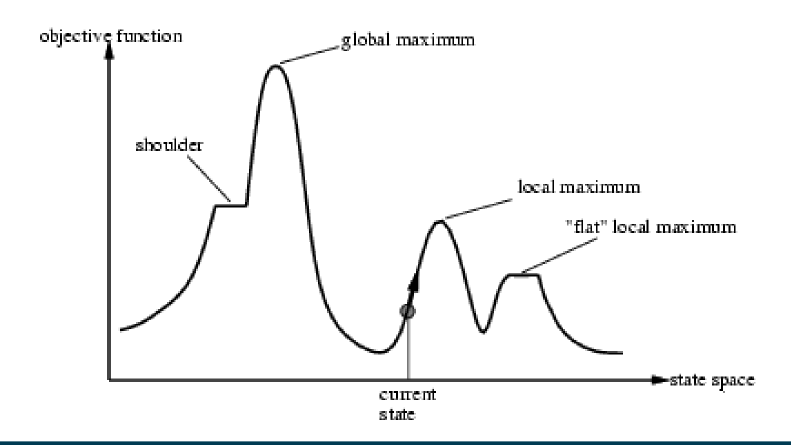
#### Hill-Climbing Search

- a.k.a. greedy local search
- always search in the direction of the steepest ascend
  - "Like climbing Everest in thick fog with amnesia"

```
function Hill-Climbing (problem) returns a state that is a local maximum inputs: problem, a problem local variables: current, a node neighbor, \text{ a node} current \leftarrow \text{Make-Node}(\text{Initial-State}[problem]) loop do neighbor \leftarrow \text{a highest-valued successor of } current if \text{Value}[\text{neighbor}] \leq \text{Value}[\text{current}] then \text{return State}[current] current \leftarrow neighbor
```

# **Hill-Climbing Search**

 Problem: depending on initial state, one can get stuck in local maxima



# Hill Climbing Search

- Given our previous problem
  - we end up at the optimum in three out of nine cases
  - but the local optimum (p=0,q=0) is reached in six out of nine cases!
  - on average, we investigate 2.1 solutions

	p=0	p=1	p=2
q=0	0,5	0.4	0.1
q=1	04	0.3	0,2
q=2	0.1	0.2	0.7

#### Variations of Hill Climbing Search

- Stochastic hill climbing
  - random selection among the uphill moves
  - the selection probability can vary with the steepness of the uphill move
- First-choice hill climbing
  - generating successors randomly until a better one is found, then pick that one
- Random-restart hill climbing
  - run hill climbing with different seeds
  - tries to avoid getting stuck in local maxima

#### **Local Beam Search**

- Keep track of k states rather than just one
- Start with k randomly generated states
- At each iteration, all the successors of all k states are generated
- Select the k best successors from the complete list and repeat

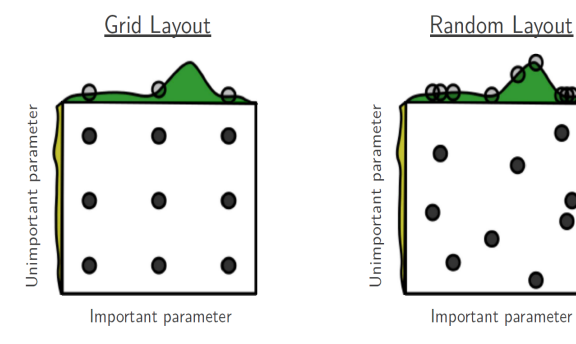


#### Grid Search vs. Random Search

- All the examples discussed so far use fixed grids
  - e.g., an interval from 0 to 1 with a step size of 0.05
- Challenges:
  - some hyperparameters are pretty sensitive
    - e.g., 0.02 is a good value, but 0 and 0.05 are not
  - others have little influence
    - but it is hard to know upfront which

#### Grid Search vs. Random Search

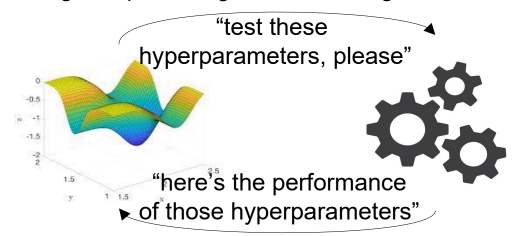
- Paper from 2012 (Bergstra and Bengio):
  - grid search may easily miss best parameters
  - random search often yields better results



Bergstra & Bengio: Random Search for Hyper-Parameter Optimization, JMLR, 2012

#### **Learning Hyperparameters**

- Hyperparameter tuning as a learning problem:
  - Given a set of hyperparameters H, predict performance p of model
  - The prediction model is referred to as a surrogate model or oracle
  - Rationale:
    - Training and evaluating an actual model is costly
    - Learning and predicting with the surrogate model is fast

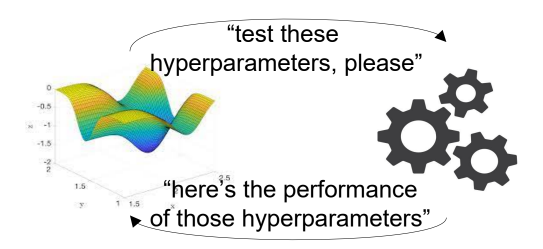


Surrogate Model

**Actual Model** 

#### **Learning Hyperparameters**

- Note:
  - we want to use not too many runs of the actual model
  - i.e., the surrogate model will have few training points
    - use a simple model
  - Most well-known: bayesian optimization



Surrogate Model

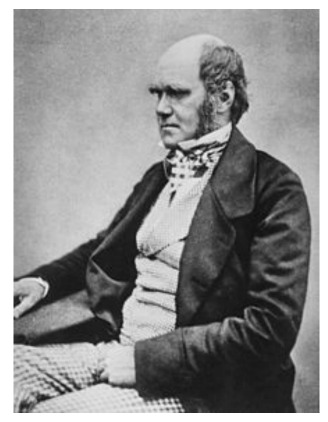
**Actual Model** 

#### Summary: Grid Search, Random Search, etc.

- Problems of grid search
  - Inefficient
  - Fixed grid sizes may miss good parameters
    - Smaller grid sizes would be even less efficient!
- Random search
  - Often finds good solutions in less time
- Learning hyperparameters / bayesian optimization
  - Sucessively tests hyperparameters close to local optima
  - Similar to hill climbing
    - Difference: explicit surrogate model

#### **Genetic Algorithms**

- Inspired by evolution
- Overall idea:
  - use a population of individuals (solutions)
  - create new individuals by crossover
  - introduce random mutations
  - from each generation, keep only the best solutions ("survival of the fittest")
- Developed in the 1970s
- John H. Holland:
  - Standard Genetic Algorithm (SGA)



Charles Darwin (1809-1882)

#### **Genetic Algorithms**

- Basic ingredients:
  - individuals: the solutions
    - hyperparameter tuning: a hyperparameter setting
  - a fitness function
    - hyperparameter tuning: performance of a hyperparameter setting (i.e., run learner with those parameters)
  - a crossover method
    - hyperparameter tuning: create a new setting from two others
  - a mutation method
    - hyperparameter tuning: change one parameter
  - survivor selection

# **SGA Reproduction Cycle**

- Select parents for the mating pool
   (size of mating pool = population size)
- 2. Shuffle the mating pool
- 3. For each consecutive pair apply crossover with probability p<sub>c</sub>, otherwise copy parents
- 4. For each offspring apply mutation (bit-flip with probability p<sub>m</sub> independently for each bit)
- 5. Replace the whole population with the resulting offspring

# **SGA Operators: 1-point crossover**

- Choose a random point on the two parents
- Split parents at this crossover point
- Create children by exchanging tails
- P<sub>c</sub> typically in range (0.6, 0.9)

Hyperparameter/ Solution	hp1	hp2	hp3	hp4	hp5	hp6	hp7
s1	true	0.87	0.75	0.01	sgd	0.05	0.72
s2	false	0.75	0.83	0.04	adam	0.04	0.53
Hyperparameter/ Solution	hp1	hp2	hp3	hp4	hp5	hp6	hp7
s1'	true	0.87	0.75	0.01	adam	0.04	0.53
s2'	false	0.75	0.83	0.04	sgd	0.05	0.72

#### **SGA Operators: Mutation**

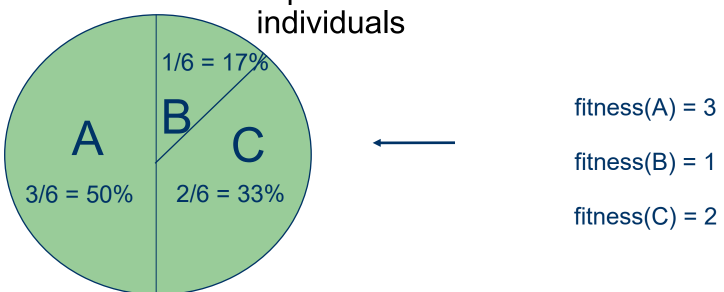
- Alter each gene independently with a probability  $p_m$ 
  - Sample from entire value range!
- $p_m$  is called the mutation rate
  - Typically between 1/pop\_size and 1/ chromosome\_length

Typically between 1/pop_cize and 1/ circumocome_length							
Hyperparameter/ Solution	hp1	hp2	hp3	hp4	hp5	hp6	hp7
s1	true	0.87	0.75	0.01	sgd	0.05	0.72
s2	false	0.75	0.83	0.04	adam	0.04	0.53
Hyperparameter/	hp1	hp2	bp3	hp4	hp5	hp6	hp7

Hyperparameter/ Solution	hp1	hp2	hp3	hp4	hp5	hp6	hp7
s1	true	0.87	0.75	0.01	adagrad	0.05	0.72
s2	false	0.75	0.86	0.04	adam	0.04	0.53

#### SGA Operators: Selection

- Main idea: better individuals get higher chance
  - Chances proportional to fitness
  - Implementation: roulette wheel technique
    - » Assign to each individual a part of the roulette wheel
    - » Spin the wheel n times to select n



#### **Crossover OR Mutation?**

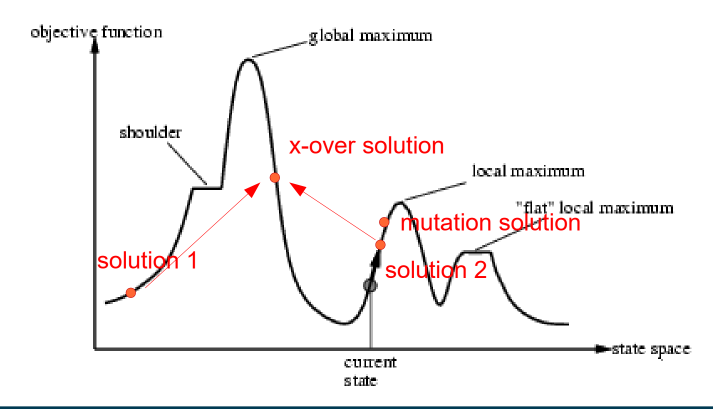
- Decade long debate: which one is better / necessary ...
- Answer (at least, rather wide agreement):
  - it depends on the problem, but
  - in general, it is good to have both
  - both have another role
  - mutation-only-EA is possible, crossover-only-EA would not work

## **Crossover OR Mutation? (cont'd)**

- Exploration: Discovering promising areas in the search space, i.e. gaining information on the problem
- Exploitation: Optimising within a promising area, i.e. using information
- There is co-operation AND competition between them
  - Crossover is explorative, it makes a big jump to an area somewhere "in between" two (parent) areas
  - Mutation is exploitative, it creates random small diversions, thereby staying near (in the area of) the parent

## **Crossover OR Mutation? (cont'd)**

- Recall the solution space example from Hill Climbing
  - crossover makes big jumps
  - mutation makes small steps



## **Crossover OR Mutation? (cont'd)**

- Only crossover can combine information from two parents
- Only mutation can introduce new information (alleles)
  - Remember: sample from entire value range
- To hit the optimum you often need a 'lucky' mutation

#### **Genetic Feature Subset Selection**

- Feature Subset Selection
  - can also be solved by Genetic Programming
- Individuals: feature subsets
- Representation: binary
  - 1 = feature is included
  - 0 = feature is not included
- Fitness: classification performance
- Crossover: combine selections of two subsets
- Mutation: flip bits

#### **Selecting a Learner**

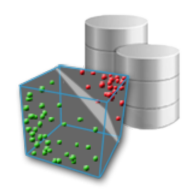
- So far, we have looked at finding good parameters for a learner
  - the learner was always fixed
- A similar problem is selecting a learner for the task at hand
- Again, we could go with search
- Another approach is meta learning

#### Selecting a Learner by Meta Learning

- Meta Learning
  - i.e., learning about learning
- Goal: learn how well a learner will perform on a given dataset
  - features: dataset characteristics, learning algorithm
  - prediction target: accuracy, RMSE, ...

#### Selecting a Learner by Meta Learning

- Also known as AutoML
- Basic idea: train a regression model
  - data points: individual datasets plus ML approach
  - target: expected accuracy/RMSE etc.



- Examples for features
  - number of instances/attributes
  - fraction of nominal/numerical attributes
  - min/max/average entropy of attributes
  - skewness of classes

**–** ...

## Selecting a Learner by Meta Learning

auto-sklearn 0.12.6 Start Releases Installation Manual Examples API Extending Search

auto-sklearn Example

Manual

License

Citing auto-sklearn

Contributing

#### auto-sklearn

auto-sklearn is an automated machine learning toolkit and a drop-in replacement for a scikit-learn estimator:

```
>>> import autosklearn.classification
>>> cls = autosklearn.classification.AutoSklearnClassifier()
>>> cls.fit(X_train, y_train)
>>> predictions = cls.predict(X_test)
```

auto-sklearn frees a machine learning user from algorithm selection and hyperparameter tuning. It leverages recent advantages in *Bayesian optimization*, meta-learning and ensemble construction. Learn more about the technology behind auto-sklearn by reading our paper published at NIPS 2015.

#### NEW: Auto-sklearn 2.0

Auto-sklearn 2.0 includes latest research on automatically configuring the AutoML system itself and contains a multitude of improvements which speed up the fitting the AutoML system.

auto-sklearn 2.0 works the same way as regular auto-sklearn and you can use it via

```
>>> from autosklearn.experimental.askl2 import AutoSklearn2Classifier
```

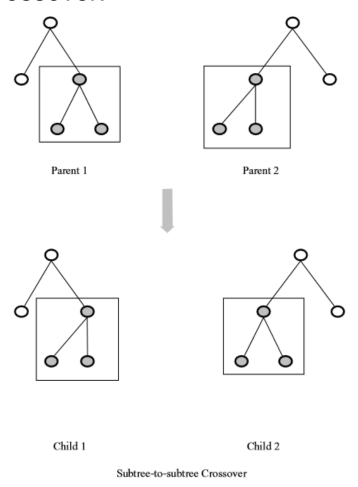
A paper describing our advances is available on arXiv.

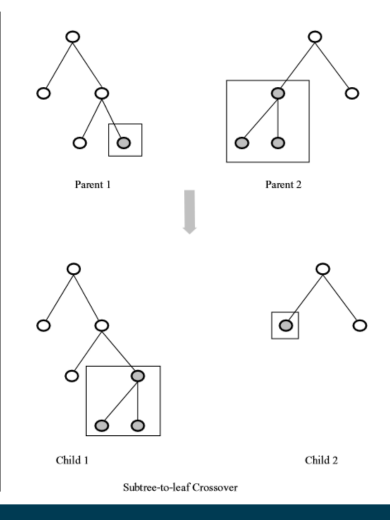
#### ...and now for something completely different.

- Recap: search heuristics are good for problems where...
  - finding an optimal solution is difficult
  - evaluating a solution candidate is easy
  - the search space of possible solutions is large
- Possible solution: genetic programming
- We have encountered such problems quite frequently
- Example: learning an optimal decision tree from data

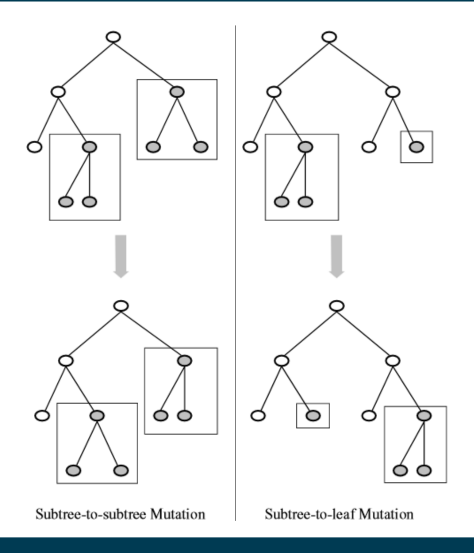
- e.g., GAIT (Fu et al., 2003)
  - also the source of the pictures on the following slides
- Population: candidate decision trees
  - initialization: e.g., trained on small subsets of data
- Create new decision trees by means of
  - crossover
  - mutation
- Fitness function: e.g., accuracy

Crossover:

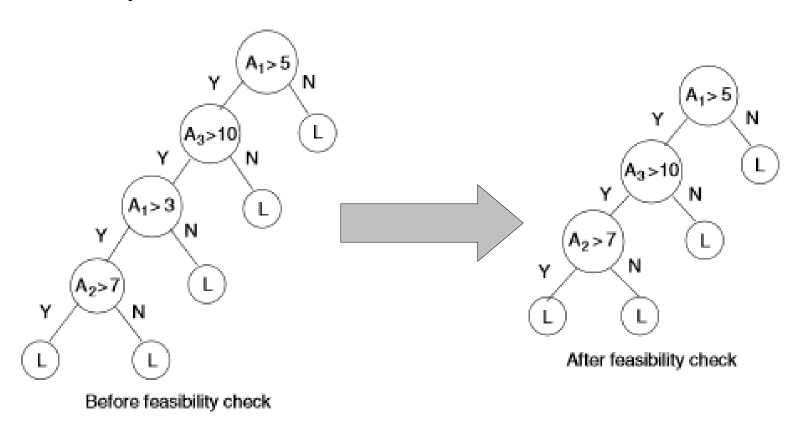




Mutation:



Feasibility Check:

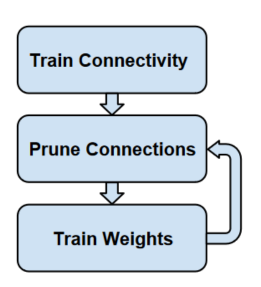


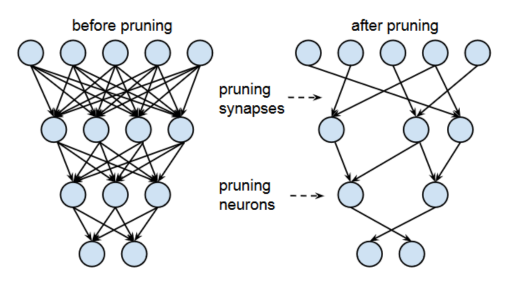
## **Combination of GP with other Learning Methods**

- Rule Learning ("Learning Classifier Systems"), since late 70s
  - Population: set of rule sets (!)
  - Crossover: combining rules from two sets
  - Mutation: changing a rule
- Artificial Neural Networks
  - Easiest solution: fixed network layout
  - The network is then represented as an ordered set (vector) of weights e.g., [0.8, 0.2, 0.5, 0.1, 0.1, 0.2]
  - Crossover and mutation are straight forward
  - Variant: AutoMLP
    - Searches for best combination of hidden layers and learning rate

# Hyperparameter Optimization vs. Pruning

- Architecture of a neural network can be seen as parameters
  - How many hidden layers? Which size?
- Pruning approaches: train large network, then start eliminating connections





Han et al. (2015): Learning both Weights and Connections for Efficient Neural Network

#### Wrap-Up

- Hyperparameter tuning is important
  - many learning methods work poorly with standard hyperparameters
  - often no global optimum, dataset dependent
- Hyperparameter tuning has a large search space
  - trying all combinations is infeasible
  - interaction effects do not allow for one-by-one tuning
- State of the art
  - Grid search, random search, bayesian optimization

#### Wrap-Up

- Heuristic Methods
  - Hill climbing with variations
  - Beam search
  - Simulated Annealing
  - Genetic Programming
  - Random search
  - Hyperparameter learning
- Other uses of genetic programming
  - Feature subset selection
  - Model fitting

#### Hyperparameter Tuning: Criticism

Just let those numbers sink...

— ...think: carbon footprint

– ...think: fair chances?

Consumption	CO <sub>2</sub> e (lbs)
Air travel, 1 passenger, NY↔SF	1984
Human life, avg, 1 year	11,023
American life, avg, 1 year	36,156
Car, avg incl. fuel, 1 lifetime	126,000
Training one model (GPU)	
Training one model (GPU)  NLP pipeline (parsing, SRL)	39
	39 78,468
NLP pipeline (parsing, SRL)	

Table 1: Estimated CO <sub>2</sub> emissions from training com-
mon NLP models, compared to familiar consumption. <sup>1</sup>

		Estimated cost (USD)	
Models	Hours	Cloud compute	Electricity
1	120	\$52-\$175	\$5
24	2880	\$1238-\$4205	\$118
4789	239,942	\$103k-\$350k	\$9870

Table 4: Estimated cost in terms of cloud compute and electricity for training: (1) a single model (2) a single tune and (3) all models trained during R&D.

Strubell et al. (2019): Energy and Policy Considerations for Deep Learning in NLP

### **Questions?**







**Heiko Paulheim**