

5. Simulated Annealing

5.1 Basic Concepts



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Outline

- **Introduction**
- **Real Annealing and Simulated Annealing**
- **Metropolis Algorithm**
- **Template of SA**
- **A Simple Example**
- **References**



Introduction

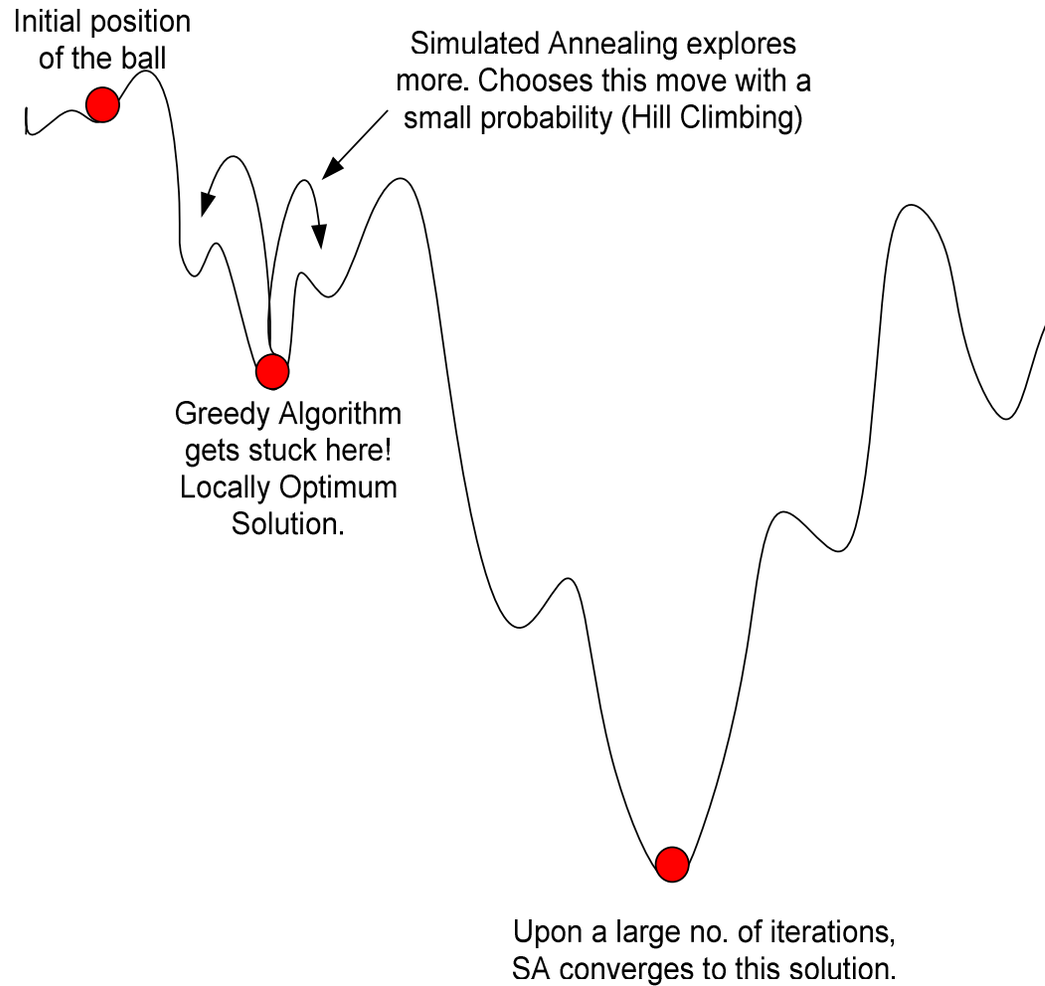


What Is Simulated Annealing?

- **Simulated Annealing (SA)**
 - SA is applied to solve optimization problems
 - SA is a **stochastic algorithm**
 - SA is escaping from local optima by allowing worsening moves
 - SA is a **memoryless algorithm**, the algorithm does not use any information gathered during the search
 - SA is applied for both **combinatorial** and **continuous** optimization problems
 - SA is simple and easy to implement.
 - SA is motivated by the physical annealing process

Simulated Annealing: Part 1

SA vs Greedy Algorithms: Ball on terrain example



History

- **Numerical simulation of annealing**, Metropolis et al. 1953.

N. Metropolis, A. Rosenbluth, M. Rosenbluth, A. Teller, and E. Teller. Equation of state calculations by fast computing machines. *Journal of Chemical Physics*, 21:1087–1092, 1953.

History

- SA for combinatorial problems
 - Kirkpatrick et. al, 1986
 - Cerny, 1985

S. Kirkpatrick, C. D. Gelatt, and M. P. Vecchi. Optimization by simulated annealing. *Science*, 220(4598):671–680, 1983.

V. Cerny, Thermodynamical approach to the traveling salesman problem : an efficient simulation algorithm. *J. of Optimization Theory and Applications*, 45(1):41–51, 1985.

History

- Originally, the use of simulated annealing in combinatorial optimization
- In the 1980s, SA had a major impact on the field of heuristic search for its simplicity and efficiency in solving **combinatorial optimization problems**.
- Then, it has been extended to deal with **continuous optimization problems**
- SA was inspired by an analogy between the **physical annealing process of solids** and the problem of **solving large combinatorial optimization problems**.

Applications

- **Basic problems**
 - Traveling Salesman Problem
 - Graph partitioning
 - Matching prob.
 - Quadratic Assignment
 - Linear Arrangement
 - Scheduling
 -

Applications

- **Engineering problem**
 - VLSI: Placement, routing...
 - Facilities layout
 - Image processing
 - Code design
 - Biology
 - Physics
 -

Real Annealing and Simulated Annealing



Real Annealing Technique

- **Annealing Technique** is known as a thermal process for **obtaining low-energy state** of a solid in a heat bath.
- The process consists of the following two steps:
 - **Increasing temperature:** Increase the **temperature** of the heat bath to a maximum value at which the solid melts.
 - **Decreasing temperature:** Decrease carefully the temperature of the heat bath until the **particles** arrange themselves in the **ground state** of the solid.

Real Annealing Technique

- In the **liquid phase** all **particles** arrange themselves randomly, whereas in the ground state of the solid, the particles are arranged in a highly structured lattice, for which the corresponding energy is minimal.
- The **ground state** of the solid is obtained only if:
 - the maximum value of the temperature is sufficiently high and
 - the cooling is done sufficiently slow.
- Strong solid are grown from careful and slow cooling.

Real Annealing Technique

- **Metastable states**
 - If the initial temperature is not sufficiently high or a fast cooling is applied, **metastable states** (imperfections) are obtained.
- **Quenching**
 - The process that leads to metastable states is called **quenching**
- **Thermal equilibrium**
 - If the **lowering of the temperature** is done sufficiently slow, the solid can reach **thermal equilibrium** at each temperature.

Simulated Annealing: Part 1

Real Annealing and Simulated Annealing

- The analogy between the physical system and the optimization problem.

Physical System		Optimization Problem
System state	↔	Solution
Molecular positions	↔	Decision variables
Energy	↔	Objective function
Minimizing energy	↔	Minimizing cost
Ground state	↔	Global optimal solution
Metastable state	↔	Local optimum
Quenching	↔	Local search
Temperature	↔	Control parameter T
Real annealing	↔	Simulated annealing

Real Annealing and Simulated Annealing

- The objective function of the problem is analogous to the energy state of the system.
- A solution of the optimization problem corresponds to a system state.
- The decision variables associated with a solution of the problem are analogous to the molecular positions.
- The global optimum corresponds to the ground state of the system.
- Finding a local minimum implies that a metastable state has been reached.

Metropolis Algorithm



Metropolis Algorithm

- In 1958 Metropolis et al. introduced a simple algorithm for simulating the evolution of a solid in a heat bath to **thermal equilibrium**.
- Their algorithm is based on Monte Carlo techniques, and generates a sequence of states of the solid in the following way.
- Given a current state i of the solid with energy E_i , a subsequent state j is generated by applying a **perturbation mechanism** that transforms the current state into a next state by a small distortion, for instance, by a displacement of a single particle.

Metropolis Algorithm

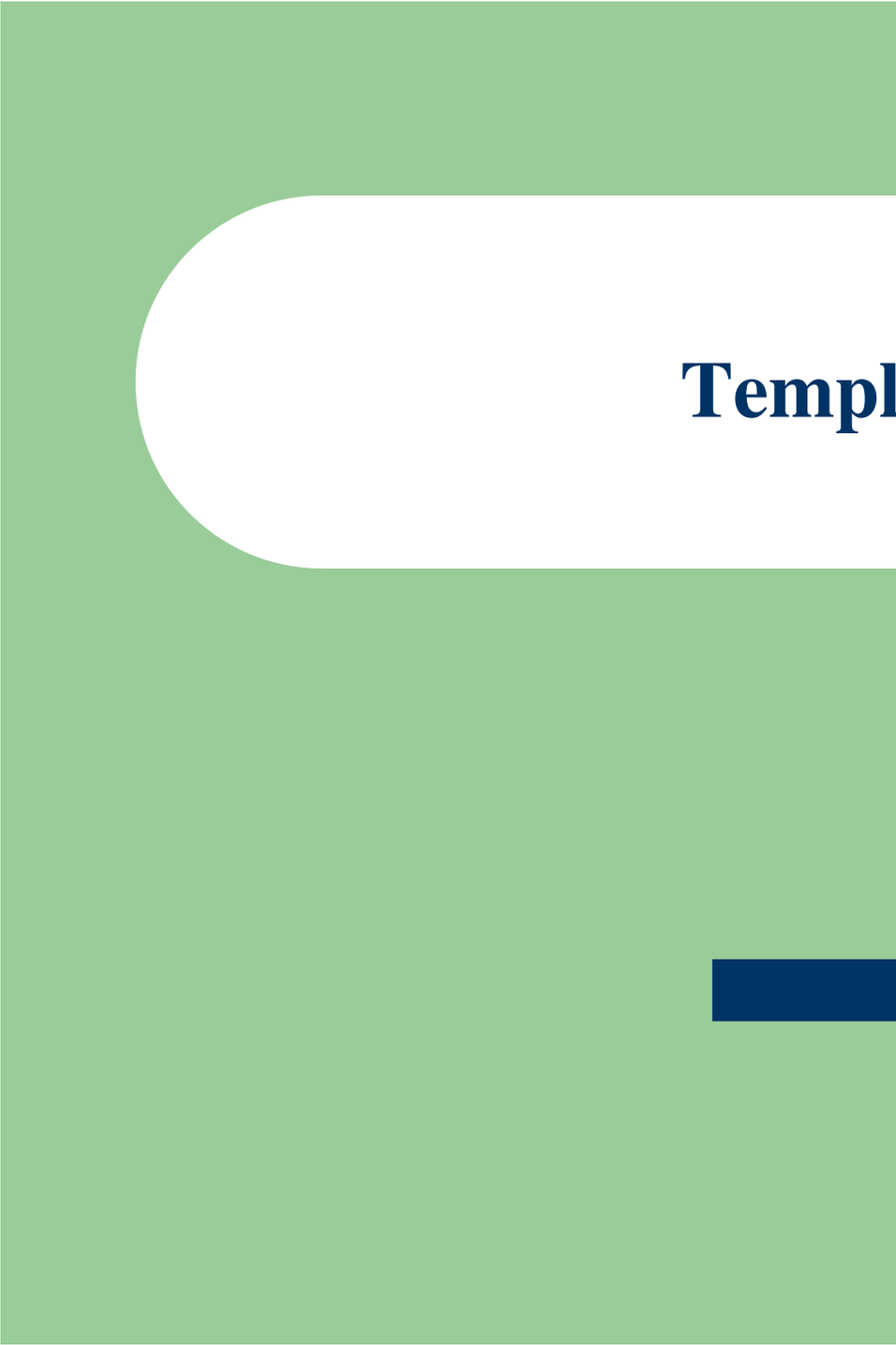
- The energy of the next state is E_j . (E_j)
- If the energy difference, $E_j - E_i$, is less than or equal to 0, the state j is accepted as the current state.
- If the energy difference is greater than 0, then state j is accepted with a probability given by

$$\exp\left(\frac{E_i - E_j}{k_B T}\right)$$

- where T denotes the temperature of the heat bath and
- k_B a physical constant known as the **Boltzmann constant**.

Metropolis Algorithm

- The acceptance rule described above is known as the **Metropolis criterion (Metropolis rule)** and the algorithm that goes with it is known as the **Metropolis algorithm**.
- In the Metropolis algorithm **thermal equilibrium** is achieved by generating a large number of transitions at a given temperature value.



Template of SA



Template of SA

- On the basis of a given, the system is subjected to an elementary,
- if this modification causes a decrease in the objective function of the system, it is accepted;
- if it causes an increase ΔE of the objective function, it is also accepted, but with a probability

$$e^{\frac{-\Delta E}{T}}$$

Template of SA

- By repeatedly observing this Metropolis rule of acceptance, a sequence of configurations is generated
- With this formalism in place, it is possible to show that, when the chain is of infinite length (in practical consideration, of “sufficient” length. . .), the system can reach (in practical consideration, can approach) **thermodynamic balance (Equilibrium State)** at the temperature considered

Simulated Annealing: Part 1

Template of SA

- At high temperature, $e^{\frac{-\Delta E}{T}}$ is close to 1,
 - therefore the majority of the moves are accepted and the algorithm becomes equivalent to a simple random walk in the configuration space .
- At low temperature, $e^{\frac{-\Delta E}{T}}$ is close to 0,
 - therefore the majority of the moves increasing energy is refused.
- At an intermediate temperature,
 - the algorithm intermittently authorizes the transformations that degrade the objective function

Template of SA

- SA can be viewed as **a sequence of Metropolis algorithms**, evaluated at decreasing values of the temperature.

Template of SA

- From an initial solution, SA proceeds in several iterations.
- At each iteration, a random neighbor is generated.
- Moves that improve the cost function are always accepted.
- Otherwise, the neighbor is selected with a given probability that depends on the current temperature and the amount of degradation ΔE of the objective function.
- ΔE represents the difference in the objective value (energy) between the current solution and the generated neighboring solution.

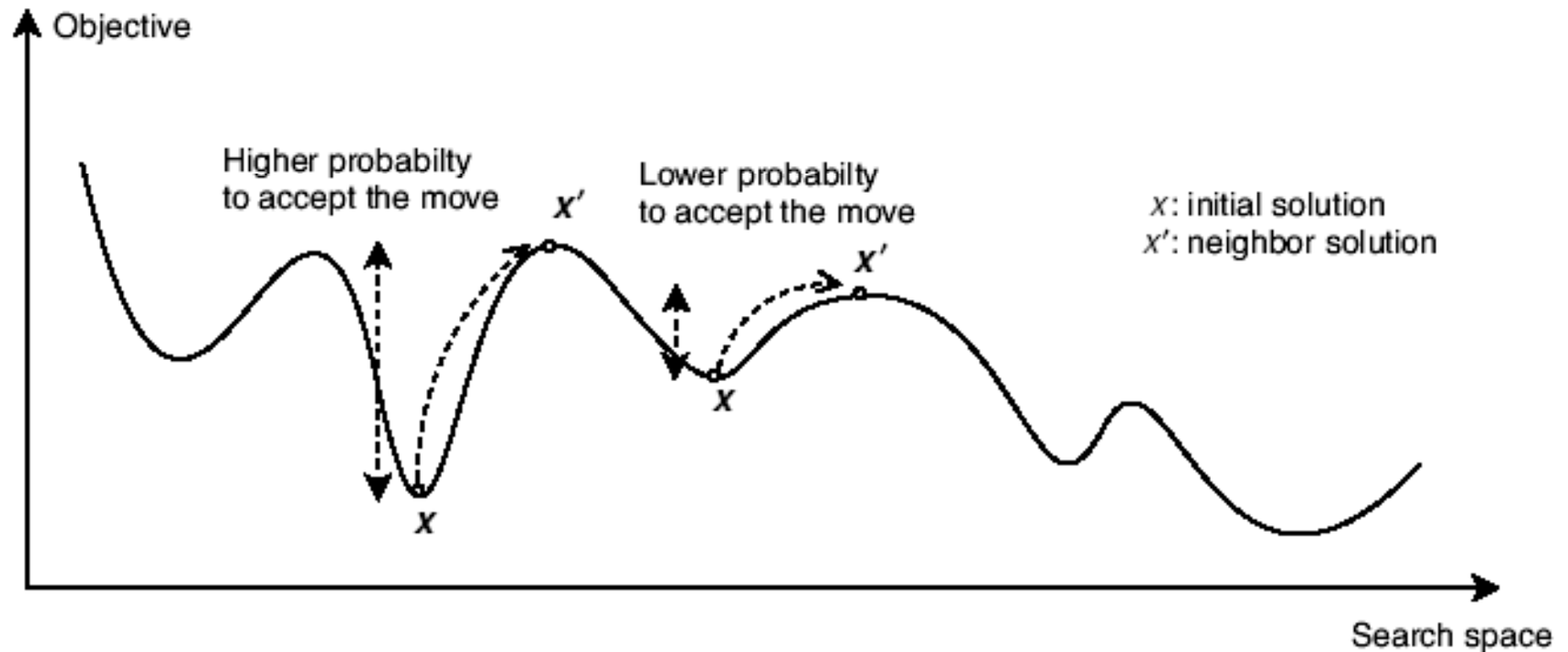
Template of SA

- The higher the temperature, the more significant the probability of accepting a worst move.
- At a given temperature, the lower the increase of the objective function, the more significant the probability of accepting the move.

Simulated Annealing: Part 1

Template of SA

- As the algorithm progresses, the probability that such moves are accepted decreases.



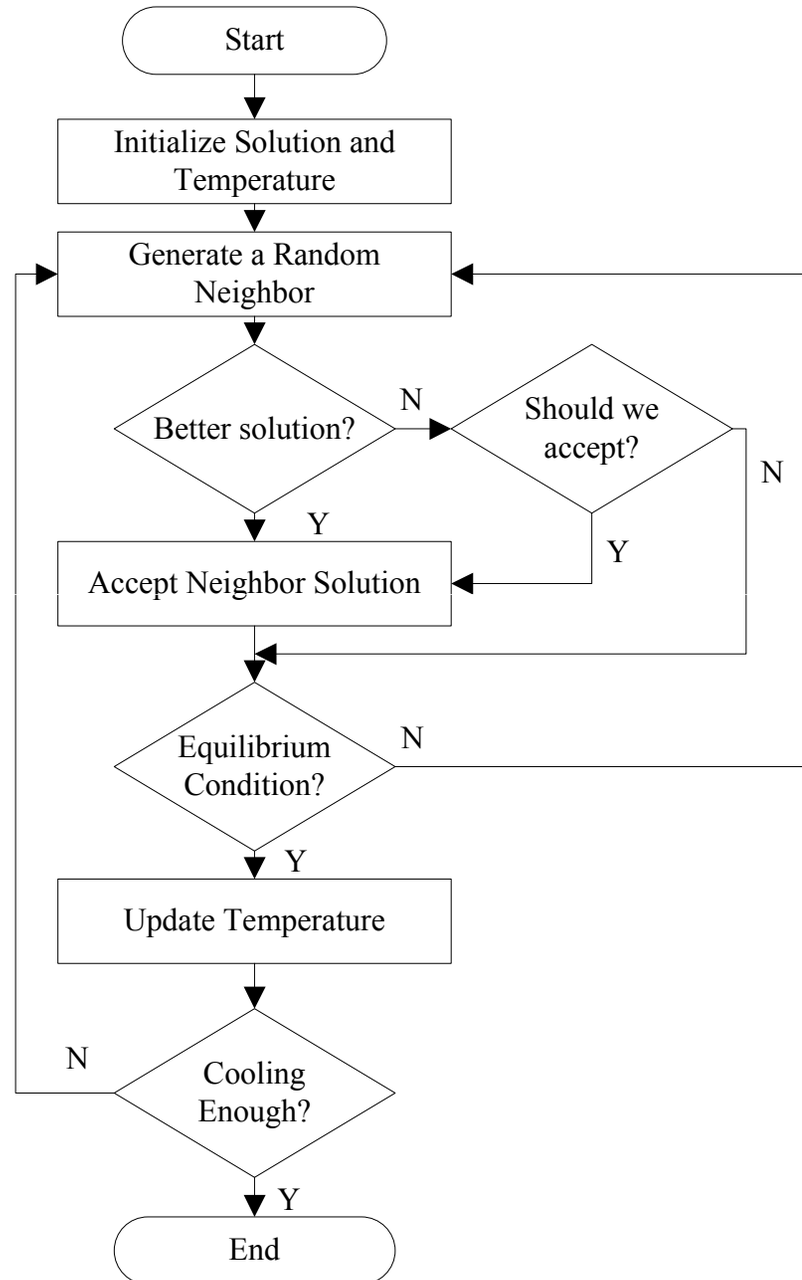
Template of SA

- **The acceptance probability function**, in general, the Boltzmann distribution:

$$P(\Delta E, T) = e^{-\frac{f(s') - f(s)}{T}}$$

- It uses a **control parameter**, called **temperature**, to determine the probability of accepting nonimproving solutions.
- At a particular level of temperature, many trials are explored.
- Once an equilibrium state is reached, the temperature is gradually decreased according to a cooling schedule such that few nonimproving solutions are accepted at the end of the search.

Simulated Annealing: Part 1



Simulated Annealing: Part 1

Template of SA

Input: Cooling schedule.

$s = s_0$; /* Generation of the initial solution */

$T = T_{max}$; /* Starting temperature */

Repeat

Repeat /* At a fixed temperature */

 Generate a random neighbor s' ;

$\Delta E = f(s') - f(s)$;

If $\Delta E \leq 0$ **Then** $s = s'$ /* Accept the neighbor solution */

Else Accept s' with a probability $e^{\frac{-\Delta E}{T}}$;

Until Equilibrium condition

 /* e.g. a given number of iterations executed at each temperature T */

$T = g(T)$; /* Temperature update */

Until Stopping criteria satisfied /* e.g. $T < T_{min}$ */

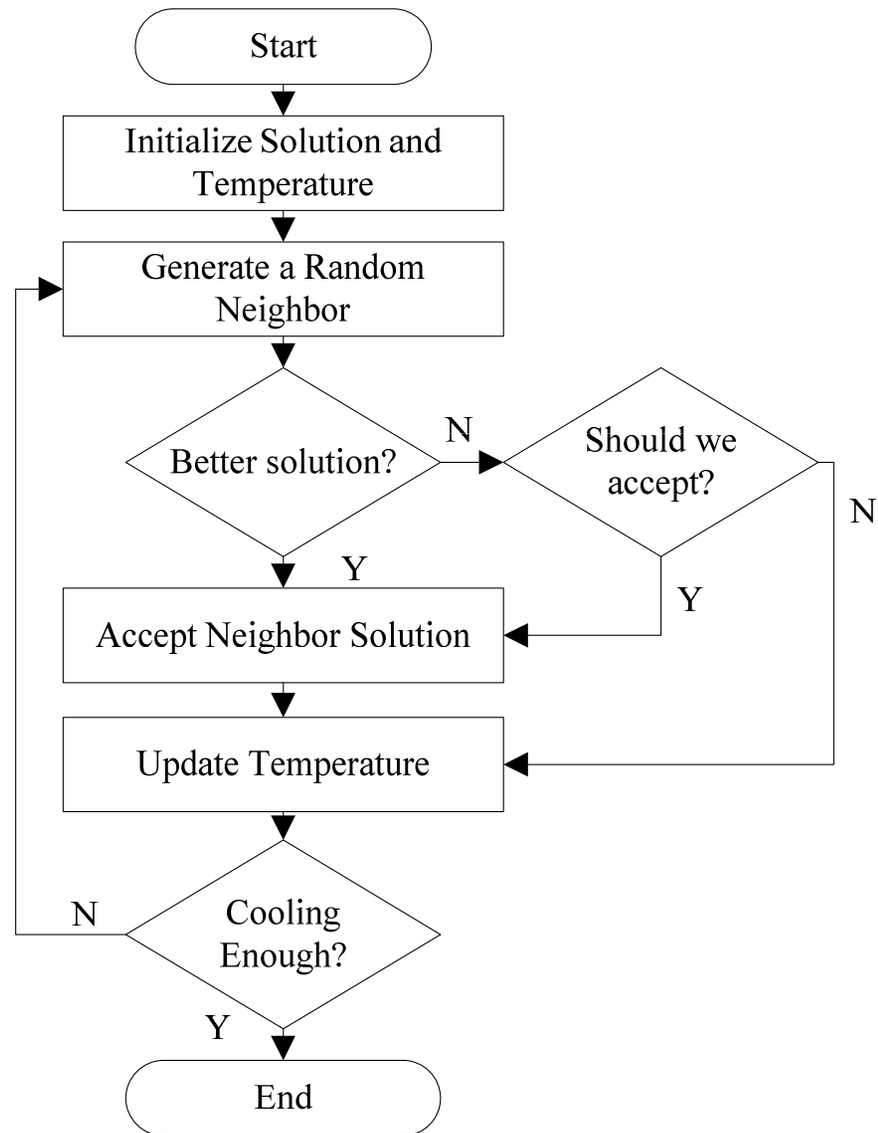
Output: Best solution found.

Inhomogeneous vs. Homogeneous Algorithm

- SA has two variants:
 - **Homogeneous variant**
 - Previous algorithm is the homogeneous variant
 - T is kept constant in the inner loop and is only decreased in the outer loop
 - **Inhomogeneous variant**
 - There is only one loop
 - T is decreased each time in the loop, but only very slightly

Simulated Annealing: Part 1

Inhomogeneous variant



Simulated Annealing: Part 1

Inhomogeneous variant

Input: Cooling schedule.

$s = s_0$; /* Generation of the initial solution */

$T = T_{max}$; /* Starting temperature */

Repeat

 Generate a random neighbor s' ;

$\Delta E = f(s') - f(s)$;

If $\Delta E \leq 0$ **Then** $s = s'$ /* Accept the neighbor solution */

Else Accept s' with a probability $e^{\frac{-\Delta E}{T}}$;

$T = g(T)$; /* Temperature update */

Until Stopping criteria satisfied /* e.g. $T < T_{min}$ */

Output: Best solution found.

Cooling Schedule

- The cooling schedule defines for each step of the algorithm i the temperature T_i .
- It has a great impact on the success of the SA optimization algorithm.
- The parameters to consider in defining a cooling schedule are:
 - the starting temperature,
 - the equilibrium state,
 - a cooling function, and
 - the final temperature that defines the stopping criteria

Template of SA

- Main components of SA:
 - Acceptance Function
 - Initial Temperature
 - Equilibrium State
 - Cooling Function
 - Stopping Condition



A Simple Example



Simulated Annealing: Part 1

A Simple Example

- Let us maximize the continuous function
 $f(x) = x^3 - 60x^2 + 900x + 100$.
- A solution x is represented as **a string of 5 bits**.
- The neighborhood consists in flipping randomly a bit.
- The initial solution is 10011 ($x = 19$, $f(x) = 2399$)
- Testing two sceneries:
 - First scenario: initial temperature T_0 equal to 500.
 - Second scenario: initial temperature T_0 equal to 100.
- Cooling: $T = 0.9 \cdot T$

A Simple Example

- In addition to the current solution, the best solution found since the beginning of the search is stored.
- Few parameters control the progress of the search, which are:
 - The temperature
 - The number of iterations performed at each temperature

Simulated Annealing: Part 1

A Simple Example

- First Scenario $T = 500$ and Initial Solution (10011)

T	Move	Solution	f	Δf	Move?	New Neighbor Solution
500	1	00011	2287	112	Yes	00011
450	3	00111	3803	<0	Yes	00111
405	5	00110	3556	247	Yes	00110
364.5	2	01110	3684	<0	Yes	01110
328	4	01100	3998	<0	Yes	01100
295.2	3	01000	3972	16	Yes	01000
265.7	4	01010	4100	<0	Yes	01010
239.1	5	01011	4071	29	Yes	01011
215.2	1	11011	343	3728	No	01011

Simulated Annealing: Part 1

A Simple Example

- Second Scenario: $T = 100$ and Initial Solution (10011).

T	Move	Solution	f	Δf	Move?	New Neighbor Solution
100	1	00011	2287	112	No	10011
90	3	10111	1227	1172	No	10011
81	5	10010	2692	< 0	Yes	10010
72.9	2	11010	516	2176	No	10010
65.6	4	10000	3236	< 0	Yes	10000
59	3	10100	2100	1136	Yes	10000

- When Temperature is not High Enough, Algorithm Gets Stuck



References



Simulated Annealing: Part 1

References

- El-Ghazali Talbi, **Metaheuristics : From Design to Implementation, John Wiley & Sons, 2009.**
- J. Dreco A. Petrowski, P. Siarry E. Taillard, **Metaheuristics for Hard Optimization, Springer-Verlag, 2006.**



The End

