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LAB RECORD

Bio Inspired Systems (23CS5BSBIS)

Submitted by

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in partial fulfillment for the award of the degree of

BACHELOR OF ENGINEERING in COMPUTER SCIENCE AND ENGINEERING



B.M.S. COLLEGE OF ENGINEERING
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CERTIFICATE

This is to certify that the Lab work entitled "Bio Inspired Systems (23CS5BSBIS)" carried out by **Shreyas Gouda M (1BM22CS270)**, who is bonafide student of **B.M.S.** College of **Engineering.** It is in partial fulfillment for the award of **Bachelor of Engineering in Computer Science and Engineering** of the Visvesvaraya Technological University, Belgaum. The Lab report has been approved as it satisfies the academic requirements of the above mention subject and the work prescribed for the said degree.

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Github Link:

https://github.com/SHREYASONI28/BIS-Lab

Genetic Algorithm for Optimization Problems

Genetic Algorithms (GA) are inspired by the process of natural selection and genetics, where the fittest individuals are selected for reproduction to produce the next generation. GAs are widely used for solving optimization and search problems. Implement a Genetic Algorithm using Python to solve a basic optimization problem, such as finding the maximum value of a mathematical function.

Implementation Steps:

- 1. Define the Problem: Create a mathematical function to optimize.
- 2. Initialize Parameters: Set the population size, mutation rate, crossover rate, and number of generations.
- 3. Create Initial Population: Generate an initial population of potential solutions.
- 4. Evaluate Fitness: Evaluate the fitness of each individual in the population.
- 5. Selection: Select individuals based on their fitness to reproduce.
- 6. Crossover: Perform crossover between selected individuals to produce offspring.
- 7. Mutation: Apply mutation to the offspring to maintain genetic diversity.
- 8. Iteration: Repeat the evaluation, selection, crossover, and mutation processes for a fixed number of generations or until convergence criteria are met.
- 9. Output the Best Solution: Track and output the best solution found during the generations.

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```
Code:
#GENETIC ALGORITHM
import numpy as np
import random
# Define the fitness function
def fitness function(x):
  return x ** 2
# Initialize parameters
population size = 100
mutation rate = 0.1
num generations = 50
bounds = (-10, 10)
# Step 1: Create initial population
def create initial population(size, bounds):
  return [random.uniform(bounds[0], bounds[1]) for in range(size)]
# Step 2: Evaluate fitness of the population
def evaluate population(population):
  return [fitness function(individual) for individual in population]
# Step 3: Selection using roulette-wheel selection
def selection(population, fitness):
  total fitness = sum(fitness)
  selection probs = [f/total] fitness for f in fitness
  return np.random.choice(population, size=2, p=selection probs)
# Step 4: Crossover operation
def crossover(parent1, parent2):
  alpha = random.uniform(0, 1)
  offspring1 = alpha * parent1 + (1 - alpha) * parent2
  offspring2 = alpha * parent2 + (1 - alpha) * parent1
  return offspring1, offspring2
# Step 5: Mutation operation
def mutate(individual, bounds):
  if random.random() < mutation rate:
     return random.uniform(bounds[0], bounds[1])
  return individual
# Main Genetic Algorithm loop
def genetic algorithm(bounds):
  # Step 1: Create initial population
  population = create initial population(population size, bounds)
```

```
best solution = None
  best fitness = float('-inf')
  for generation in range(num generations):
    # Step 2: Evaluate fitness
    fitness = evaluate population(population)
    # Track the best solution
    current best fitness = max(fitness)
    if current best fitness > best fitness:
       best fitness = current best fitness
       best solution = population[fitness.index(current best fitness)]
    # Step 3: Create new population
    new_population = []
    while len(new population) < population size:
       parent1, parent2 = selection(population, fitness)
       offspring1, offspring2 = crossover(parent1, parent2)
       new population.append(mutate(offspring1, bounds))
       new population.append(mutate(offspring2, bounds))
    # Replace the old population with the new population
    population = new population[:population size]
  return best solution, best fitness
# Run the Genetic Algorithm
best solution, best fitness = genetic algorithm(bounds)
print(f"Best Solution: x = {best solution}")
print(f"Best Fitness: f(x) = \{best fitness\}")
Output:
  Best Solution: x = 9.97704555295002
```

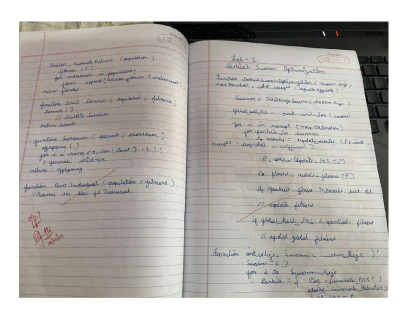
Best Fitness: f(x) = 99.54143796563977

Particle Swarm Optimization for Function Optimization

Particle Swarm Optimization (PSO) is inspired by the social behavior of birds flocking or fish schooling. PSO is used to find optimal solutions by iteratively improving a candidate solution with regard to a given measure of quality. Implement the PSO algorithm using Python to optimize a mathematical function.

Implementation Steps:

- 1. Define the Problem: Create a mathematical function to optimize.
- 2. Initialize Parameters: Set the number of particles, inertia weight, cognitive and social coefficients.
- 3. Initialize Particles: Generate an initial population of particles with random positions and velocities.
- 4. Evaluate Fitness: Evaluate the fitness of each particle based on the optimization function.
- 5. Update Velocities and Positions: Update the velocity and position of each particle based on its own best position and the global best position.
- 6. Iterate: Repeat the evaluation, updating, and position adjustment for a fixed number of iterations or until convergence criteria are met.
- 7. Output the Best Solution: Track and output the best solution found during the iterations.



Code: **#PARTICLE SWARM OPTIMIZATION** import numpy as np # Rastrigin function: A benchmark function for optimization problems def rastrigin(x): A = 10# Calculate the Rastrigin function value based on the input vector x return A * len(x) + sum(x i^{**2} - A * np.cos(2 * np.pi * x i) for x i in x) # Particle Swarm Optimization class class ParticleSwarmOptimizer: def init (self, func, n particles, n dimensions, n iterations, inertia weight=0.7, cognitive coeff=1.5, social coeff=1.5, bounds=(-5.12, 5.12)): self.func = func # The function to optimize self.n particles = n particles # Number of particles in the swarm self.n dimensions = n dimensions # Dimensions of the search space self.n iterations = n iterations # Number of iterations for the optimization self.lower bound, self.upper bound = bounds # Bounds for the search space # Initialize particle positions randomly within the specified bounds self.positions = np.random.uniform(self.lower bound, self.upper bound, (n particles, n dimensions)) # Initialize particle velocities randomly self.velocities = np.random.uniform(-1, 1, (n particles, n dimensions)) # Personal best positions and scores for each particle self.pbest positions = np.copy(self.positions) self.pbest scores = np.array([func(p) for p in self.positions]) # Evaluate initial fitness # Global best position and score among all particles self.gbest position = self.pbest positions[np.argmin(self.pbest scores)] self.gbest score = np.min(self.pbest scores) def optimize(self): # Main loop for the optimization process for in range(self.n iterations): for i in range(self.n particles): # Evaluate the fitness of the current position fitness = self.func(self.positions[i]) # Update personal best if the current fitness is better if fitness < self.pbest scores[i]: self.pbest scores[i] = fitness self.pbest positions[i] = self.positions[i] # Update global best if the current fitness is better if fitness < self.gbest score: self.gbest score = fitness self.gbest position = self.positions[i]

```
# Generate random coefficients for cognitive and social components
       r1, r2 = np.random.rand(self.n dimensions), np.random.rand(self.n dimensions)
       # Update velocities based on inertia, personal best, and global best
       self.velocities = (self.velocities * 0.7 + # Inertia weight
                   1.5 * r1 * (self.pbest positions - self.positions) + # Cognitive component
                   1.5 * r2 * (self.gbest position - self.positions)) # Social component
       # Update positions based on new velocities and clip to stay within bounds
       self.positions = np.clip(self.positions + self.velocities, self.lower bound, self.upper bound)
       # Print the best fitness found so far in this iteration
       print(f"Best Fitness: {self.gbest score}")
    # Return the best position and score found after all iterations
    return self.gbest position, self.gbest score
# Create and run the optimizer
pso = ParticleSwarmOptimizer(func=rastrigin, n particles=30, n dimensions=2, n iterations=100)
best position, best score = pso.optimize()
# Print the best position and corresponding fitness score found
print("\nBest Position Found:", best position)
print("Best Fitness Score:", best score)
```

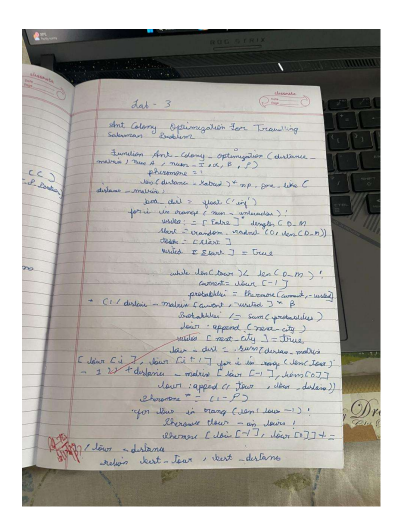
Output:

```
Best Fitness: 7.523349690449162
                                                   Best Fitness: 0.00042484148907107055
Best Fitness: 5.479012944526062
                                                   Best Fitness: 0.00042484148907107055
Best Fitness: 5.479012944526062
                                                   Best Fitness: 0.00019896058490331825
Best Fitness: 5.479012944526062
                                                   Best Fitness: 0.00019896058490331825
                                                   Best Fitness: 0.00019896058490331825
Best Fitness: 5.479012944526062
                                                   Best Fitness: 9.185587018123442e-06
                                                   Best Fitness: 9.185587018123442e-06
Best Fitness: 5.35158483420342
                                                   Best Fitness: 9.185587018123442e-06
Best Fitness: 4.23336222695108
                                                   Best Fitness: 9.185587018123442e-06
Best Fitness: 2.3059731550465656
                                                   Best Fitness: 9.185587018123442e-06
Best Fitness: 2.3059731550465656
                                                   Best Fitness: 9.185587018123442e-06
Best Fitness: 2.3059731550465656
                                                   Best Fitness: 9.185587018123442e-06
Best Fitness: 2.3059731550465656
Best Fitness: 2.2923779383497873
Best Fitness: 2.2923779383497873
Best Fitness: 2.2923779383497873
                                                   Best Fitness: 8.17354336390963e-06
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                                                   Best Fitness: 3.993851240835511e-06
                                                   Best Fitness: 3.993851240835511e-06
                                                   Best Fitness: 3.993851240835511e-06
Best Fitness: 2.2923779383497873
                                                   Best Fitness: 3.993851240835511e-06
Best Fitness: 2.2923779383497873
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Best Fitness: 2.2923779383497873
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Best Fitness: 2.2923779383497873
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Best Fitness: 2.2923779383497873
                                                   Best Fitness: 1.6462023069152565e-06
Best Fitness: 2.2923779383497873
                                                   Best Fitness: 1.6462023069152565e-06
Best Fitness: 1.4393114014934305
                                                   Best Fitness: 1.6462023069152565e-06
Best Fitness: 1.4393114014934305
                                                   Best Fitness: 1.5764355048020207e-06
Best Fitness: 1.3002025319518147
                                                   Best Position Found: [3.76308963e-05 8.08082678e-05]
Best Fitness: 1.3002025319518147
                                                   Best Fitness Score: 1.5764355048020207e-06
Best Fitness: 1.3002025319518147
```

Ant Colony Optimization for the Traveling Salesman Problem

The foraging behaviour of ants has inspired the development of optimization algorithms that can solve complex problems such as the Traveling Salesman Problem (TSP). Ant Colony Optimization (ACO) simulates the way ants find the shortest path between food sources and their nest. Implement the ACO algorithm using Python to solve the TSP, where the objective is to find the shortest possible route that visits a list of cities and returns to the origin city. Implementation Steps:

- 1. Define the Problem: Create a set of cities with their coordinates.
- 2. Initialize Parameters: Set the number of ants, the importance of pheromone (alpha), the importance of heuristic information (beta), the evaporation rate (rho), and the initial pheromone value.
- 3. Construct Solutions: Each ant constructs a solution by probabilistically choosing the next city based on pheromone trails and heuristic information.
- 4. Update Pheromones: After all ants have constructed their solutions, update the pheromone trails based on the quality of the solutions found.
- 5. Iterate: Repeat the construction and updating process for a fixed number of iterations or until convergence criteria are met.
- 6. Output the Best Solution: Keep track of and output the best solution found during the iterations.



```
Code:
#ANT COLONY OPTIMIZATION
import random
import numpy as np
# Distance calculation (Euclidean distance)
def euclidean distance(city1, city2):
  return np.sqrt((city1[0] - city2[0])**2 + (city1[1] - city2[1])**2)
# Ant Colony Optimization Algorithm
class ACO:
  def init (self, cities, num ants=10, num iterations=100, alpha=1.0, beta=2.0, rho=0.5, Q=100):
    self.cities = cities
    self.num ants = num ants
    self.num iterations = num iterations
    self.alpha = alpha
                           # Importance of pheromone
                          # Importance of heuristic information (distance)
    self.beta = beta
                         # Pheromone evaporation rate
    self.rho = rho
                         # Total pheromone deposited per ant per tour
    self.Q = Q
    self.num cities = len(cities)
    # Initialize pheromone matrix (for each pair of cities)
     self.pheromone = np.ones((self.num cities, self.num cities)) / self.num cities
     self.distances = np.zeros((self.num cities, self.num cities))
    # Compute distance matrix
     for i in range(self.num cities):
       for j in range(i + 1, self.num cities):
         self.distances[i][j] = self.distances[j][i] = euclidean distance(cities[i], cities[j])
  def choose next city(self, ant, visited):
    # Calculate the probability of moving to each city
    current city = ant[-1]
    probabilities = []
     for i in range(self.num cities):
       if i not in visited:
         pheromone = self.pheromone[current city][i] ** self.alpha
         heuristic = (1.0 / self.distances[current city][i]) ** self.beta
         probabilities.append(pheromone * heuristic)
         probabilities.append(0)
    # Normalize probabilities
    total = sum(probabilities)
    if total == 0: # In case there's no valid path (shouldn't happen with good settings)
```

```
return random.choice([i for i in range(self.num cities) if i not in visited])
  probabilities = [prob / total for prob in probabilities]
  # Choose next city based on probabilities
  next city = random.choices(range(self.num cities), probabilities)[0]
  return next city
def construct solution(self):
  # Each ant starts at a random city
  ant = [random.randint(0, self.num cities - 1)]
  visited = set(ant)
  while len(ant) < self.num cities:
    next city = self. choose next city(ant, visited)
    ant.append(next city)
    visited.add(next city)
  # Return to the starting city
  ant.append(ant[0])
  return ant
def evaluate solution(self, solution):
  # Calculate the total distance of the tour
  total distance = 0
  for i in range(len(solution) - 1):
    total distance += self.distances[solution[i]][solution[i + 1]]
  return total distance
def update pheromone(self, all solutions):
  # Initialize pheromone update matrix
  pheromone delta = np.zeros((self.num cities, self.num cities))
  # For each solution, deposit pheromone
  for solution in all solutions:
    tour length = self. evaluate solution(solution)
     for i in range(len(solution) - 1):
       pheromone delta[solution[i]][solution[i+1]] += self.Q / tour length
  # Evaporate pheromone
  self.pheromone = (1 - self.rho) * self.pheromone + pheromone delta
def solve(self):
  best solution = None
  best distance = float('inf')
```

```
for iteration in range(self.num iterations):
       all solutions = []
       # Each ant constructs a solution
       for ant in range(self.num ants):
          solution = self. construct solution()
          all solutions.append(solution)
          tour length = self. evaluate solution(solution)
          # Update best solution if necessary
          if tour length < best distance:
            best solution = solution
            best distance = tour length
       # Update pheromones based on solutions found
       self. update pheromone(all solutions)
       print(f"Iteration {iteration + 1}, Best Distance: {best distance}")
     return best solution, best distance
# Function to take user input for cities
def get user input():
  num cities = int(input("Enter the number of cities: "))
  cities = []
  print("Enter the coordinates of each city (x, y):")
  for i in range(num cities):
    x, y = map(float, input(f''City \{i+1\}: ").split())
     cities.append((x, y))
  return cities
# Example usage:
if __name__== "__main__":
  # Take user input for cities
  cities = get user input()
  # Take user input for ACO parameters
  num ants = int(input("Enter the number of ants: "))
  num iterations = int(input("Enter the number of iterations: "))
  alpha = float(input("Enter the value of alpha (pheromone importance): "))
  beta = float(input("Enter the value of beta (distance importance): "))
  rho = float(input("Enter the value of rho (pheromone evaporation rate): "))
  Q = float(input("Enter the value of Q (pheromone deposit per ant): "))
  # Create an instance of ACO and solve the problem
```

```
aco = ACO(cities, num_ants, num_iterations, alpha, beta, rho, Q)
best_solution, best_distance = aco.solve()
print(f"\nBest Solution (Tour): {best_solution}")
print(f"Best Distance: {best_distance}")
```

Output:

```
Enter the number of cities: 5
Enter the coordinates of each city (x, y):
City 1: 0 0
City 2: 1 3
City 3: 4 3
City 4: 6 1
City 5: 3 0
Enter the number of ants: 10
Enter the number of iterations: 100
Enter the value of alpha (pheromone importance): 1.0
Enter the value of beta (distance importance): 2.0
Enter the value of rho (pheromone evaporation rate): 0.5
Enter the value of Q (pheromone deposit per ant): 100
```

```
Best Solution (Tour): [1, 0, 4, 3, 2, 1]
Best Distance: 15.15298244508295
```

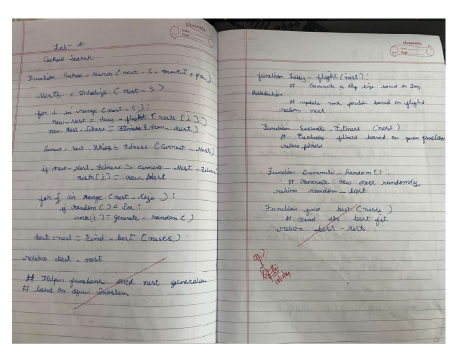
Cuckoo Search (CS)

Cuckoo Search (CS) is a nature-inspired optimization algorithm based on the brood parasitism of some cuckoo species. This behavior involves laying eggs in the nests of other birds, leading to the optimization of survival strategies. CS uses Lévy flights to generate new solutions, promoting global search capabilities and avoiding local minima. The algorithm is widely used for solving continuous optimization problems and has applications in various domains, including engineering design, machine learning, and data mining.

Implementation Steps:

- 1. Define the Problem: Create a mathematical function to optimize.
- 2. Initialize Parameters: Set the number of nests, the probability of discovery, and the number of iterations.
- 3. Initialize Population: Generate an initial population of nests with random positions.
- 4. Evaluate Fitness: Evaluate the fitness of each nest based on the optimization function.
- 5. Generate New Solutions: Create new solutions via Lévy flights.
- 6. Abandon Worst Nests: Abandon a fraction of the worst nests and replace them with new random positions.
- 7. Iterate: Repeat the evaluation, updating, and replacement process for a fixed number of iterations or until convergence criteria are met.
- 8. Output the Best Solution: Track and output the best solution found during the iterations.

Algorithm:



Code: #CUCKOO SEARCH

import numpy as np

```
import math # Import the standard math module def levy_flight(Lambda): sigma = (math.gamma(1 + Lambda) * math.sin(math.pi * Lambda / 2) / \\ (math.gamma((1 + Lambda) / 2) * Lambda * 2 ** ((Lambda - 1) / 2))) ** (1 / Lambda) \\ u = np.random.normal(0, sigma, 1) \\ v = np.random.normal(0, 1, 1) \\ step = u / abs(v) ** (1 / Lambda) \\ return step
```

```
def cuckoo search(obj function, bounds, n=25, pa=0.25, max iter=100):
  # Initialize nests
  dim = len(bounds)
  nests = np.random.rand(n, dim)
  for i in range(dim):
     nests[:, i] = nests[:, i] * (bounds[i][1] - bounds[i][0]) + bounds[i][0]
  fitness = np.array([obj function(nest) for nest in nests])
  # Start optimization
  for in range(max iter):
     for i in range(n):
       # Generate a new solution via Levy flight
       new nest = nests[i] + levy flight(1.5) * np.random.randn(dim)
       # Apply bounds
       new nest = np.clip(new nest, [b[0]] for b in bounds], [b[1]] for b in bounds])
       new fitness = obj function(new nest)
       # Update if new solution is better
       if new fitness < fitness[i]:
          nests[i] = new nest
          fitness[i] = new fitness
     # Abandon some nests and create new ones
     abandon idx = np.random.rand(n) < pa
     for i in np.where(abandon idx)[0]:
       nests[i] = np.random.rand(dim) * (np.array([b[1] for b in bounds]) - np.array([b[0] for b in bounds])) - np.array([b[0] for b in bounds]))
bounds])) + np.array([b[0] for b in bounds])
       fitness[i] = obj function(nests[i])
  # Return the best solution
  best idx = np.argmin(fitness)
  return nests[best idx], fitness[best idx]
# Example usage: Minimize f(x) = x^2
def objective(x):
  return sum(xi**2 for xi in x)
bounds = [(-10, 10), (-10, 10)] # 2D problem
best solution, best fitness = cuckoo search(objective, bounds)
print("Best Solution:", best solution)
print("Best Fitness:", best fitness)
```

Output:

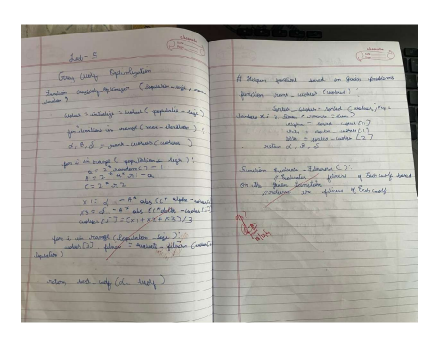
Best Solution: [-0.14023741 0.59049343]
Best Fitness: 0.36834901994989167

Grey Wolf Optimizer (GWO)

The Grey Wolf Optimizer (GWO) algorithm is a swarm intelligence algorithm inspired by the social hierarchy and hunting behavior of grey wolves. It mimics the leadership structure of alpha, beta, delta, and omega wolves and their collaborative hunting strategies. The GWO algorithm uses these social hierarchies to model the optimization process, where the alpha wolves guide the search process while beta and delta wolves assist in refining the search direction. This algorithm is effective for continuous optimization problems and has applications in engineering, data analysis, and machine learning.

Implementation Steps:

- 1. Define the Problem: Create a mathematical function to optimize.
- 2. Initialize Parameters: Set the number of wolves and the number of iterations.
- 3. Initialize Population: Generate an initial population of wolves with random positions.
- 4. Evaluate Fitness: Evaluate the fitness of each wolf based on the optimization function.
- 5. Update Positions: Update the positions of the wolves based on the positions of alpha, beta, and delta wolves.
- 6. Iterate: Repeat the evaluation and position updating process for a fixed number of iterations or until convergence criteria are met.
- 7. Output the Best Solution: Track and output the best solution found during the iterations.



```
Code:
#Grey Wolf Optimizer (GWO)
import numpy as np
def objective function(x):
  """Example objective function: Sphere function."""
  return sum(x^{**}2)
definitialize population(dim, n wolves, bounds):
  """Initialize the positions of the wolves randomly within the given bounds."""
  return np.random.uniform(bounds[0], bounds[1], (n wolves, dim))
def gwo(objective function, bounds, dim, n wolves, n iterations):
  # Initialize population
  wolves = initialize population(dim, n wolves, bounds)
  fitness = np.apply along axis(objective function, 1, wolves)
  # Initialize alpha, beta, and delta
  alpha, beta, delta = np.argsort(fitness)[:3]
  alpha pos, alpha score = wolves[alpha], fitness[alpha]
  beta pos, beta score = wolves[beta], fitness[beta]
  delta pos, delta score = wolves[delta], fitness[delta]
  # Main optimization loop
  for iteration in range(n iterations):
    a = 2 - 2 * (iteration / n iterations) # Linearly decreasing a
     for i in range(n wolves):
       for j in range(dim):
```

```
# Update each wolf's position
          r1, r2 = np.random.rand(), np.random.rand()
          A1, C1 = 2 * a * r1 - a, 2 * r2
          D alpha = abs(C1 * alpha pos[j] - wolves[i, j])
          X1 = alpha pos[i] - A1 * D alpha
          r1, r2 = np.random.rand(), np.random.rand()
          A2, C2 = 2 * a * r1 - a, 2 * r2
          D beta = abs(C2 * beta pos[i] - wolves[i, i])
          X2 = beta pos[i] - A2 * D beta
          r1, r2 = np.random.rand(), np.random.rand()
          A3, C3 = 2 * a * r1 - a, 2 * r2
          D delta = abs(C3 * delta pos[j] - wolves[i, j])
          X3 = delta pos[i] - A3 * D delta
          # Average position update
          wolves[i, j] = (X1 + X2 + X3) / 3.0
       # Enforce bounds
       wolves[i, :] = np.clip(wolves[i, :], bounds[0], bounds[1])
     # Evaluate fitness and update alpha, beta, delta
     fitness = np.apply along axis(objective function, 1, wolves)
     sorted indices = np.argsort(fitness)
     alpha, beta, delta = sorted indices[:3]
     alpha pos, alpha score = wolves[alpha], fitness[alpha]
     beta pos, beta score = wolves[beta], fitness[beta]
     delta pos, delta score = wolves[delta], fitness[delta]
  return alpha pos, alpha score
# Example usage
dim = 5 # Number of dimensions
bounds = (-10, 10) # Search space bounds
n wolves = 30 # Number of wolves
n iterations = 100 # Number of iterations
best solution, best score = gwo(objective function, bounds, dim, n wolves, n iterations)
print(f"Best solution: {best solution}")
print(f"Best score: {best score}")
Output:
```

Best solution: [-1.48263895e-11 -1.24732979e-11 1.51277899e-11 1.54330567e-11

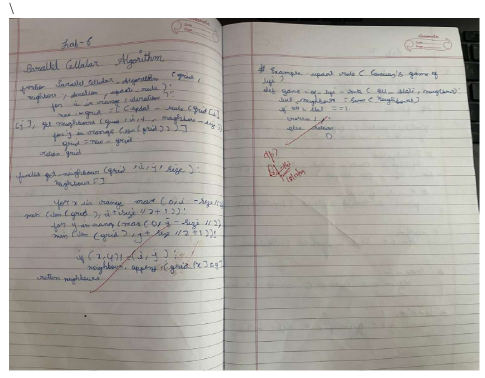
1.16834722e-11]

Best score: 9.78937775690888e-22

Parallel Cellular Algorithms and Programs

Parallel Cellular Algorithms are inspired by the functioning of biological cells that operate in a highly parallel and distributed manner. These algorithms leverage the principles of cellular automata and parallel computing to solve complex optimization problems efficiently. Each cell represents a potential solution and interacts with its neighbors to update its state based on predefined rules. This interaction models the diffusion of information across the cellular grid, enabling the algorithm to explore the search space effectively. Parallel Cellular Algorithms are particularly suitable for large-scale optimization problems and can be implemented on parallel computing architectures for enhanced performance. Implementation Steps:

- 1. Define the Problem: Create a mathematical function to optimize.
- 2. Initialize Parameters: Set the number of cells, grid size, neighborhood structure, and number of iterations.
- 3. Initialize Population: Generate an initial population of cells with random positions in the solution space.
- 4. Evaluate Fitness: Evaluate the fitness of each cell based on the optimization function.
- 5. Update States: Update the state of each cell based on the states of its neighboring cells and predefined update rules.
- 6. Iterate: Repeat the evaluation and state updating process for a fixed number of iterations or until convergence criteria are met.
- 7. Output the Best Solution: Track and output the best solution found during the iterations.



```
Code:
#Parallel Cellular Algorithms and Programs
import numpy as np
# Define the optimization function
def fitness function(x):
  return x^{**}2
# Initialize parameters
num cells = 10
grid size = 1.0
iterations = 100
neighborhood_size = 1
# Initialize population
cells = np.random.uniform(-grid size, grid size, num cells)
# Main loop
for in range(iterations):
  # Evaluate fitness
  fitness = np.array([fitness function(cell) for cell in cells])
  # Update states
  new cells = np.copy(cells)
  for i in range(num_cells):
    # Get neighbors
    neighbors = cells[max(0, i-neighborhood size):min(num cells, i+neighborhood size+1)]
    # Update cell based on neighbors
    new cells[i] = np.mean(neighbors) + np.random.uniform(-0.1, 0.1) # Add some noise
  cells = new cells
# Output the best solution
best cell = cells[np.argmin(fitness)]
print(f"Best solution found: {best cell}")
print(f"Fitness: {fitness function(best cell)}")
Output:
Best solution found: -0.11165744078455692
Fitness: 0.012467384082556834
```

Optimization via Gene Expression Algorithms

Gene Expression Algorithms (GEA) are inspired by the biological process of gene expression in living organisms. This process involves the translation of genetic information encoded in DNA into functional proteins. In GEA, solutions to optimization problems are encoded in a manner similar to genetic sequences. The algorithm evolves these solutions through selection, crossover, mutation, and gene expression to find optimal or near-optimal solutions. GEA is effective for solving complex optimization problems in various domains, including engineering, data analysis, and machine learning.

Implementation Steps:

- 1. Define the Problem: Create a mathematical function to optimize.
- 2. Initialize Parameters: Set the population size, number of genes, mutation rate, crossover rate, and number of generations.
- 3. Initialize Population: Generate an initial population of random genetic sequences.
- 4. Evaluate Fitness: Evaluate the fitness of each genetic sequence based on the optimization function.
- 5. Selection: Select genetic sequences based on their fitness for reproduction.
- 6. Crossover: Perform crossover between selected sequences to produce offspring.
- 7. Mutation: Apply mutation to the offspring to introduce variability.
- 8. Gene Expression: Translate genetic sequences into functional solutions.
- 9. Iterate: Repeat the selection, crossover, mutation, and gene expression processes for a fixed number of generations or until convergence criteria are met.
- 10. Output the Best Solution: Track and output the best solution found during the iterations.

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```
Code:
#Optimization via Gene Expression Algorithms
import numpy as np
# Define the optimization function
def fitness function(x):
  return x^{**}2
# Convert binary string to decimal
def binary to decimal(binary str):
  return int(binary_str, 2) / (2**len(binary_str) - 1) * 10 - 5 # Scale to [-5, 5]
# Initialize parameters
population size = 20
num genes = 10
mutation rate = 0.1
crossover rate = 0.7
generations = 100
# Initialize population
population = [".join(np.random.choice(['0', '1'], num genes)) for in range(population size)]
# Main loop
for in range(generations):
  # Evaluate fitness
```

```
fitness = [fitness function(binary to decimal(ind)) for ind in population]
  # Selection (roulette wheel)
  total fitness = sum(fitness)
  probabilities = [f / total fitness for f in fitness]
  selected = np.random.choice(population, size=population size, p=probabilities)
  # Crossover
  offspring = []
  for i in range(0, population size, 2):
     if np.random.rand() < crossover rate:
       point = np.random.randint(1, num genes)
       offspring.append(selected[i][:point] + selected[i+1][point:])
       offspring.append(selected[i+1][:point] + selected[i][point:])
     else:
       offspring.append(selected[i])
       offspring.append(selected[i+1])
  # Mutation
  for i in range(population size):
     if np.random.rand() < mutation rate:
       point = np.random.randint(num genes)
       offspring[i] = offspring[i][:point] + ('1' if offspring[i][point] == '0' else '0') +
offspring[i][point+1:]
  population = offspring
# Output the best solution
best individual = min(population, key=lambda ind: fitness function(binary to decimal(ind)))
best fitness = fitness function(binary to decimal(best individual))
print(f"Best solution found: {binary to decimal(best individual)}")
print(f" Fitness: {best fitness}")
Output:
 Best solution found: -4.872922776148583
   Fitness: 23.74537638230761
```