SET OF PROBLEMS

// Initialize particles

```
1. Particle Motion Simulator
Description:
Simulate the motion of particles in a two-dimensional space under the influence of forces.
Specifications:
Structure: Represents particle properties (mass, position, velocity).
Array: Stores the position and velocity vectors of multiple particles.
Union: Handles force types (gravitational, electric, or magnetic).
Strings: Define force types applied to particles.
const Pointers: Protect particle properties.
Double Pointers: Dynamically allocate memory for the particle system.
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
// Define constants for forces
#define GRAVITY 9.8
#define ELECTRIC CONSTANT 8.99e9
#define MAGNETIC CONSTANT 1.0
// Force types enumeration
typedef enum {
  GRAVITATIONAL,
  ELECTRIC.
  MAGNETIC
} ForceType;
// Particle structure
typedef struct {
  double mass:
                    // Mass of the particle (kg)
  double position[2]; // Position in 2D (x, y)
  double velocity[2]; // Velocity in 2D (vx, vy)
} Particle;
// Union for force types
typedef union {
  double gravitational_force;
  double electric force;
  double magnetic_force;
} Force:
// Function prototypes
void updateParticlePosition(Particle* particle, double time_interval);
void applyForce(Particle* particle, ForceType force_type, double force_magnitude);
void printParticle(Particle* particle);
int main() {
  // Dynamically allocate memory for an array of particles using double pointers
  int num_particles = 3; // Example for 3 particles
  Particle** particles = (Particle**) malloc(num_particles * sizeof(Particle*));
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for (int i = 0; i < num particles; <math>i++) {
     particles[i] = (Particle*) malloc(sizeof(Particle));
     particles[i]->mass = 1.0 + i; // Assign mass of each particle
     particles[i]->position[0] = i * 1.0; // Assign initial position
     particles[i]->position[1] = 0.0;
     particles[i]->velocity[0] = 0.0; // Initial velocity
     particles[i]->velocity[1] = 0.0;
  }
  // Apply forces to particles and update positions
  applyForce(particles[0], GRAVITATIONAL, 9.8); // Gravitational force on particle 0
  applyForce(particles[1], ELECTRIC, 5.0); // Electric force on particle 1
  applyForce(particles[2], MAGNETIC, 3.0); // Magnetic force on particle 2
  // Update particle positions based on forces and time intervals
  double time_interval = 1.0; // 1 second time step
  for (int i = 0; i < num_particles; i++) {
     updateParticlePosition(particles[i], time_interval);
     printParticle(particles[i]);
  }
  // Free allocated memory
  for (int i = 0; i < num particles; <math>i++) {
     free(particles[i]);
  free(particles);
  return 0;
}
// Function to update particle position based on velocity
void updateParticlePosition(Particle* particle, double time_interval) {
  particle->position[0] += particle->velocity[0] * time interval;
  particle->position[1] += particle->velocity[1] * time_interval;
}
// Function to apply a force to a particle
void applyForce(Particle* particle, ForceType force_type, double force_magnitude) {
  switch (force_type) {
     case GRAVITATIONAL:
        particle->velocity[1] += force magnitude / particle->mass; // Simplified gravity
       break;
     case ELECTRIC:
       // Simple electric force, assuming particle charge = 1 (for simplicity)
        particle->velocity[0] += force magnitude / particle->mass;
       break;
     case MAGNETIC:
       // Simplified magnetic force, assuming perpendicular motion to magnetic field
       particle->velocity[0] += force_magnitude / particle->mass; // Simplification
       break;
}
// Function to print the particle details
void printParticle(Particle* particle) {
```

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printf("Particle - Mass: %.2f, Position: (%.2f, %.2f), Velocity: (%.2f, %.2f)\n",
       particle->mass, particle->position[0], particle->position[1],
       particle->velocity[0], particle->velocity[1]);
}
2. Electromagnetic Field Calculator
Description:
Calculate the electromagnetic field intensity at various points in space.
Specifications:
Structure: Stores field parameters (electric field, magnetic field, and position).
Array: Holds field values at discrete points.
Union: Represents either electric or magnetic field components.
Strings: Represent coordinate systems (Cartesian, cylindrical, spherical).
const Pointers: Prevent modification of field parameters.
Double Pointers: Manage memory for field grid allocation dynamically.
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
#define MAX POINTS 100
// Define a union to hold either electric or magnetic field components
union FieldComponent {
  double electric[3]; // Electric field components (Ex, Ey, Ez)
  double magnetic[3]; // Magnetic field components (Bx, By, Bz)
};
// Struct to hold field parameters and the position
struct Field {
  union FieldComponent field; // Electric or magnetic field
  double position[3];
                        // Coordinates (x, y, z)
};
// Function to print the field at a given point
void printField(struct Field *field, const char *fieldType, const char *coordSys) {
  printf("%s field at position (%lf, %lf, %lf) in %s coordinates:\n", fieldType, field->position[0],
field->position[1], field->position[2], coordSys);
  for (int i = 0; i < 3; i++) {
     if (fieldType == "Electric") {
        printf("E%d: %lf\n", i+1, field->field.electric[i]);
     } else if (fieldType == "Magnetic") {
       printf("B%d: %lf\n", i+1, field->field.magnetic[i]);
     }
  }
}
// Function to calculate the intensity of the electromagnetic field at a given point
double calculateIntensity(struct Field *field) {
  double intensity = 0.0;
  // Calculate intensity as the magnitude of the field vector (E or B)
  for (int i = 0; i < 3; i++) {
     intensity += field->field.electric[i] * field->field.electric[i]; // For electric field
     // intensity += field->field.magnetic[i] * field->field.magnetic[i]; // Uncomment for magnetic field
  }
```

```
return sqrt(intensity);
int main() {
  // Array of field data at discrete points
  struct Field *fields = malloc(MAX POINTS * sizeof(struct Field));
  // Example: Set electric and magnetic fields at points (this is just for illustration)
  fields[0].position[0] = 1.0; fields[0].position[1] = 2.0; fields[0].position[2] = 3.0;
  fields[0].field.electric[0] = 5.0; fields[0].field.electric[1] = 3.0; fields[0].field.electric[2] = 1.0;
  // Print electric field at point (1,2,3) in Cartesian coordinates
  printField(&fields[0], "Electric", "Cartesian");
  // Calculate and print the intensity of the electric field at point (1,2,3)
  double intensity = calculateIntensity(&fields[0]);
  printf("Intensity of Electric field: %If\n", intensity);
  // Free dynamically allocated memory
  free(fields);
  return 0:
}
3. Atomic Energy Level Tracker
Description:
Track the energy levels of atoms and the transitions between them.
Specifications:
Structure: Contains atomic details (element name, energy levels, and transition probabilities).
Array: Stores energy levels for different atoms.
Union: Represents different energy states.
Strings: Represent element names.
const Pointers: Protect atomic data.
Double Pointers: Allocate memory for dynamically adding new elements.
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
// Define a constant for maximum number of energy levels
#define MAX_ENERGY_LEVELS 10
// Define a structure to store energy level information
typedef struct {
  double energy[MAX_ENERGY_LEVELS]; // Energy levels for the atom
  double transitionProbabilities[MAX_ENERGY_LEVELS - 1]; // Transition probabilities between levels
  int numLevels; // Number of energy levels for this atom
} EnergyStates;
// Define a structure to represent atomic details
typedef struct {
  const char *elementName; // Element name (string is constant)
  EnergyStates *energyStates; // Pointer to energy state data (dynamically allocated)
} Atom;
// Function to create a new atom
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Atom *createAtom(const char *elementName, int numLevels) {
  Atom *newAtom = (Atom *)malloc(sizeof(Atom));
  newAtom->elementName = elementName; // Constant string, no need to allocate memory
  // Allocate memory for energy levels
  newAtom->energyStates = (EnergyStates *)malloc(sizeof(EnergyStates));
  newAtom->energyStates->numLevels = numLevels:
  // Initialize energy levels and transition probabilities to zero
  for (int i = 0; i < numLevels; i++) {
     newAtom->energyStates->energy[i] = 0.0;
     if (i < numLevels - 1) {
       newAtom->energyStates->transitionProbabilities[i] = 0.0;
     }
  }
  return newAtom;
// Function to free memory used by an atom
void freeAtom(Atom *atom) {
  if (atom) {
     free(atom->energyStates); // Free energy states memory
     free(atom); // Free atom memory
  }
}
// Function to set energy levels for a given atom
void setEnergyLevels(Atom *atom, double *energies, int numLevels) {
  for (int i = 0; i < numLevels; i++) {
     atom->energyStates->energy[i] = energies[i];
  }
}
// Function to set transition probabilities between energy levels
void setTransitionProbabilities(Atom *atom, double *probabilities, int numLevels) {
  for (int i = 0; i < numLevels - 1; i++) {
     atom->energyStates->transitionProbabilities[i] = probabilities[i];
}
// Function to print atom details (energy levels and transition probabilities)
void printAtomDetails(Atom *atom) {
  printf("Element: %s\n", atom->elementName);
  printf("Energy Levels: \n");
  for (int i = 0; i < atom->energyStates->numLevels; i++) {
     printf("Level %d: %.2f eV\n", i + 1, atom->energyStates->energy[i]);
  }
  printf("Transition Probabilities: \n");
  for (int i = 0; i < atom->energyStates->numLevels - 1; i++) {
     printf("Transition %d -> %d: %.4\ln, i + 1, i + 2, atom->energyStates->transitionProbabilities[i]);
  }
}
```

```
// Main function to test the implementation
int main() {
  // Example of creating an atom and setting its energy levels and transition probabilities
  Atom *hydrogen = createAtom("Hydrogen", 3);
  double hydrogenEnergies[] = {13.6, 3.4, 1.5}; // Energy levels in eV
  setEnergyLevels(hydrogen, hydrogenEnergies, 3):
  double hydrogenTransitions[] = {0.98, 0.95}; // Transition probabilities
  setTransitionProbabilities(hydrogen, hydrogenTransitions, 3);
  // Print details of the hydrogen atom
  printAtomDetails(hydrogen);
  // Free memory
  freeAtom(hydrogen);
  return 0;
}
4. Quantum State Representation System
Description:
Develop a program to represent quantum states and their evolution over time.
Specifications:
Structure: Holds state properties (wavefunction amplitude, phase, and energy).
Array: Represents the wavefunction across multiple points.
Union: Stores amplitude or phase information.
Strings: Describe state labels (e.g., "ground state," "excited state").
const Pointers: Protect state properties.
Double Pointers: Manage quantum states dynamically.
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
#include <string.h>
// Define constants for quantum states
#define NUM POINTS 10 // Number of points for wavefunction representation
// Union to store amplitude or phase
union QuantumInfo {
  double amplitude; // Wavefunction amplitude
  double phase; // Phase of the wavefunction
};
// Structure to represent a quantum state
struct QuantumState {
  union QuantumInfo *wavefunction; // Pointer to an array of quantum information (amplitude or phase)
  double energy:
                            // Energy of the state
  const char *stateLabel;
                               // Label describing the quantum state
};
// Function to create a quantum state
struct QuantumState *createQuantumState(const char *label, double energy, double *amplitudeData,
double *phaseData) {
```

```
struct QuantumState *state = (struct QuantumState *)malloc(sizeof(struct QuantumState));
  state->stateLabel = label;
  state->energy = energy;
  // Dynamically allocate memory for wavefunction data (amplitude or phase)
  state->wavefunction = (union QuantumInfo *)malloc(NUM POINTS * sizeof(union QuantumInfo));
  for (int i = 0; i < NUM POINTS; i++) {
     // Assign amplitude and phase to the wavefunction based on data passed in
     state->wavefunction[i].amplitude = amplitudeData[i];
     state->wavefunction[i].phase = phaseData[i];
  }
  return state;
// Function to update quantum state dynamically (e.g., time evolution)
void evolveQuantumState(struct QuantumState *state, double time) {
  // For simplicity, we simulate a basic time evolution where phase shifts with time
  for (int i = 0; i < NUM POINTS; i++) {
     state->wavefunction[i].phase += 0.1 * time; // Update phase (just a simple model)
  }
}
// Function to print the quantum state
void printQuantumState(struct QuantumState *state) {
  printf("Quantum State: %s\n", state->stateLabel);
  printf("Energy: %.2f\n", state->energy);
  printf("Wavefunction:\n");
  for (int i = 0: i < NUM_{POINTS}: i++) {
     printf("Point %d: Amplitude = %.2f, Phase = %.2f\n", i, state->wavefunction[i].amplitude,
state->wavefunction[i].phase);
}
int main() {
  // Sample data for quantum states
  double amplitudeData[NUM_POINTS] = {0.5, 0.7, 1.0, 0.8, 0.6, 0.3, 0.9, 0.4, 0.2, 0.1};
  double phaseData[NUM_POINTS] = {0.0, 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 4.5};
  // Create a quantum state for the "ground state"
  struct QuantumState *groundState = createQuantumState("Ground State", 0.0, amplitudeData,
phaseData);
  // Print the initial state
  printQuantumState(groundState);
  // Simulate the evolution of the quantum state over time (e.g., t = 2.0)
  double time = 2.0;
  evolveQuantumState(groundState, time);
  // Print the evolved state
  printf("\nAfter %f units of time:\n", time);
  printQuantumState(groundState);
```

```
// Free allocated memory
  free(groundState->wavefunction);
  free(groundState);
  return 0;
}
5. Optics Simulation Tool
Description:
Simulate light rays passing through different optical elements.
Specifications:
Structure: Represents optical properties (refractive index, focal length).
Array: Stores light ray paths.
Union: Handles lens or mirror parameters.
Strings: Represent optical element types.
const Pointers: Protect optical properties.
Double Pointers: Manage arrays of optical elements dynamically.
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#define MAX_RAYS 100
// Optical Element Type: Lens or Mirror
typedef enum {
  LENS,
  MIRROR
} OpticalElementType;
// Structure to represent an optical element
typedef struct {
  OpticalElementType type;
  double focal length; // Focal length of lens or mirror
  double refractive index; // Refractive index for lenses
  char name[50]: // Name or type of the optical element (for display purposes)
} OpticalElement;
// Structure to represent a light ray
typedef struct {
  double x; // Position along the x-axis
  double y; // Position along the y-axis
  double angle; // Angle of the ray in degrees
} LightRay;
// Array of light rays
LightRay lightRays[MAX_RAYS];
// Function to simulate light ray passing through optical elements
void simulateLightRays(OpticalElement* element, LightRay* rays, int numRays) {
  for (int i = 0; i < numRays; i++) {
     // Simple simulation logic based on type of optical element (Lens or Mirror)
     if (element->type == LENS) {
       // Ray passing through a lens
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printf("Ray %d passing through Lens %s with focal length %.2f\n", i+1, element->name,
element->focal length);
       // Modify ray position and angle based on lens properties (simplified)
       rays[i].angle = rays[i].angle - (1 / element->focal_length);
     } else if (element->type == MIRROR) {
       // Ray reflecting from a mirror
       printf("Ray %d reflecting from Mirror %s with focal length %.2f\n", i+1, element->name,
element->focal length);
       // Modify ray angle based on mirror properties (simplified)
       rays[i].angle = -rays[i].angle;
     }
     // Output the updated ray angle
     printf("Updated ray angle: %.2f degrees\n", rays[i].angle);
  }
// Function to dynamically allocate memory for optical elements
OpticalElement* createOpticalElement(OpticalElementType type, double focalLength, double
refractiveIndex, const char* name) {
  OpticalElement* newElement = (OpticalElement*)malloc(sizeof(OpticalElement));
  newElement->type = type;
  newElement->focal_length = focalLength;
  newElement->refractive index = refractiveIndex;
  strncpy(newElement->name, name, sizeof(newElement->name) - 1);
  newElement->name[sizeof(newElement->name) - 1] = '\0'; // Ensure null-termination
  return newElement;
}
// Function to simulate the entire optics system
void simulateOpticsSystem(OpticalElement** elements, int numElements) {
  for (int i = 0; i < numElements; i++) {
     simulateLightRays(elements[i], lightRays, MAX_RAYS);
  }
}
int main() {
  // Create some optical elements (Lens and Mirror)
  OpticalElement* lens1 = createOpticalElement(LENS, 50.0, 1.5, "Convex Lens");
  OpticalElement* mirror1 = createOpticalElement(MIRROR, 100.0, 1.0, "Concave Mirror");
  // Array of pointers to optical elements
  OpticalElement* elements[] = {lens1, mirror1};
  // Initialize some light rays
  for (int i = 0; i < MAX RAYS; i++) {
     lightRays[i].x = 0.0;
     lightRays[i].y = 0.0;
     lightRays[i].angle = i * 1.0; // Rays with different angles
  }
  // Simulate the optics system
  simulateOpticsSystem(elements, 2);
  // Free dynamically allocated memory
  free(lens1);
```

```
free(mirror1);
  return 0;
}
6. Thermodynamics State Calculator
Description:
Calculate thermodynamic states of a system based on input parameters like pressure, volume, and
temperature.
Specifications:
Structure: Represents thermodynamic properties (P, V, T, and entropy).
Array: Stores states over a range of conditions.
Union: Handles dependent properties like energy or entropy.
Strings: Represent state descriptions.
const Pointers: Protect thermodynamic data.
Double Pointers: Allocate state data dynamically for simulation.
#include <stdio.h>
#include <stdlib.h>
// Define a structure for storing thermodynamic properties
typedef struct {
  double pressure; // Pressure (P)
  double volume;
                    // Volume (V)
  double temperature; // Temperature (T)
  double entropy; // Entropy (S), dependent property
  union {
     double energy; // Internal energy (U), dependent property
     double enthalpy: // Enthalpy (H), dependent property
  };
} ThermoState;
// Function to calculate entropy based on P, V, T (simplified model)
double calculate_entropy(double P, double V, double T) {
  return (P * V) / T; // Example relationship (not physically accurate, for demonstration)
}
// Function to calculate internal energy based on P, V, T (simplified model)
double calculate_energy(double P, double V, double T) {
  return P * V / T; // Example relationship (not physically accurate, for demonstration)
}
// Function to create a state dynamically
ThermoState* create_state(double P, double V, double T) {
  ThermoState* state = (ThermoState*)malloc(sizeof(ThermoState));
  if (state == NULL) {
     printf("Memory allocation failed\n");
     exit(1);
  }
  state->pressure = P:
  state->volume = V;
  state->temperature = T;
  state->entropy = calculate_entropy(P, V, T);
  state->energy = calculate_energy(P, V, T);
  return state:
```

```
}
// Function to free dynamically allocated state
void free_state(ThermoState* state) {
  free(state);
}
// Function to print thermodynamic state
void print_state(ThermoState* state) {
  printf("Pressure: %.2f Pa\n", state->pressure);
  printf("Volume: %.2f m^3\n", state->volume);
  printf("Temperature: %.2f K\n", state->temperature);
  printf("Entropy: %.2f J/K\n", state->entropy);
  printf("Energy: %.2f J\n", state->energy);
}
int main() {
  // Example of dynamically allocating memory for a state
  ThermoState* state1 = create_state(101325, 0.1, 300.0); // P = 101325 Pa, V = 0.1 m^3, T = 300K
  // Print the state information
  print_state(state1);
  // Free allocated memory
  free_state(state1);
  return 0;
7. Nuclear Reaction Tracker
Description:
Track the parameters of nuclear reactions like fission and fusion processes.
Specifications:
Structure: Represents reaction details (reactants, products, energy released).
Array: Holds data for multiple reactions.
Union: Represents either energy release or product details.
Strings: Represent reactant and product names.
const Pointers: Protect reaction details.
Double Pointers: Dynamically allocate memory for reaction data.
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
// Define a Union to store either energy released or product details
union ReactionDetails {
  double energyReleased; // Energy released in the reaction
  char *productDetails; // Product details (name of the product)
};
// Define a Structure for a nuclear reaction
struct NuclearReaction {
  char *reactant1;
                          // Name of the first reactant
  char *reactant2;
                          // Name of the second reactant (for fusion)
  union ReactionDetails details; // Details (either energy or products)
```

```
int isFission;
                       // Flag to differentiate between fission (0) or fusion (1)
};
// Function to create a new reaction dynamically
struct NuclearReaction* createReaction(char *reactant1, char *reactant2, double energyReleased, int
isFission) {
  struct NuclearReaction *reaction = (struct NuclearReaction *) malloc(sizeof(struct NuclearReaction));
  reaction->reactant1 = (char *) malloc(strlen(reactant1) + 1);
  strcpy(reaction->reactant1, reactant1);
  if (isFission == 0) {
     reaction->reactant2 = NULL; // Fission doesn't require a second reactant
  } else {
     reaction->reactant2 = (char *) malloc(strlen(reactant2) + 1);
     strcpy(reaction->reactant2, reactant2);
  }
  // Set energy released or product details
  if (isFission == 0) {
     reaction->details.energyReleased = energyReleased;
  } else {
     reaction->details.productDetails = (char *) malloc(100 * sizeof(char)); // Assume product name is
<100 chars
     strcpy(reaction->details.productDetails, "Helium, Neutron, Energy");
  }
  reaction->isFission = isFission;
  return reaction;
}
// Function to print the details of a reaction
void printReactionDetails(struct NuclearReaction *reaction) {
  if (reaction->isFission == 0) {
     printf("Fission Reaction:\n");
     printf("Reactant: %s + %s\n", reaction->reactant1, reaction->reactant2? reaction->reactant2:
"N/A");
     printf("Energy Released: %.2lf MeV\n", reaction->details.energyReleased);
  } else {
     printf("Fusion Reaction:\n");
     printf("Reactant: %s + %s\n", reaction->reactant1, reaction->reactant2);
     printf("Products: %s\n", reaction->details.productDetails);
  }
}
// Function to free the memory allocated for a reaction
void freeReaction(struct NuclearReaction *reaction) {
  free(reaction->reactant1);
  if (reaction->reactant2) {
     free(reaction->reactant2);
  if (reaction->isFission == 1) {
     free(reaction->details.productDetails);
```

```
free(reaction);
int main() {
  // Create a few example nuclear reactions
  struct NuclearReaction *fissionReaction = createReaction("Uranium-235", "Neutron", 200.0, 0);
  struct NuclearReaction *fusionReaction = createReaction("Deuterium", "Tritium", 0.0, 1);
  // Print the reaction details
  printReactionDetails(fissionReaction);
  printf("\n");
  printReactionDetails(fusionReaction);
  // Free allocated memory
  freeReaction(fissionReaction):
  freeReaction(fusionReaction);
  return 0;
}
8. Gravitational Field Simulation
Description:
Simulate the gravitational field of massive objects in a system.
Specifications:
Structure: Contains object properties (mass, position, field strength).
Array: Stores field values at different points.
Union: Handles either mass or field strength as parameters.
Strings: Represent object labels (e.g., "Planet A," "Star B").
const Pointers: Protect object properties.
Double Pointers: Dynamically allocate memory for gravitational field data.
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
// Constants
#define G 6.67430e-11 // Gravitational constant (m^3 kg^-1 s^-2)
// Structure to store object properties (mass, position, field strength)
typedef struct {
                     // Name/label of the object (e.g., "Planet A")
  char label[50];
                     // Mass of the object in kilograms
  double mass:
  double position[3]; // Position of the object in space (x, y, z coordinates)
  double field strength[3]; // Gravitational field strength at the object's position (x, y, z components)
} GravitationalObject;
// Union to handle either mass or field strength as parameters
typedef union {
  double mass:
  double field strength[3];
} Gravitational Parameter;
// Function to calculate gravitational field at a given point due to an object
void calculate gravitational field(GravitationalObject* obj, double* point, double* field) {
  // Vector from the object to the point
```

```
double dx = point[0] - obj->position[0];
  double dy = point[1] - obj->position[1];
  double dz = point[2] - obj->position[2];
  // Calculate the distance between the object and the point
  double r = sqrt(dx * dx + dy * dy + dz * dz);
  // Gravitational field strength calculation
  if (r != 0) {
     double field magnitude = G * obj->mass / (r * r);
     // Field components (directional)
     field[0] = field magnitude * dx / r;
     field[1] = field magnitude * dy / r;
     field[2] = field magnitude * dz / r;
  } else {
     field[0] = field[1] = field[2] = 0;
}
// Function to simulate and display the gravitational field at various points in space
void simulate gravitational field(GravitationalObject* objects, int num objects, double** points, int
num points) {
  // Allocate memory for field values (dynamically)
  double** field values = (double**)malloc(num points * sizeof(double*));
  for (int i = 0; i < num points; i++) {
     field values[i] = (double*)malloc(3 * sizeof(double)); // 3 components for x, y, z
  }
  // For each point, calculate the gravitational field due to all objects
  for (int i = 0; i < num points; i++) {
     for (int j = 0; j < 3; j++) {
        field values[i][j] = 0; // Initialize field to zero
     }
     // Sum the contributions of all objects to the field at this point
     for (int j = 0; j < num objects; j++) {
        double field[3]:
        calculate gravitational field(&objects[i], points[i], field);
        // Accumulate the field contributions
        for (int k = 0; k < 3; k++) {
          field values[i][k] += field[k];
     }
     // Display the resulting field at the current point
     printf("Gravitational field at point (%f, %f, %f): (%f, %f, %f)\n",
           points[i][0], points[i][1], points[i][2],
          field values[i][0], field values[i][1], field values[i][2]);
  }
  // Free dynamically allocated memory
  for (int i = 0; i < num points; i++) {
     free(field values[i]);
```

```
free(field values);
int main() {
  // Define some sample objects (e.g., planets, stars)
  GravitationalObject objects[2] = {
     {"Planet A", 5.972e24, {0, 0, 0}, {0, 0, 0}}, // Earth-like object
     {"Star B", 1.989e30, {1000, 1000, 1000}, {0, 0, 0}} // Sun-like object at (1000, 1000, 1000)
  };
  // Define some points in space where we want to calculate the gravitational field
  double points[2][3] = \{
     {10, 10, 10}, // Point 1 (x, y, z)
     {200, 200, 200} // Point 2 (x, y, z)
  };
  // Call the simulation function
  simulate gravitational field(objects, 2, (double**)points, 2);
  return 0;
}
9. Wave Interference Analyzer
Description:
Analyze interference patterns produced by waves from multiple sources.
Specifications:
Structure: Represents wave properties (amplitude, wavelength, and phase).
Array: Stores wave interference data at discrete points.
Union: Handles either amplitude or phase information.
Strings: Represent wave source labels.
const Pointers: Protect wave properties.
Double Pointers: Manage dynamic allocation of wave sources.
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
#define MAX SOURCES 10 // You can define a limit for the number of sources
// Structure to hold wave properties (amplitude, wavelength, phase)
typedef struct {
  double amplitude;
  double wavelength;
  double phase;
} WaveProperties;
// Union to store either amplitude or phase information
typedef union {
  double amplitude;
  double phase;
} WaveUnion;
// Structure to represent a single wave source
typedef struct {
```

```
char* label;
                      // Label for the wave source
  WaveProperties properties; // Wave properties: amplitude, wavelength, phase
  WaveUnion wave info; // Union for amplitude or phase information
} WaveSource;
// Double pointer to manage dynamic allocation for an array of wave sources
typedef struct {
  WaveSource** sources; // Array of wave sources
                    // Number of sources
  int num sources;
} WaveSourceManager;
// Function to initialize the WaveSourceManager
WaveSourceManager* createWaveSourceManager(int num_sources) {
  WaveSourceManager* manager = (WaveSourceManager*)malloc(sizeof(WaveSourceManager));
  manager->sources = (WaveSource**)malloc(num sources * sizeof(WaveSource*));
  manager->num_sources = num_sources;
  return manager;
}
// Function to initialize a single wave source
void initWaveSource(WaveSource* source, const char* label, double amplitude, double wavelength,
double phase) {
  source->label = label;
  source->properties.amplitude = amplitude;
  source->properties.wavelength = wavelength;
  source->properties.phase = phase;
}
// Function to calculate interference pattern at a point (for simplicity)
double calculateInterferencePattern(WaveSourceManager* manager, double position) {
  double total amplitude = 0.0;
  double total phase = 0.0;
  for (int i = 0; i < manager->num sources; i++) {
    WaveSource* source = manager->sources[i];
    // Calculate interference for each wave based on its properties
    total_amplitude += source->properties.amplitude * cos(source->properties.phase + (2 * M PI *
position / source->properties.wavelength));
    total phase += source->properties.phase;
  }
  // In this simple case, we return the total amplitude
  return total amplitude;
}
// Clean up memory for wave sources
void freeWaveSourceManager(WaveSourceManager* manager) {
  free(manager->sources);
  free(manager);
}
int main() {
  // Create a manager for 2 wave sources
  WaveSourceManager* manager = createWaveSourceManager(2);
```

```
// Initialize wave sources
  WaveSource wave1:
  initWaveSource(&wave1, "Wave1", 1.0, 2.0, 0.0);
  WaveSource wave2;
  initWaveSource(&wave2, "Wave2", 0.5, 3.0, M PI / 2);
  // Assign sources to the manager
  manager->sources[0] = &wave1;
  manager->sources[1] = &wave2;
  // Analyze interference pattern at position x = 5.0
  double interference = calculateInterferencePattern(manager, 5.0);
  printf("Interference pattern at x = 5.0: %f\n", interference);
  // Clean up
  freeWaveSourceManager(manager);
  return 0;
10. Magnetic Material Property Database
Description:
Create a database to store and retrieve properties of magnetic materials.
Specifications:
Structure: Represents material properties (permeability, saturation).
Array: Stores data for multiple materials.
Union: Handles temperature-dependent properties.
Strings: Represent material names.
const Pointers: Protect material data.
Double Pointers: Allocate material records dynamically.
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
// Union to handle temperature-dependent properties
union TempDependentProperties {
  double curieTemperature; // Curie temperature (for ferromagnetic materials)
  double temperatureSaturation; // Temperature-dependent saturation value
};
// Structure to represent a magnetic material's properties
typedef struct {
  char name[50];
                               // Material name (e.g., "Iron")
  double permeability;
                                // Permeability ( )
  double saturation;
                                // Saturation (B s)
  union TempDependentProperties tempProperties; // Union for temperature-related properties
} Material:
// Function to create a new material
Material* createMaterial(const char* name, double permeability, double saturation, double curieTemp) {
  Material* newMaterial = (Material*)malloc(sizeof(Material)); // Dynamically allocate memory for a new
material
```

```
if (newMaterial == NULL) {
     printf("Memory allocation failed.\n");
     exit(1);
  strncpy(newMaterial->name, name, sizeof(newMaterial->name)); // Copy the name
  newMaterial->permeability = permeability;
                                                           // Set permeability
  newMaterial->saturation = saturation:
                                                         // Set saturation
  newMaterial->tempProperties.curieTemperature = curieTemp; // Set Curie temperature
  return newMaterial:
}
// Function to display material information
void displayMaterial(const Material* material) {
  printf("Material: %s\n", material->name);
  printf("Permeability: %.2lf\n", material->permeability);
  printf("Saturation: %.2lf\n", material->saturation);
  printf("Curie Temperature: %.2lf K\n", material->tempProperties.curieTemperature);
}
// Function to free dynamically allocated memory for materials
void freeMaterial(Material* material) {
  free(material); // Free the memory allocated for the material
}
int main() {
  // Dynamically allocate memory for an array of materials (using double pointer)
  Material** materials = (Material**)malloc(3 * sizeof(Material*)); // Example with 3 materials
  if (materials == NULL) {
     printf("Memory allocation failed.\n");
     return 1;
  }
  // Create materials and store in the array
  materials[0] = createMaterial("Iron", 1.26e-6, 2.2, 1043.0);
  materials[1] = createMaterial("Nickel", 6.0e-6, 0.6, 358.0);
  materials[2] = createMaterial("Cobalt", 1.2e-5, 1.4, 1388.0);
  // Display all materials
  for (int i = 0; i < 3; i++) {
     displayMaterial(materials[i]);
     printf("\n");
  }
  // Free the dynamically allocated memory for each material
  for (int i = 0; i < 3; i++) {
     freeMaterial(materials[i]);
  }
  // Free the array of pointers
  free(materials);
  return 0;
}
```

```
Description:
Simulate the behavior of plasma under various conditions.
Specifications:
Structure: Represents plasma parameters (density, temperature, and electric field).
Array: Stores simulation results.
Union: Handles either density or temperature data.
Strings: Represent plasma types.
const Pointers: Protect plasma parameters.
Double Pointers: Manage dynamic allocation for simulation data.
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#define MAX PLASMA TYPES 5
// Define a union for storing either density or temperature data.
typedef union {
  double density;
  double temperature;
} PlasmaData;
// Structure to represent plasma parameters.
typedef struct {
                    // Plasma type (e.g., "Ionized Gas", "Hot Plasma")
  char type[50];
  double electricField; // Electric field in the plasma
  PlasmaData data; // Either density or temperature
} Plasma;
// Structure to hold simulation results.
typedef struct {
  Plasma* plasmas;
                      // Array of plasma objects
  size t numPlasmas; // Number of plasmas in the simulation
} PlasmaSimulation;
// Function to initialize the plasma array.
void initPlasmaSimulation(PlasmaSimulation* sim, size_t numPlasmas) {
  sim->plasmas = (Plasma*)malloc(numPlasmas * sizeof(Plasma));
  sim->numPlasmas = numPlasmas:
  if (!sim->plasmas) {
     printf("Memory allocation failed!\n");
     exit(1);
  }
}
// Function to set plasma data.
void setPlasmaData(Plasma* plasma, const char* type, double electricField, double value, int isDensity) {
  strncpy(plasma->type, type, sizeof(plasma->type) - 1);
  plasma->electricField = electricField;
  if (isDensity) {
     plasma->data.density = value;
  } else {
     plasma->data.temperature = value;
}
```

```
// Function to display the plasma simulation results.
void displaySimulationResults(const PlasmaSimulation* sim) {
  for (size_t i = 0; i < sim->numPlasmas; ++i) {
     printf("Plasma Type: %s\n", sim->plasmas[i].type);
     printf("Electric Field: %.2f\n", sim->plasmas[i].electricField);
     if (sim->plasmas[i].data.density) {
       printf("Density: %.2f\n", sim->plasmas[i].data.density);
     } else {
       printf("Temperature: %.2f\n", sim->plasmas[i].data.temperature);
     printf("-----\n");
}
// Function to free the memory used by the simulation.
void freePlasmaSimulation(PlasmaSimulation* sim) {
  free(sim->plasmas);
}
int main() {
  PlasmaSimulation sim;
  size t numPlasmas = 3;
  // Initialize simulation with 3 plasma types.
  initPlasmaSimulation(&sim, numPlasmas);
  // Set data for each plasma (density for first and temperature for others).
  setPlasmaData(&sim.plasmas[0], "Ionized Gas", 5.0, 1.2, 1); // Set density
  setPlasmaData(&sim.plasmas[1], "Hot Plasma", 10.0, 15000.0, 0); // Set temperature
  setPlasmaData(&sim.plasmas[2], "Cold Plasma", 8.0, 1000.0, 0); // Set temperature
  // Display simulation results.
  displaySimulationResults(&sim);
  // Free the dynamically allocated memory.
  freePlasmaSimulation(&sim):
  return 0;
}
12. Kinematics Equation Solver
Description:
Solve complex kinematics problems for objects in motion.
Specifications:
Structure: Represents object properties (initial velocity, acceleration, displacement).
Array: Stores time-dependent motion data.
Union: Handles either velocity or displacement equations.
Strings: Represent motion descriptions.
const Pointers: Protect object properties.
Double Pointers: Dynamically allocate memory for motion data.
#include <stdio.h>
#include <stdlib.h>
```

```
// Structure to represent object properties (initial velocity, acceleration, displacement)
typedef struct {
  double initial velocity; // Initial velocity (m/s)
  double acceleration: // Acceleration (m/s^2)
  double displacement;
                          // Displacement (m)
} ObjectProperties;
// Union to handle either velocity or displacement equations
typedef union {
  double velocity;
                        // Final velocity (m/s)
  double displacement; // Displacement (m)
} MotionData;
// Structure to represent kinematics equations (using both structure and union)
typedef struct {
  ObjectProperties obj_props; // Object properties
  double time:
                         // Time (s)
  MotionData motion;
                             // Union for either velocity or displacement
} KinematicsSolver;
// Function to solve for final velocity (v = u + at)
double solve_velocity(KinematicsSolver *ks) {
  return ks->obj props.initial velocity + ks->obj props.acceleration * ks->time;
}
// Function to solve for displacement (s = ut + 0.5 * a * t^2)
double solve displacement(KinematicsSolver *ks) {
  return ks->obj_props.initial_velocity * ks->time + 0.5 * ks->obj_props.acceleration * ks->time *
ks->time:
}
// Function to display motion data
void display motion data(KinematicsSolver *ks) {
  printf("Initial Velocity (u): %.2f m/s\n", ks->obj_props.initial_velocity);
  printf("Acceleration (a): %.2f m/s^2\n", ks->obj props.acceleration);
  printf("Time (t): %.2f s\n", ks->time);
}
int main() {
  // Dynamically allocate memory for motion data (array for storing times and velocities)
  int num samples = 5;
  double *velocities = (double*)malloc(num_samples * sizeof(double));
  if (velocities == NULL) {
     printf("Memory allocation failed\n");
     return 1;
  }
  // Sample kinematic values for the object
  KinematicsSolver ks = {
     .obj props = \{50.0, 9.8, 0.0\}, // Initial velocity = 50 \text{ m/s}, acceleration = 9.8 \text{ m/s}^2, displacement = 0
     .time = 10.0
  };
  // Display initial object properties
  display motion data(&ks);
```

```
// Calculate final velocity and displacement
  ks.motion.velocity = solve velocity(&ks);
  ks.motion.displacement = solve_displacement(&ks);
  printf("\nFinal velocity (v): %.2f m/s\n", ks.motion.velocity);
  printf("Displacement (s): %.2f m\n", ks.motion.displacement);
  // Store velocity data at different time intervals
  for (int i = 0; i < num samples; <math>i++) {
     ks.time = (i + 1) * 2.0; // Time steps at 2s intervals
     velocities[i] = solve_velocity(&ks);
  }
  // Display velocities at different time intervals
  printf("\nVelocities at different time intervals:\n");
  for (int i = 0; i < num\_samples; i++) {
     printf("t = \%.2f s, v = \%.2f m/s\n", (i + 1) * 2.0, velocities[i]);
  }
  // Free dynamically allocated memory
  free(velocities);
  return 0;
}
13. Spectral Line Database
Description:
Develop a database to store and analyze spectral lines of elements.
Specifications:
Structure: Represents line properties (wavelength, intensity, and element).
Array: Stores spectral line data.
Union: Handles either intensity or wavelength information.
Strings: Represent element names.
const Pointers: Protect spectral line data.
Double Pointers: Allocate spectral line records dynamically.
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
// Define the maximum size for element names
#define MAX ELEMENT NAME 50
// Union to hold either wavelength or intensity information
union SpectralData {
  double wavelength;
  double intensity;
};
// Structure to store spectral line data
struct SpectralLine {
  char element[MAX_ELEMENT_NAME]; // Element name (e.g., "Hydrogen")
                                 // Wavelength or Intensity
  union SpectralData data;
  int isWavelength;
                              // Flag to check if the data is wavelength (1) or intensity (0)
```

```
};
// Function to create a new spectral line record dynamically
void createSpectralLine(struct SpectralLine **line, const char *element, double value, int isWavelength) {
  *line = (struct SpectralLine *)malloc(sizeof(struct SpectralLine)); // Allocate memory for a new record
  if (*line == NULL) {
     printf("Memory allocation failed!\n");
     exit(1);
  }
  // Set the element name
  strncpy((*line)->element, element, MAX_ELEMENT_NAME - 1);
  (*line)->element[MAX_ELEMENT_NAME - 1] = '\0'; // Ensure null termination
  // Store either wavelength or intensity based on the flag
  if (isWavelength) {
     (*line)->data.wavelength = value;
     (*line)->isWavelength = 1;
  } else {
     (*line)->data.intensity = value;
     (*line)->isWavelength = 0;
  }
}
// Function to display a spectral line record
void displaySpectralLine(const struct SpectralLine *line) {
  if (line->isWavelength) {
     printf("Element: %s, Wavelength: %.2f nm\n", line->element, line->data.wavelength);
  } else {
     printf("Element: %s, Intensity: %.2f\n", line->element, line->data.intensity);
  }
}
// Function to free allocated memory for spectral line records
void freeSpectralLine(struct SpectralLine *line) {
  free(line);
}
int main() {
  struct SpectralLine *line1, *line2;
  // Create a spectral line for Hydrogen with wavelength
  createSpectralLine(&line1, "Hydrogen", 656.3, 1); // Wavelength in nm
  // Create a spectral line for Oxygen with intensity
  createSpectralLine(&line2, "Oxygen", 12345.67, 0); // Intensity in arbitrary units
  // Display the spectral lines
  displaySpectralLine(line1);
  displaySpectralLine(line2);
  // Free allocated memory
  freeSpectralLine(line1);
  freeSpectralLine(line2);
```

return 0;

```
14. Projectile Motion Simulator
Description:
Simulate and analyze projectile motion under varying conditions.
Specifications:
Structure: Stores projectile properties (mass, velocity, and angle).
Array: Stores motion trajectory data.
Union: Handles either velocity or displacement parameters.
Strings: Represent trajectory descriptions.
const Pointers: Protect projectile properties.
Double Pointers: Manage trajectory records dynamically.
#include <stdio.h>
#include <math.h>
#include <stdlib.h>
#define GRAVITY 9.81 // m/s^2
// Structure to hold projectile properties
typedef struct {
  double mass; // Mass of the projectile (kg)
  double velocity; // Initial velocity (m/s)
  double angle; // Launch angle (degrees)
} Projectile;
// Union to handle velocity or displacement data
typedef union {
  double velocity;
  double displacement;
} TrajectoryData;
// Structure to store trajectory data
typedef struct {
  double time; // Time at this point in the trajectory double height; // Height of the projectile at this time
  double horizontal; // Horizontal displacement at this time
} TrajectoryPoint;
// Function to calculate the trajectory
TrajectoryPoint *calculateTrajectory(Projectile *p, int *pointsCount) {
  // Calculate the initial velocity components
  double angle_rad = p->angle * (M_PI / 180.0);
  double v x = p->velocity * cos(angle rad);
  double v_y = p->velocity * sin(angle_rad);
  // Time of flight (T)
  double flightTime = (2 * v_y) / GRAVITY;
  // Max number of points
  int maxPoints = 100;
  *pointsCount = maxPoints;
  // Dynamically allocate memory for trajectory points
```

}

```
TrajectoryPoint *trajectory = (TrajectoryPoint *)malloc(maxPoints * sizeof(TrajectoryPoint));
  // Calculate the trajectory points
  for (int i = 0; i < maxPoints; i++) {
     double t = flightTime * i / (maxPoints - 1); // Normalize time
     // Calculate position at time t
     trajectory[i].time = t;
     trajectory[i].horizontal = v_x * t;
     trajectory[i].height = v_y * t - 0.5 * GRAVITY * t * t;
     // If the projectile hits the ground, stop calculating
     if (trajectory[i].height < 0) {
        *pointsCount = i;
        break:
     }
  }
  return trajectory;
// Function to print trajectory points
void printTrajectory(TrajectoryPoint *trajectory, int pointsCount) {
  printf("Time (s)\tHorizontal Displacement (m)\tHeight (m)\n");
  for (int i = 0; i < pointsCount; i++) {
     printf("%.2f\t\t%.2f\n", trajectory[i].time, trajectory[i].horizontal, trajectory[i].height);
  }
}
int main() {
  // Initialize projectile properties
  Projectile p;
  p.mass = 1.0; // Mass (kg)
  p.velocity = 50.0; // Initial velocity (m/s)
  p.angle = 45.0; // Launch angle (degrees)
  // Calculate the trajectory
  int pointsCount = 0;
  TrajectoryPoint *trajectory = calculateTrajectory(&p, &pointsCount);
  // Print the results
  printTrajectory(trajectory, pointsCount);
  // Free dynamically allocated memory
  free(trajectory);
  return 0;
}
15. Material Stress-Strain Analyzer
Description:
Analyze the stress-strain behavior of materials under different loads.
Specifications:
Structure: Represents material properties (stress, strain, modulus).
```

```
Array: Stores stress-strain data.
Union: Handles dependent properties like yield stress or elastic modulus.
Strings: Represent material names.
const Pointers: Protect material properties.
Double Pointers: Allocate stress-strain data dynamically.
#include <stdio.h>
#include <stdlib.h>
#include <string.h>
#define MAX MATERIALS 10
// Union to store dependent properties like yield stress or elastic modulus
union MaterialProperties {
  double yieldStress;
  double elasticModulus;
};
// Structure to represent the material's stress and strain data
typedef struct {
  char name[50]:
                          // Material name (String)
                          // Pointer to stress data (dynamically allocated)
  double* stress;
                         // Pointer to strain data (dynamically allocated)
  double* strain:
                         // Number of data points for stress-strain data
  int dataPoints:
  union Material Properties properties; // Union to hold dependent material properties
} Material;
// Function to allocate and initialize a material's stress-strain data
void initMaterial(Material* material, const char* name, int dataPoints) {
  strncpy(material->name, name, sizeof(material->name) - 1);
  material->name[sizeof(material->name) - 1] = '\0'; // Ensure null-terminated string
  material->dataPoints = dataPoints;
  // Dynamically allocate memory for stress and strain data
  material->stress = (double*)malloc(dataPoints * sizeof(double));
  material->strain = (double*)malloc(dataPoints * sizeof(double));
  if (!material->stress || !material->strain) {
     printf("Memory allocation failed for stress or strain data.\n");
     exit(1);
}
// Function to deallocate material data
void freeMaterial(Material* material) {
  free(material->stress);
  free(material->strain);
}
// Function to display stress-strain data
void displayStressStrainData(Material* material) {
  printf("Material: %s\n", material->name);
  printf("Stress (Pa) | Strain\n");
  for (int i = 0; i < material->dataPoints; i++) {
     printf("%.2f
                    | %.5f\n", material->stress[i], material->strain[i]);
```

```
}
// Function to set yield stress or elastic modulus
void setMaterialProperties(Material* material, double value, int isElasticModulus) {
  if (isElasticModulus) {
     material->properties.elasticModulus = value;
  } else {
     material->properties.yieldStress = value;
}
// Function to analyze stress-strain behavior (basic analysis: display yield stress or modulus)
void analyzeStressStrain(Material* material) {
  printf("Analyzing stress-strain for material: %s\n", material->name);
  printf("Elastic Modulus: %.2f MPa\n", material->properties.elasticModulus);
  printf("Yield Stress: %.2f MPa\n", material->properties.yieldStress);
}
int main() {
  Material materials[MAX_MATERIALS];
  // Example material 1
  initMaterial(&materials[0], "Steel", 5);
  materials[0].stress[0] = 100.0; materials[0].strain[0] = 0.002;
  materials[0].stress[1] = 200.0; materials[0].strain[1] = 0.004;
  materials[0].stress[2] = 300.0; materials[0].strain[2] = 0.006;
  materials[0].stress[3] = 400.0; materials[0].strain[3] = 0.008;
  materials[0].stress[4] = 500.0; materials[0].strain[4] = 0.01;
  setMaterialProperties(&materials[0], 210000, 1); // Elastic Modulus (MPa)
  setMaterialProperties(&materials[0], 250, 0); // Yield Stress (MPa)
  displayStressStrainData(&materials[0]);
  analyzeStressStrain(&materials[0]);
  // Example material 2
  initMaterial(&materials[1], "Aluminum", 5);
  materials[1].stress[0] = 50.0; materials[1].strain[0] = 0.001;
  materials[1].stress[1] = 100.0; materials[1].strain[1] = 0.002;
  materials[1].stress[2] = 150.0; materials[1].strain[2] = 0.003;
  materials[1].stress[3] = 200.0; materials[1].strain[3] = 0.004;
  materials[1].stress[4] = 250.0; materials[1].strain[4] = 0.005;
  setMaterialProperties(&materials[1], 70000, 1); // Elastic Modulus (MPa)
  setMaterialProperties(&materials[1], 150, 0); // Yield Stress (MPa)
  displayStressStrainData(&materials[1]);
  analyzeStressStrain(&materials[1]);
  // Clean up dynamic memory
  freeMaterial(&materials[0]);
  freeMaterial(&materials[1]);
  return 0;
```

}		