

Chem 30324, Spring 2019, Homework 10

Due April 17, 2019

The two-state system.

Consider a closed system containing N objects, each of which can be in one of two energy states, of energy either 0 or ε . The total internal energy U of the box is the sum of the energies of the individual objects.

1. Write down all the possible microstates for a box in which $N = 4$ and the internal energy $U = 2\varepsilon$.
2. What does the postulate of *equal a priori probabilities* say about the relative likelihood of occurrence of any one of these microstates?
3. What is the entropy of the box? (Thank you, Ludwig Boltzmann.)
4. Suppose two identical such boxes are brought into thermal contact and allowed to come to equilibrium. Calculate the change in internal energy ΔU and in entropy ΔS associated with this process.

The canonical ensemble.

The energy spectrum of some molecule is described by the diagram below.

 Energy States

5. Write the partition function q for the molecule at thermal equilibrium at a temperature $\beta = 1/k_B T$.
6. Plot the probability for the molecule to be in each of the three energy states vs. temperature. Be sure to indicate the probabilities in the limits of $T \rightarrow 0$ and $T \rightarrow \infty$.

7. Derive an expression for the energy U per molecule by summing over the possible microstates weighted by their probabilities. Plot the average energy vs. temperature.
8. Derive an expression for the energy U per molecule by taking the appropriate derivative of the partition function from problem 5 (*Hint: it is easier to work with the expressions in term of β than in T .*) Does your result agree with that from problem 7?
9. Derive an expression for the Helmholtz energy A per molecule from the partition function. Plot A vs. temperature, assuming $\epsilon/k_B = 300$ K.
10. Derive an expression for the entropy S per molecules and plot vs. temperature, again assuming $\epsilon/k_B = 300$ K.
11. In class we took the First Law as a postulate and demonstrated the Second Law. Look at your results for Problems 6 and 10. Can you use them to rationalize the Third Law? Explain your answer.

Thermodynamics from scratch.

Let's calculate the thermodynamic properties of an ideal gas of CO molecules at 1 bar pressure. CO has a rotational constant $B = 1.931 \text{ cm}^{-1}$ and vibrational frequency $\nu = 2156.6 \text{ cm}^{-1}$. Suppose you have a 20 dm^3 cubic bottle containing 1 mole of CO gas that you can consider to behave ideally.

12. The characteristic temperature Θ of a particular degree of freedom is the characteristic quantum of energy for the degree of freedom divided by k_B . Calculate the characteristic translational, rotational, and vibrational temperatures of CO.
13. Calculate the *translational partition function* of a CO molecule in the bottle at 298 K. What is the unit of the partition function?
14. Plot the *rotational and vibrational partition functions* of a CO molecule in the bottle from $T = 200$ to 2000 K (assume the CO remains a gas over the whole range). *Hint: Use your answer to Problem 12 to simplify calculating the rotational partition function.*

15. Plot the *total translational, rotational, and vibrational energies* of CO in the bottle from $T = 200$ to 2000 K (assume the CO remains a gas over the whole range). Which (if any) of the three types of motions dominate the total energy?

16. Plot the *total translational, rotational, and vibrational constant volume molar heat capacities* of CO in the bottle from $T = 200$ to 2000 K . Which (if any) of the three types of motions dominate the heat capacity?

17. Plot the *total translational, rotational, and vibrational Helmholtz energies* of CO in the bottle from $T = 200$ to 2000 K . Which (if any) of the three types of motions dominate the Helmholtz energy?

18. Use your formulas to calculate ΔP , ΔU , ΔA , and ΔS associated with isothermally expanding the gas from 20 dm^3 to 40 dm^3 .