# Experimental Rate Laws

Rate laws (differential rate laws) provide a mathematical description of how changes in the concentration of a substance affect the rate of a chemical reaction. Rate laws are determined experimentally and cannot be predicted by reaction stoichiometry. The order of reaction describes how much a change in the concentration of each substance affects the overall rate, and the overall order of a reaction is the sum of the orders for each substance present in the reaction. Reaction orders are typically first order, second order, or zero order, but fractional and even negative orders are possible.

### Learning Objectives

By the end of this section, you will be able to:

* Explain the form and function of a rate law
* Use rate laws to calculate reaction rates
* Use rate and concentration data to identify reaction orders and derive rate laws

As described in the previous module, the rate of a reaction is often affected by the concentrations of reactants. Rate laws (sometimes called differential rate laws) or rate equations are mathematical expressions that describe the relationship between the rate of a chemical reaction and the concentration of its reactants. As an example, consider the reaction described by the chemical equation

aA+bB⟶productsaA+bB⟶products

where a and b are stoichiometric coefficients. The rate law for this reaction is written as:

rate=k[A]m[B]nrate=k[A]m[B]n

in which [A] and [B] represent the molar concentrations of reactants, and k is the rate constant, which is specific for a particular reaction at a particular temperature. The exponents m and n are the reaction orders and are typically positive integers, though they can be fractions, negative, or zero. The rate constant k and the reaction orders m and n must be determined experimentally by observing how the rate of a reaction changes as the concentrations of the reactants are changed. The rate constant k is independent of the reactant concentrations, but it does vary with temperature.

The reaction orders in a rate law describe the mathematical dependence of the rate on reactant concentrations. Referring to the generic rate law above, the reaction is m order with respect to A and n order with respect to B. For example, if m = 1 and n = 2, the reaction is first order in A and second order in B. The overall reaction order is simply the sum of orders for each reactant. For the example rate law here, the reaction is third order overall (1 + 2 = 3). A few specific examples are shown below to further illustrate this concept.

The rate law:

rate=k[H2O2]rate=k[H2O2]

describes a reaction that is first order in hydrogen peroxide and first order overall. The rate law:

rate=k[C4H6]2rate=k[C4H6]2

describes a reaction that is second order in C4H6 and second order overall. The rate law:

rate=k[H+][OH−]rate=k[H+][OH−]

describes a reaction that is first order in H+, first order in OH−, and second order overall.

### EXAMPLE: 7.0.1 Writing Rate Laws from Reaction Orders

An experiment shows that the reaction of nitrogen dioxide with carbon monoxide:

NO2(g)+CO(g)⟶NO(g)+CO2(g)NO2(g)+CO(g)⟶NO(g)+CO2(g)

is second order in NO2 and zero order in CO at 100 °C. What is the rate law for the reaction?

#### Solution

The reaction will have the form:

rate=k[NO2]m[CO]nrate=k[NO2]m[CO]n

The reaction is second order in NO2; thus m = 2. The reaction is zero order in CO; thus n = 0. The rate law is:

rate=k[NO2]2[CO]0=k[NO2]2rate=k[NO2]2[CO]0=k[NO2]2

Remember that a number raised to the zero power is equal to 1, thus [CO]0 = 1, which is why the CO concentration term may be omitted from the rate law: the rate of reaction is solely dependent on the concentration of NO2. A later chapter section on reaction mechanisms will explain how a reactant’s concentration can have no effect on a reaction rate despite being involved in the reaction.

#### Check Your Learning

The rate law for the reaction:

H2(g)+2NO(g)⟶N2O(g)+H2O(g)H2(g)+2NO(g)⟶N2O(g)+H2O(g)

has been determined to be rate = k[NO]2[H2]. What are the orders with respect to each reactant, and what is the overall order of the reaction?

### Answer

order in NO = 2; order in H2 = 1; overall order = 3

#### Check Your Learning

In a transesterification reaction, a triglyceride reacts with an alcohol to form an ester and glycerol. Many students learn about the reaction between methanol (CH

3

OH) and ethyl acetate (CH

3

CH

2

OCOCH

3

) as a sample reaction before studying the chemical reactions that produce biodiesel:

CH3OH+CH3CH2OCOCH3⟶CH3OCOCH3+CH3CH2OHCH3OH+CH3CH2OCOCH3⟶CH3OCOCH3+CH3CH2OH

The rate law for the reaction between methanol and ethyl acetate is, under certain conditions, determined to be:

rate=k[CH3OH]rate=k[CH3OH]

What is the order of reaction with respect to methanol and ethyl acetate, and what is the overall order of reaction?

### Answer

order in CH3OH = 1; order in CH3CH2OCOCH3 = 0; overall order = 1

A common experimental approach to the determination of rate laws is the method of initial rates. This method involves measuring reaction rates for multiple experimental trials carried out using different initial reactant concentrations. Comparing the measured rates for these trials permits determination of the reaction orders and, subsequently, the rate constant, which together are used to formulate a rate law. This approach is illustrated in the next two example exercises.

### EXAMPLE: 7.0.2 Determining a Rate Law from Initial Rates

Ozone in the upper atmosphere is depleted when it reacts with nitrogen oxides. The rates of the reactions of nitrogen oxides with ozone are important factors in deciding how significant these reactions are in the formation of the ozone hole over Antarctica ([Figure 7.1](#p_WBmd)). One such reaction is the combination of nitric oxide, NO, with ozone, O3:

Figure 7.1

A contour map showing stratospheric ozone concentration and the “ozone hole” that occurs over Antarctica during its spring months. (credit: modification of work by NASA)

NO(g)+O3(g)⟶NO2(g)+O2(g)NO(g)+O3(g)⟶NO2(g)+O2(g)

This reaction has been studied in the laboratory, and the following rate data were determined at 25 °C.

|  |  |  |  |
| --- | --- | --- | --- |
| Trial | [NO] (mol/L) | [O3] (mol/L) | Δ[NO2]Δt(molL−1s−1)Δ[NO2]Δt(molL−1s−1) |
| 1 | 1.00 ×× 10−6 | 3.00 ×× 10−6 | 6.60 ×× 10−5 |
| 2 | 1.00 ×× 10−6 | 6.00 ×× 10−6 | 1.32 ×× 10−4 |
| 3 | 1.00 ×× 10−6 | 9.00 ×× 10−6 | 1.98 ×× 10−4 |
| 4 | 2.00 ×× 10−6 | 9.00 ×× 10−6 | 3.96 ×× 10−4 |
| 5 | 3.00 ×× 10−6 | 9.00 ×× 10−6 | 5.94 ×× 10−4 |

Determine the rate law and the rate constant for the reaction at 25 °C.

#### Solution

The rate law will have the form:

rate=k[NO]m[O3]nrate=k[NO]m[O3]n

Determine the values of m, n, and k from the experimental data using the following three-part process:

1. Step 1.
2. Step 2.
3. Step 3.

#### Check Your Learning

Acetaldehyde decomposes when heated to yield methane and carbon monoxide according to the equation:

CH3CHO(g)⟶CH4(g)+CO(g)CH3CHO(g)⟶CH4(g)+CO(g)

Determine the rate law and the rate constant for the reaction from the following experimental data:

|  |  |  |
| --- | --- | --- |
| Trial | [CH3CHO] (mol/L) | −Δ[CH3CHO]Δt(molL−1s−1)−Δ[CH3CHO]Δt(molL−1s−1) |
| 1 | 1.75 ×× 10−3 | 2.06 ×× 10−11 |
| 2 | 3.50 ×× 10−3 | 8.24 ×× 10−11 |
| 3 | 7.00 ×× 10−3 | 3.30 ×× 10−10 |

### Answer

rate=k[CH3CHO]2rate=k[CH3CHO]2 with k = 6.73 ×× 10−6 L/mol/s

### EXAMPLE: 7.0.3 Determining Rate Laws from Initial Rates

Using the initial rates method and the experimental data, determine the rate law and the value of the rate constant for this reaction:

2NO(g)+Cl2(g)⟶2NOCl(g)2NO(g)+Cl2(g)⟶2NOCl(g)

|  |  |  |  |
| --- | --- | --- | --- |
| Trial | [NO] (mol/L) | [Cl2] (mol/L) | −Δ[NO]Δt(molL−1s−1)−Δ[NO]Δt(molL−1s−1) |
| 1 | 0.10 | 0.10 | 0.00300 |
| 2 | 0.10 | 0.15 | 0.00450 |
| 3 | 0.15 | 0.10 | 0.00675 |

#### Solution

The rate law for this reaction will have the form:

rate=k[NO]m[Cl2]nrate=k[NO]m[Cl2]n

As in [Example 7.2](#Example_7.2), approach this problem in a stepwise fashion, determining the values of m and n from the experimental data and then using these values to determine the value of k. In this example, however, an explicit algebraic approach (vs. the implicit approach of the previous example) will be used to determine the values of m and n:

1. Step 1.
2. Step 2.
3. Step 3.

#### Check Your Learning

Use the provided initial rate data to derive the rate law for the reaction whose equation is:

OCl−(aq)+I−(aq)⟶OI−(aq)+Cl−(aq)OCl−(aq)+I−(aq)⟶OI−(aq)+Cl−(aq)

|  |  |  |  |
| --- | --- | --- | --- |
| Trial | [OCl−] (mol/L) | [I−] (mol/L) | Initial Rate (mol/L/s) |
| 1 | 0.0040 | 0.0020 | 0.00184 |
| 2 | 0.0020 | 0.0040 | 0.00092 |
| 3 | 0.0020 | 0.0020 | 0.00046 |

Determine the rate law expression and the value of the rate constant k with appropriate units for this reaction.

### Answer

 rate 2rate 3=0.000920.00046=k(0.0020)x(0.0040)yk(0.0020)x(0.0020)yrate 2rate 3=0.000920.00046=k(0.0020)x(0.0040)yk(0.0020)x(0.0020)y
2.00 = 2.00y
y = 1
 rate 1rate 2=0.001840.00092=k(0.0040)x(0.0020)yk(0.0020)x(0.0040)yrate 1rate 2=0.001840.00092=k(0.0040)x(0.0020)yk(0.0020)x(0.0040)y
 2.00=2x2y2.00=2x214.00=2xx=22.00=2x2y2.00=2x214.00=2xx=2
Substituting the concentration data from trial 1 and solving for k yields:
 rate=k[OCl−]2[I−]10.00184=k(0.0040)2(0.0020)1k=5.75×104mol−2L2s−1rate=k[OCl−]2[I−]10.00184=k(0.0040)2(0.0020)1k=5.75×104mol−2L2s−1

### 7.0.4 Reaction Order and Rate Constant Units

In some of our examples, the reaction orders in the rate law happen to be the same as the coefficients in the chemical equation for the reaction. This is merely a coincidence and very often not the case.

Rate laws may exhibit fractional orders for some reactants, and negative reaction orders are sometimes observed when an increase in the concentration of one reactant causes a decrease in reaction rate. A few examples illustrating these points are provided:

NO2+CO⟶NO+CO2rate=k[NO2]2CH3CHO⟶CH4+COrate=k[CH3CHO]22N2O5⟶NO2+O2rate=k[N2O5]2NO2+F2⟶2NO2Frate=k[NO2][F2]2NO2Cl⟶2NO2+Cl2rate=k[NO2Cl]NO2+CO⟶NO+CO2rate=k[NO2]2CH3CHO⟶CH4+COrate=k[CH3CHO]22N2O5⟶NO2+O2rate=k[N2O5]2NO2+F2⟶2NO2Frate=k[NO2][F2]2NO2Cl⟶2NO2+Cl2rate=k[NO2Cl]

It is important to note that rate laws are determined by experiment only and are not reliably predicted by reaction stoichiometry.

The units for a rate constant will vary as appropriate to accommodate the overall order of the reaction. The unit of the rate constant for the second-order reaction described in [Example 7.2](#Example_7.2) was determined to be Lmol−1s−1.Lmol−1s−1. For the third-order reaction described in [Example 7.3](#Example_7.3), the unit for k was derived to be L2mol−2s−1.L2mol−2s−1. Dimensional analysis requires the rate constant unit for a reaction whose overall order is x to be Lx−1mol1−xs−1.Lx−1mol1−xs−1.[Table 7.1](#Table_7.1) summarizes the rate constant units for common reaction orders.

Table 7.2

Rate Constant Units for Common Reaction Orders

|  |  |
| --- | --- |
| Overall Reaction Order (x) | Rate Constant Unit (Lx−1 mol1−x s−1) |
| 0 (zero) | mol L−1 s−1 |
| 1 (first) | s−1 |
| 2 (second) | L mol−1 s−1 |
| 3 (third) | L2 mol−2 s−1 |

Note that the units in this table were derived using specific units for concentration (mol/L) and time (s), though any valid units for these two properties may be used.

### 7.0.5 Order with Respect to a Reactant

### Example: 7.0.6 Determination of Order

Read this online at <https://edtechbooks.org/general_college_chemistry_2/experimental_rate_la>