The decomposition of NH₃ has been studied in great detail and a voluminous literature is available on this subject. Little information is available, however, concerning the homogeneous reaction rates. In an early paper, Hinshelwood and Burk investigated the homogeneous decomposition and noted that no measurable reaction occurred. Recently Sage, using a ballistic piston, has observed the reaction to proceed at temperatures above 1500 K. More recently, Hinshelwood and Burk have shown the reaction to proceed at temperatures between 1500 and 4500 K although no rate data were derived from these experiments.

The advent of the shock tube and "fast" infrared detectors has now made possible a method for the measurement of the homogeneous decomposition rate. In this paper we report the results of preliminary activation-energy measurements for the homogeneous decomposition of NH₃.

The use of the shock tube for emission measurements has been described by several authors. The present investigations are probably the first published data on reaction-rate measurements using infrared emission. Many shock-tube rate measurements have, however, been made by monitoring light intensities in the visible and ultraviolet regions of the spectrum. If it is assumed that radiative equilibrium is attained in times short compared with the chemical decomposition times after passage of the shock wave and, furthermore, that the NH₃ behaves as a transparent gas, then it is possible to measure the change in NH₃ concentration directly by following the infrared emission intensity as a function of time. Radiation of intermediates may be minimized by measuring only the initial concentration changes.

Rough calculations, based on the absolute intensity data of McKean and Schatz, indicate that the most advantageous band for a shock-tube study, considering available detectors and window materials, is the one centered near three microns. For judiciously chosen optical depths, the assumption of transparency may then be shown to be quite reasonable.
(99.99 per cent) and Matheson standard grade Ar (99.998 per cent). Before each run, a gas sample was taken directly from the low-pressure end of the tube for chemical analysis. For data reduction, only those runs which fell within about 3 per cent of the desired composition were used.

**Calculation of Thermodynamic Properties Behind the Shock**

From measured initial conditions and shock velocity, the thermodynamic parameters after the passage of the shock may be calculated by the use of the well-known conservation relations. Because the kinetic measurements were to be restricted to initial rates, only properties immediately behind the shock front were required. For the mixture containing 1 per cent NH₃, the calculations were performed neglecting the presence of the NH₃. Calculations for the 8 per cent mixture were carried out by assuming equilibrium before and after the shock wave with respect to the internal degrees of freedom but not with respect to chemical composition. Enthalpies for NH₃ were computed in the 2000 to 3000 K range with standard methods.¹²

**Experimental Results**

The initial rate of decomposition of NH₃ in Ar has been measured between 2000 and 3000 K by following the rate of decrease of emission intensity in the 2.7 to 3.2 μ wavelength region. A typical experimental record is shown in Figure 2. As was pointed out earlier, the intensity of the emission is assumed proportional to the concentration of NH₃. From the record it may be seen that the emission intensity jumps very rapidly to a maximum value after passage of the shock wave and then decays, approaching the zero level asymptotically.

![Figure 2](image-url)  
**Fig. 2.** Typical experimental record. Vertical lines are at 20 μsec intervals. \( X_{NH_3} = 0.08, \ T = 2720 \text{K}. \)
For reduction of the data it was assumed that

\[
-\frac{d(NH_3)}{dt} = k(NH_3)(M)
\]

where \((M)\) represents either Ar or NH\(_3\).

In terms of the NH\(_3\) partial pressure, \(p_{NH_3}\), and the mole fractions \(X_M\) and \(X_{NH_3}\), Equation (1) may be written as

\[
-\frac{RT}{X_{NH_3}p_T^2} \frac{dp_{NH_3}}{dt} = kX_M.
\]

Here \(p_T\) is the total pressure and \(T\) the temperature after the passage of the shock; \(t\) represents the actual time for the decomposition reaction and may be obtained from the “oscilloscope time” by multiplying by an appropriate density ratio. For transparent gases, \(p_{NH_3}\) is directly proportional to the emission intensity and, with properly designed electronic circuits, directly proportional to the oscilloscope voltage \(V\) registered by the detector. Hence

\[V = \text{const} \ p_{NH_3}\]

or

\[
\frac{dV}{dt} = \text{const} \ \frac{dp_{NH_3}}{dt}.
\]

The constant of proportionality may be evaluated at \(t = 0\) from the oscilloscope trace and the known partial pressure of NH\(_3\) immediately behind the shock front. Thus, in terms of the initial voltage \(V_0\) and mole fraction \(X_{NH_3}\), and with an assumed Arrhenius temperature dependence, Equation (2) becomes

\[
-\frac{RT}{X_{NH_3}p_T^2} \frac{dp_{NH_3}}{dt} = \frac{X_M}{X_{NH_3}} \frac{dV}{dt} = Be^{-E/RT}.
\]

For the initial stages of the reaction, \(T\) and \(p_T\) are approximately constant and \(X_{NH_3} \approx X_{NH_3}\). On this basis, an activation energy has been determined from Equation (4). The logarithm of the left-hand side of Equation (4) has been plotted against \(1/T\) for our measurements and the resulting graph is shown in Figure 3. Least-square straight-line fits of these data yield apparent activation energies of 51 kcal/mole for the mixture containing 1 per cent NH\(_3\) and 53 kcal/mole for the mixture containing 8 per cent NH\(_3\).

Fig. 3. The quantity \(\beta = \log \frac{(RT/X_{NH_3}p_T^2)(dp_{NH_3}/dt)}{dV/dt}\) versus \((1/T)\). The term \(dp_{NH_3}/dt\) has been evaluated for the initial decrease in NH\(_3\) emission.
Conclusions

It is natural to assume that the initial step in the homogeneous decomposition of NH₃ is

$$\text{NH}_3 + M \rightarrow \text{NH}_2 + H + M \quad (5)$$

where $M$ is either Ar or NH₃. Comparing our measured activation energy of about 52 kcal with the experimental value of about 105 kcal for the NH₂-H bond strength, it is clear that our data do not represent the initial decomposition rate. No doubt we are measuring the early rate and a chain mechanism is a reasonable explanation of our low-activation energy. We shall not attempt to guess the details of the actual mechanism but shall simply note that a chain with bimolecular termination would involve the square root of the initial rate constant, indicating an initial activation energy of about twice our measured value. This result follows, at least qualitatively, from the assumption that the reaction rate may be written

$$\text{rate of reaction} \propto r_p \left( \frac{r_i}{r_b} \right)^{1/2} \quad (6)$$

where $r_p$, $r_i$ and $r_b$ are the rates of chain propagation, initiation and breaking, respectively. For simple Arrhenius temperature dependence the apparent activation energy of Equation (6) would be

$$E = E_p + \frac{1}{2}(E_i - E_b) \quad (7)$$

where the subscripts have the same meaning as for the rates in Equation (6). For $E_p$ and $E_b$ small, we have $E_i \approx 2E$ in accord with our experimental findings.

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