

Evaluated Gas Phase Basicities and Proton Affinities of Molecules; Heats of Formation of Protonated Molecules

Cite as: Journal of Physical and Chemical Reference Data **13**, 695 (1984); <https://doi.org/10.1063/1.555719>
Published Online: 15 October 2009

Sharon G. Lias, Joel F. Liebman, and Rhoda D. Levin



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Evaluated Gas Phase Basicities and Proton Affinities of Molecules; Heats of Formation of Protonated Molecules

Sharon G. Lias

Center for Chemical Physics, National Bureau of Standards, Gaithersburg, MD 20899

Joel F. Liebman

Department of Chemistry, University of Maryland Baltimore County, Catonsville, MD 21228

and

Rhoda D. Levin

Center for Chemical Physics, National Bureau of Standards, Gaithersburg, MD 20899

The available data on gas phase basicities and proton affinities of molecules are compiled and evaluated. Tables giving the molecules ordered (1) according to proton affinity and (2) according to empirical formula, sorted alphabetically are provided. The heats of formation of the molecules and the corresponding protonated species are also listed.

Key words: basicity, heats of formation, ion-molecule reaction, proton affinity, proton transfer.

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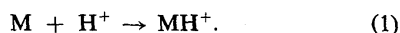
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1. Introduction

1.1. Definitions

The gas phase basicity and proton affinity of a molecule, M, are both defined in terms of the hypothetical reaction:

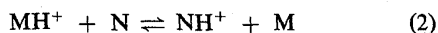


The gas phase basicity is the negative of the free energy change associated with this reaction, while the proton affinity is the negative of the corresponding enthalpy change.

1.2. Proton Affinities and Gas Phase Basicities: An Overview

Proton transfer reactions are of considerable importance in chemistry. Acid-base reactions have been studied extensively in solution for decades, but it is only in the past twenty years or so that experimental techniques have been devised to permit the quantitative study of the thermochemistry of proton transfer reactions in the gas phase. Particularly in the years since 1971, when the first gas phase ion-molecule equilibrium studies on proton transfer appeared, there has been a burgeoning of data in the literature, and consequently, a great interest in using information from such gas phase studies to distinguish between the chemical effects of solvation and the effects of intrinsic molecular properties. To date, although several excellent reviews of this active field of research have appeared¹⁻⁷, and two unevaluated compilations of data are available⁸⁻⁹, there was, until the current publication, no single reference which presented a comprehensive collection of data on gas phase proton affinities evaluated for internal consistency.

The vast majority of proton affinities presented here are based on measurements of the equilibrium constants of gas phase proton transfer reactions:



where:

$$-RT \ln K_{eq} = \Delta G_{Rn} = \Delta H_{Rn} - T\Delta S_{Rn} \quad (3)$$

and the equilibrium constant for reaction 2 is obtained from a mass spectrometric observation of the relative abundances of equilibrated ions, MH^+ and NH^+ , in a mixture of known composition of compounds M and N:

$$K_{eq} = [NH^+]/[MH^+][M]/[N] \quad (4)$$

The observed equilibrium constant of reaction 2 directly leads to a value for the Gibbs free energy

change of reaction, which represents the relative gas phase basicities of compounds M and N; if the entropy change of the reaction is determined, or can be reliably estimated, a value for the enthalpy change of reaction, or relative proton affinity, is obtained. Thus, the extensive scales of data presented here based on equilibrium constant measurements provide relative gas phase basicities and relative proton affinities, but do not directly give any information about absolute values of proton affinities, i.e., about the actual enthalpy changes of reaction 1 for the various compounds. Absolute values must be assigned based on some comparison standard incorporated in the thermochemical ladder for which heats of formation of both M and MH^+ are independently available. These species will necessarily be limited to those for which an MH^+ ion of known structure can be generated in a mass spectrometer, so that a reliable independent gas phase heat of formation will be known.

Thus, it must be emphasized that for data derived from equilibrium constant measurements, absolute values for proton affinities cited by authors depend on the proton affinity value selected for a comparison standard, and these may vary considerably from year to year, and from paper to paper. (For example, proton affinities varying from 202 kcal/mol to 210 kcal/mol have been cited for ammonia, which is often used as a comparison standard for scales of proton affinities). Although researchers working in the field are usually acutely aware of these arbitrary fluctuations in absolute values assigned to gas phase proton affinities, workers in related fields who seek to use these data often fall unwarily into the trap of using proton affinities from the literature without realizing that the experimental results actually lead only to relative thermochemical relationships, and that the absolute values assigned to proton affinities may have changed because of subsequent re-evaluations of the thermochemistry of a primary standard. The current publication seeks to provide a complete and internally-consistent set of gas phase proton affinity values based mainly on the vast body of data generated by equilibrium constant measurements, and to assign absolute values to the resulting proton affinity scale using the best current information about the thermochemistry of positive ions, imposing the requirement of internal consistency.

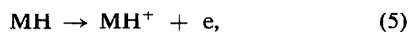
2. Types of Experiment Leading to Proton Affinity/Gas Basicity Data

In this compilation, as discussed above, most of the information is derived from scales of relative values for the gas basicities and proton affinities, based on (1) measurements of equilibrium constants of proton transfer reactions, or (2) use of the so-called "bracketing" technique. Absolute values are assigned to the scale using certain primary standards (discussed in

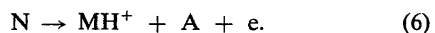
detail below), whose position in the proton affinity scale is known and for which well-established values of the heats of formation of both M and MH^+ are available.

2.1. Absolute Proton Affinity Values from Ionization Thresholds

Experiments in which the heat of formation of MH^+ is determined lead directly to values for the proton affinity when combined with a heat of formation of the corresponding neutral molecule, M. If MH is a sufficiently stable species that it can be introduced into a mass spectrometer or be generated in situ, or if MH^+ exists as a fragmentation product from some larger molecular species, absolute values for the heat of formation of MH^+ may be obtained through determinations of the ionization potential of MH :



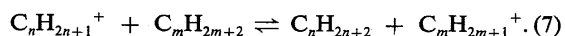
or the appearance potential of MH^+ from a larger molecule:



Because the heats of formation of MH^+ are known from ionization potential or appearance potential determinations, such "absolute" proton affinity determinations are labelled "Threshold Value" in Table 1. Such absolute values for proton affinities are available for only a very few species of the 780 compounds on which data is reported here, largely because in most cases the neutral MH molecule does not exist, and the energetics of formation of an MH^+ ion of the required structure in ionic fragmentation processes is not reliably established.

2.2. Proton Affinity Values from Thermochemical Information Derived from Hydride Transfer Equilibrium Constant Determinations

Values for heats of formation of a number of carbonium ions, $C_nH_{2n+1}^+$, are known from measurements of hydride transfer equilibrium constants in alkane mixtures:



In systems for which the heats of formation of three reactant species have been independently established, the heat of formation of the fourth can be determined. Since a knowledge of the heat of formation of the $C_mH_{2m+1}^+$ carbonium ion can be used to calculate an absolute proton affinity value for the corresponding olefin, C_mH_{2m} , several values based on hydride transfer equilibria are also included in the compilation; the heats of formation of ions from such experiments are all related to the heat of formation of the tert-butyl ion as

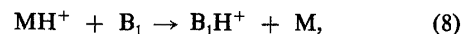
a primary standard. (See below for a discussion of the heat of formation of this standard.)

2.3. Relative Gas Basicity Values from Proton Transfer Equilibrium Constants

During the past decade, there has been extensive use of gas phase ion-molecule equilibrium constant measurements to establish the relative basicities of chemical compounds in the absence of solvent effects. In these studies, carried out in a number of laboratories using mainly high pressure mass spectrometry, ion cyclotron resonance spectroscopy, and flowing afterglow experiments, the equilibrium constants associated with proton transfer reactions such as (2) are measured, yielding the relative gas basicities of M and N from the free energy change associated with the reaction at a given temperature (Eq. 3). If assumptions are made about the entropy changes associated with reaction 2, or if experimental determinations of the entropy change have been made (Sec. 3.1.1), values for relative gas phase basicities obtained in this way can be translated into scales of relative proton affinities. Although these equilibrium constants are determined mass spectrometrically at pressures of 10^{-6} torr to approximately 1 torr, there is abundant evidence¹⁻⁷ that thermodynamic equilibrium is achieved in this type of experiment.

2.4. Relative Gas Basicity/Proton Affinity Values from "Bracketing"

There are certain species for which measurements of equilibrium constants for proton transfer reactions are difficult or impossible. These include free radicals and species for which the MH^+ ion undergoes fast reactions with M or alternate reaction channels with N. In these cases, the basicity is usually estimated using the technique known as "bracketing". In this approach, one reacts MH^+ with a series of bases, observing whether or not proton transfer occurs:



The basicity of M is assumed to lie between those of B_1 and B_2 , and where these species have known gas basicity values, a value can be assigned to M. Results obtained through bracketing experiments are generally less reliable than those obtained through other methods because of numerous possible complications. For instance, exothermic proton transfer reactions sometimes do not occur when there is an energetically favorable alternate channel open to the reactants. Furthermore, endothermic proton transfer reactions can be observed if the reaction has a negative Gibbs free

energy change, or if the reaction is not endothermic by more than ~ 8 kJ/mol (~ 2 kcal/mol). Finally, if there are several isomeric structures of B_1H^+ , the observed proton transfer reaction may involve a rearrangement of the B_1 or B_1H^+ species in the reaction complex to a more stable structure, so that the observed "bracketing" does not necessarily reflect the thermochemistry of a straightforward or even unique proton transfer reaction.

2.5. Other Sources of Proton Affinity/Gas Basicity Data

The compilation includes a few pieces of data originating from measurements which do not fall into any of the above categories. Quantitative information about relative proton affinities has been obtained through the determination of the energy barrier associated with endothermic proton transfer reactions through an Arrhenius treatment of the temperature dependence of the rate constants. Also, determinations of the equilibrium constants of processes such as:



lead to values of heats of formation of the condensation or association ions, ABH^+ , if the heats of formation of AH^+ and B are known; this information can then be used to derive the proton affinity of AB .

Qualitative information about relative proton affinities has been derived from observations of the modes of dissociation of $(ABH)^+$ ions into $(AH^+ + B)$ or $(BH^+ + A)$; a semi-quantitative relationship between the ratios of the two product ions and relative proton affinities has been reported¹⁰.

3. Description of the Evaluation

3.1. Thermodynamic Ladders

In the compilation of gas phase basicity and proton affinity data presented here, the available thermodynamic ladders giving relative gas phase basicities of series of compounds were made the initial basis for the evaluation, using consistency between sets of data from different laboratories and matching of the intervals of the scale with those predicted from widely separated primary standards as the criteria for judgement. To minimize problems due to uncertainties in the temperature at which measurements were made, the thermodynamic ladders were related to as many local standards over the course of the scale as possible. Results obtained from "bracketing" experiments were then related to the evaluated scale. A specific example of this procedure and the mode of presentation in Table 1 is given in Sec. 5.1. Finally, the experimentally-determined scale of relative gas basicities was translated

into a scale of proton affinities by evaluating the entropy change associated with reaction 1.

3.1.1. Entropy Changes

For most polyatomic species, the entropy change of the half reaction ($M \rightarrow MH^+$) was estimated by making the usual simplifying assumption¹⁻⁷ that this quantity can be adequately approximated by the expression:

$$\Delta S = R \ln[\sigma(M)/\sigma(MH^+)] \quad (11)$$

(where $\sigma(M)$ and $\sigma(MH^+)$ are the rotational symmetry numbers of M and MH^+). The use of this expression is obviously inexact, in some cases enough so that a more exact calculation of the entropy change from the complete partition function is warranted. This is especially true for species in which internal rotations are lost or gained upon protonation, or in which there is a significant difference in the moments of inertia of the protonated and unprotonated species. In such cases, the entropy change was calculated from the complete partition function. In all cases for which experimental entropy change determinations of reaction 2 have been made, the experimental results were made the basis of the evaluation of the relevant entropy changes.

3.1.2. Temperature

Several of the data sets constituting the backbone thermodynamic ladder for Table 1 are very extensive, covering energy ranges of up to 300 kJ/mol (72 kcal/mol). Uncertainties of only a few percent in the operating temperature used in the original experiments may translate into differences of tens of kJ (or several kcal) in the absolute values assigned to gas basicities or proton affinities when applied across such an energy range relative to a single absolute standard. One must be particularly aware of this problem in dealing with data sets from ion cyclotron resonance experiments originally reported to have been carried out at "300 K". More recent work has shown that unless special precautions are taken, the operating temperature of an ICR cell at "ambient" temperature is higher than 300 K by 20 to 50 degrees. In such cases, the free energy changes originally calculated from the observed equilibrium constant (Eq. 3) have been corrected by multiplying by $T(\text{corrected})/300$. Where such changes in the original data have been made, the magnitude of the correction factor is indicated in the notes under the literature citation. A comparison of data from these sets with analogous results from experiments in which the temperature was measured generally confirms that the magnitudes of these corrections are at least approximately correct. (It is obvious that for the data sets considered here, the distinction between "300 K" and the temperature corresponding to STP conditions,

298.15 K, is essentially irrelevant; the two numbers, 298 and 300, will be used in this discussion and in the tables essentially interchangeably, depending mainly on which number was used in a particular paper.)

3.2. Assignment of Absolute Values to the Scale: Reference Standards

Although absolute values have been assigned to the relative proton affinity scale here, these values can be considered really well-established only for the lower part of the scale (i.e., for proton affinities lower than about 750 kJ/mol or 180 kcal/mol), which is based on numerous primary comparison standards. The upper part of the scale is based primarily on the proton affinity of isobutene, which in turn depends on the heat of formation of the tert-butyl ion; other standards, which appear to give a consistent assignment to the scale, are ketene and ammonia. However, none of these standards can be considered to have a firmly-enough established proton affinity that it can be stated with certainty that future variations in absolute assignments will not occur.

To minimize problems due to uncertainties in the temperature, long thermodynamic ladders have been related to as many local standards over the course of the scale as possible. In such cases, the same data are listed twice in Table 1, once as originally reported and a second time in evaluated form, as described in more detail in Sec. 5.1. Similarly, in the single thermodynamic ladder available for the low end of the scale (H_2 to C_2H_6), in agreement with an evaluation of these data from the laboratory where the original work was done¹¹, we have chosen to relate segments of the scale to well-established local standards, rather than accept the originally-reported relationships between widely separated portions of the scale.

In the following section, we present details of the evaluations of the proton affinities of the compounds which served as the primary standards for assigning absolute values to the gas basicity and proton affinity scales. In this process, for all but the lowest part of the scale (H_2 to C_2H_6), the greatest weight was given to the proton affinities of ethylene, propylene, and isobutene. The heats of formation of $C_2H_5^+$ and $CH_3C^+HCH_3$ have been carefully studied in numerous laboratories, and are now rather well established. The heat of formation of the tert-butyl ion is somewhat less well established, but recent evaluations of this heat of formation give a proton affinity value for isobutene which is in fairly good agreement with information from other comparison standards in the immediate vicinity in the scale (ketene and ammonia). The relevant heats of formation, and evaluated 300 K proton affinity values of the standard compounds are summarized in Sec. 3.3. It should be noted that the heats of formation of positive ions cited here are given using the so-called "stationary electron convention" commonly used by

mass spectrometrists¹², and described in detail below (Sec. 5.2.1).

3.2.1. Ammonia

For reasons which appear to have more to do with tradition than with science, the scale of gas phase proton affinities is often related to the proton affinity of ammonia as a standard. This proton affinity is defined by the reaction:



An experimental determination of the heat of formation of NH_4^+ based on its appearance energy from a $(NH_3)_2$ dimer¹³:



leads to a value of the proton affinity of ammonia of 846.3 kJ/mol (202.3 kcal/mol) at 0 K, and assuming that all rotational degrees of freedom are excited and that no vibrational degrees of freedom are excited at 298 K, this corresponds to a value of 852 kJ/mol (203.6 kcal/mol) for the proton affinity of ammonia at 298 K. Although the authors of that paper cite error limits of ± 5 kJ/mol (1.3 kcal/mol) for this value of the proton affinity, an examination of the error limits associated with the appearance energy measurement (± 2 kJ/mol or 0.5 kcal/mol), the heat of formation of NH_2 (± 12 kJ/mol or 3 kcal/mol) and the assumed well depth for formation of the $(NH_3)_2$ dimer (± 4 kJ/mol or 1 kcal/mol) leads one to assign error limits of at least ± 13 kJ/mol (3.2 kcal/mol) to this value. Substitution of the dimer well depth used in reference 13 (-14.6 kJ/mol or 3.5 kcal/mol) with a value from a more recent ab initio calculation¹⁴ (-7.9 kJ/mol or 1.9 kcal/mol) leads to a modification of the proton affinity derived from this experimental result to a value of 845 kJ/mol (202.0 kcal/mol). More direct ab initio calculations of the proton affinity of ammonia^{15,16} lead to values of 858 kJ/mol (205.0 kcal/mol) and 879 kJ/mol (210 kcal/mol), respectively.

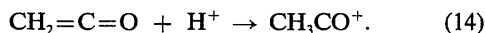
In fact, the exact value for the proton affinity of ammonia is of less importance than it is usually accorded, since the position of ammonia in the scale of relative gas phase basicities is not well established. In fact results from various laboratories show considerable variation in measurements involving this molecule. For instance, relative to the proton affinity of isobutene, one would assign values for the proton affinity of ammonia ranging from 851 to 858 kJ/mol (203.5 to 205.0 kcal/mol), depending on which set of measurements one selected. (Although this range is not really large in terms of absolute quantities, the reproducibility of measurements of proton transfer equilibrium constants often allows one to assign error limits of < 0.4 kJ/mol (0.1 kcal/mol) to any particular directly-measured

interval in the free energy scale. The apparently poor quality of the measurements involving this compound is easy to rationalize, since ammonia tends to adsorb on surfaces making accurate pressure measurements difficult.) The selected value for the proton affinity of ammonia, 853.5 kJ/mol (204.0 kcal/mol), which is in good agreement with the experimental determination¹³ and one theoretical calculation¹⁵, is an intermediate value taken from the several thermochemical ladders which have been reported relating the proton affinity of ammonia to that of the primary standard isobutene. Because the agreement between the different thermochemical scales is generally good except for the position of ammonia, the exact value of the proton affinity of ammonia was given little weight in carrying out the evaluation reported here. This proton affinity was treated as a secondary standard, and, in fact, was allowed to "float" relative to the rest of the scale in evaluating different data sets, in order to maintain the overall agreement between the different experimentally-determined thermodynamic ladders.

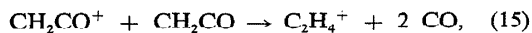
The entropy change associated with the half reaction ($\text{NH}_3 \rightarrow \text{NH}_4^+$) is -7.5 J/mol K (-1.8 cal/mol K), calculated from the complete partition function. This leads to a value of 818 kJ/mol (195.6 kcal/mol) for the 300 K gas phase basicity.

3.2.2. Ketene

The proton affinity is defined by the reaction:



A recent determination¹⁷ of the heat of formation of CH_3CO^+ at 298 K led to a value of 657 kJ/mol (157 kcal/mol). Accepting a value of -47.7 kJ/mol (-11.4 kcal/mol) for the heat of formation of ketene¹⁸, this would correspond to a proton affinity of 825.5 kJ/mol (197.3 kcal/mol). A re-evaluation of the data of that reference by the present authors gives a heat of formation for CH_3CO^+ of 653 kJ/mol (156 kcal/mol) which corresponds to a proton affinity of 830 kJ/mol (198.3 kcal/mol). However, from the onset of 9.6035 eV determined¹⁹ for the occurrence of the reaction:

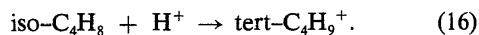


one calculates a slightly less negative heat of formation for CH_2CO , -36.4 kJ/mol (-8.7 kcal/mol), which would correspond to a proton affinity¹⁷ of 832 kJ/mol (198.8 kcal/mol), or 836 kJ/mol (199.8 kcal/mol), respectively.

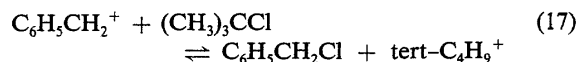
The position of ketene in the thermochemical ladder of proton affinities has been established^{12,20}, and relative to the value chosen above for isobutene, leads to a proton affinity for ketene of 828 kJ/mol (198.0 kcal/mol).

3.2.3. Isobutene

The proton affinity of isobutene is defined by the reaction:

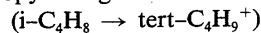


The heat of formation of the $\text{tert-C}_4\text{H}_9^+$ ion is less well established than those of the ethyl and *sec*-propyl ions discussed below. A recent analysis of the 298 K heat of formation of this ion as derived from appearance energy measurements led to a value of $<695 \pm 3 \text{ kJ/mol}$ ($166.2 \pm 0.8 \text{ kcal/mol}$)²¹. A value of 697 kJ/mol (166.5 kcal/mol) is obtained for this heat of formation taking a value of $50.2 \pm 4 \text{ kJ/mol}$ ($12.0 \pm 1 \text{ kcal/mol}$) for the heat of formation of the *tert*-butyl radical²² and a value of 6.7 eV for the ionization potential²³. However, the latter value can not be considered well-established without additional corroborating evidence, since experimentally-determined values for the heat of formation of the radical ranging from 44 to 35 kJ/mol (10.5 to 8.4 kcal/mol)²⁴⁻²⁷ have also been reported, and a value of 6.58 eV has been reported for the ionization potential²⁸. The corroborating evidence is derived from measurements²⁹⁻³¹ of the equilibrium constant of the reaction:



which give a value of -5.4 kJ/mol (-1.3 kcal/mol) for the enthalpy change at 300 K; from these data the heat of formation of the *tert*-butyl ion would be estimated to be 694 kJ/mol (165.8 kcal/mol) accepting a value of 899 kJ/mol (214.8 kcal/mol) for the heat of formation of the benzyl ion. This is based on a value for the heat of formation of the benzyl radical of 204 kJ/mol (48.8 kcal/mol)^{22,32} and an ionization potential of the radical of 7.20 eV³³. (The heat of formation of $(\text{CH}_3)_3\text{CCl}$ is taken as -182 kJ/mol or -43.5 kcal/mol ³⁴; the heat of formation of $\text{C}_6\text{H}_5\text{CH}_2\text{Cl}$ is derived from a liquid phase heat of formation³⁴ and a heat of vaporization³⁵ taken to be 4.1 kcal/mol.) Based on this result, the proton affinity of isobutene is 820 kJ/mol (195.9 kcal/mol).

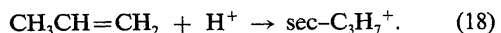
Taking into account changes in external rotational symmetry numbers, the gain of an internal rotation upon protonation, and the change in the moments of inertia, the entropy change for the half reaction



is about $+3 \text{ J/mol K}$ ($+0.7 \text{ cal/mol K}$)³⁶. This leads to a value of 784 kJ/mol (187.3 kcal/mol) for the 298 K gas phase basicity.

3.2.4. Propylene

The proton affinity of propylene is defined by the reaction:



The heat of formation of the $\text{sec-C}_3\text{H}_7^+$ ion at 298 K has been determined to be 798 kJ/mol (190.8 kcal/mol)³⁷, 799 kJ/mol (190.9 kcal/mol)³⁸, or 802.5 kJ/mol (191.8 kcal/mol)²¹. Taking a value of 799 kJ/mol (191.0 kcal/mol) for this heat of formation, the proton affinity of propylene is 751 kJ/mol (179.5 kcal/mol).

A calculation of the rotational entropy change associated with changes in the moments of inertia and rotational symmetry numbers for the half reaction ($\text{CH}_3\text{CH}=\text{CH}_2 \rightarrow \text{sec-C}_3\text{H}_7^+$) leads to a value of -3.8 J/mol K (-0.9 cal/mol K); this estimate fails to take into account the predicted positive entropy change brought about by the loss of the double bond. (The methyl group in the ion can be taken to be essentially a free rotor³⁹.) In the evaluation of the gas basicity of propylene, it has been assumed that the entropy change of this half reaction is close to zero, as discussed before³⁶. On this basis, the gas basicity of propylene is taken to be 718 kJ/mol (171.7 kcal/mol).

The gas basicity of propylene has been related through thermochemical ladders to several of the other comparison standards listed here. If one estimates the proton affinity through such data using ethylene as the primary standard for the proton affinity scale (i.e. using the data from reference 40 for the interval C_2H_4 - H_2O and from reference 41 for the interval H_2O - C_3H_6), one obtains a value of 752 kJ/mol (179.7 kcal/mol) for the proton affinity of propylene, in excellent agreement with the conclusion presented above. On the other hand, the interval in the thermochemical ladder between isobutene and propylene appears to be somewhat constricted, an effect which may result from the fact that small inaccuracies in temperature measurement may cause noticeable errors in such a scale if the scale covers a wide energy range, as discussed above, or may possibly reflect uncertainties in the heat of formation of the *tert*-butyl ion, also discussed above.

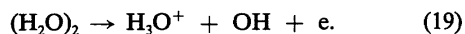
3.2.5. Formaldehyde

Approximately following the recent discussion of Collyer and McMahon⁴⁰, an analysis of the proton affinity of formaldehyde is given. Although this compound was not used as a primary reference standard in the evaluation of the thermochemical ladder, this proton affinity provides an additional reference point to verify the assignment of absolute values to the proton affinity scale. Values for the heat of formation of CH_2OH^+ derived from the appearance energy of this ion from CH_3OH range from 711 kJ/mol (170.0 kcal/mol) to 714 kJ/mol (170.7 kcal/mol)⁴²⁻⁴⁴ corresponding to proton affinity values of 711-715 kJ/mol (170-171 kcal/mol). Since the proton affinity of formaldehyde is 22 kJ/mol (5.2 kcal/mol) above that of water⁴⁵, the value predicted from the evaluated proton affinity scale is 718 kJ/mol (171.7 kcal/mol), in

reasonably good agreement with that obtained from threshold measurements.

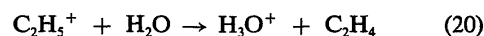
3.2.6. Water

A value for the heat of formation of H_3O^+ has been obtained from a measurement of the appearance energy of this ion from a hydrogen-bonded dimer⁴⁶:



From this experiment, one obtains a value for the proton affinity of H_2O of 694 kJ/mol (165.8 kcal/mol) at 0 K, and assuming that all rotational degrees of freedom are excited and that no vibrational degrees of freedom are excited at 298 K, this corresponds to a value of 697 kJ/mol (166.5 kcal/mol) for the proton affinity of water at 298 K. The error limits cited by the authors for this proton affinity are ± 7.5 kJ/mol (1.8 kcal/mol), with the major uncertainty being the well depth for formation of the water dimer. A recent state-of-the-art *ab initio* calculation⁴⁷, which included corrections for the change in the zero point energy upon protonation, correction to 298 K, and vibrational mode corrections, verifies the well depth of -17 kJ/mol (-4 kcal/mol) assumed in the interpretation of those results; these authors also calculate a value for the proton affinity of water of 689 kJ/mol (164.8 kcal/mol), in good agreement with the experimental result.

Recent equilibrium studies^{40,48} have tied the proton affinity of H_2O to that of ethylene. The experimental results from those studies indicate that the Gibbs free energy change of the reaction:



is -7.5 kJ/mol (-1.8 kcal/mol)⁴⁸ or -14 kJ/mol (-3.4 kcal/mol)⁴⁰. In one of these studies⁴⁸, the direct measurement of the free energy change of reaction 20 was complicated by clustering of H_3O^+ to C_2H_4 , and therefore, more weight is given to the latter result, which is based on a low pressure measurement of a thermodynamic ladder linking the gas basicities of the two compounds through several interlocking steps. Accepting a value for the proton affinity of C_2H_4 of 680 kJ/mol (162.6 kcal/mol) (see Discussion below) this result leads to a proton affinity of water of 697 kJ/mol (166.7 kcal/mol) if one calculates the entropy change for reaction 20 taking into account only the changes in the rotational symmetry numbers of the reactants and products; a more complete calculation of the entropy change for this reaction leads to a value of 696 kJ/mol (166.3 kcal/mol). Both estimates of the entropy change give proton affinity values which are in agreement with the value derived from the threshold determination of the heat of formation of H_3O^+ ⁴⁶ and with the *ab initio* calculation⁴⁷.

The entropy change associated with the half

reaction ($\text{H}_2\text{O} \rightarrow \text{H}_3\text{O}^+$) is 4.3 J/mol K (1.03 cal/mol K)⁴⁹. The evaluated proton affinity of water, 697 kJ/mol (166.5 kcal/mol), thus corresponds to a gas phase basicity of 665 kJ/mol (159.0 kcal/mol).

3.2.7. Ethylene

The proton affinity of ethylene is defined by the reaction:



The heat of formation of C_2H_5^+ at 298 K has been determined from photoion-photoelectron coincidence experiments to be 901 ± 4 kJ/mol (215.3 ± 1.0 kcal/mol)³⁷, or 903 ± 2 kJ/mol (215.9 ± 0.5 kcal/mol)³⁸, and from photoionization mass spectrometric appearance potentials, to be 904 ± 2 kJ/mol (216.0 ± 0.5 kcal/mol)²¹. In agreement with recent analyses of these data^{40,48}, we take an average value of 902 kJ/mol (215.6 kcal/mol) for this heat of formation, and on that basis, take a value for the proton affinity of ethylene of 680 kJ/mol (162.6 kcal/mol).

Assuming the C_2H_5^+ ion to have a non-classical bridged structure of C_{2v} symmetry¹⁶ (which has been calculated⁵⁰ to be 21.6 kJ/mol or 5.2 kcal/mol lower in energy than the classical structure), the entropy change calculated from the complete partition function for the half reaction ($\text{C}_2\text{H}_4 \rightarrow \text{C}_2\text{H}_5^+$) is -10 J/mol K (-2.4 cal/mol K). The gas basicity of this compound is 651 kJ/mol (155.6 kcal/mol).

3.2.8. Carbon monoxide

The heat of formation of HCO^+ derived from appearance potential measurements in formaldehyde is 824 kJ/mol (196.9 kcal/mol) corresponding to a proton affinity of 596 kJ/mol (142.4 kcal/mol)⁵¹, or 827 kJ/mol (197.6 kcal/mol) corresponding to a proton affinity of 593 kJ/mol (141.7 kcal/mol)⁵². The selected value is the average of these two and some other values from appearance potential measurements⁵³, or 594 kJ/mol (141.9 kcal/mol). Accepting a recent recommended value for the heat of formation of HCO^+ of 37 ± 5 kJ/mol (8.9 ± 1.2 kcal/mol), an ionization potential determination⁵³ for this radical of 8.27 ± 0.01 eV leads to a heat of formation of the HCO^+ ion of 835 kJ/mol (199.6 kcal/mol), and a corresponding proton affinity of 584.5 ± 6 kJ/mol (139.7 ± 1.4 kcal/mol), in good agreement with the values derived above.

Following the recommendation of Bohme et al.¹¹, the entropy change associated with the half reaction ($\text{CO} \rightarrow \text{HCO}^+$) is taken as 3 J/mol K (0.8 cal/mol K).

3.2.9. Carbon dioxide

From the appearance potential of HCO_2^+ in formic acid (12.29 eV)⁵⁴, one can derive a heat of formation of the ion of 589 kJ/mol (140.8 kcal/mol), which corresponds to a proton affinity of 548 kJ/mol (130.9 kcal/mol).

Following the recommendation of Bohme et al.¹¹, the entropy change associated with the half reaction ($\text{CO}_2 \rightarrow \text{HCO}_2^+$) is taken as 19 J/mol K (4.5 cal/mol K).

3.2.10. Oxygen atom

The heat of formation of the OH^+ ion has been determined from its appearance potential in H_2O (18.115 ± 0.008 eV) to be 1293 kJ/mol (308.96 kcal/mol)⁵⁵. A value for the ionization potential of the OH radical has been derived from appearance energy measurements in HOF to be 12.88 eV⁵⁶ in good agreement with the value of 13.01 eV⁵⁷ obtained from a direct experimental measurement. These values correspond to heats of formation of the OH^+ ion of 1281.5 or 1294 kJ/mol (306.3 or 309.3 kcal/mol), respectively. Accepting the value of 1293 kJ/mol (308.96 kcal/mol)⁵⁵, the proton affinity of the O atom is taken as 487 kJ/mol (116.3 kcal/mol).

Bohme et al.¹¹ recommend a value of 27 J/mol K (6.5 cal/mol K) for the entropy change associated with the half reaction ($\text{O} \rightarrow \text{OH}^+$). On this basis, the gas basicity of the O atom at 298 K is taken as 463 kJ/mol (110.7 kcal/mol).

3.2.11. Molecular Oxygen

The ionization potential of O_2H has been determined to be 11.35 ± 0.01 eV⁵⁸. Accepting values for the heat of formation of O_2H of 13 kJ/mol (3.2 kcal/mol) at 0 K, or 10.5 kJ/mol (2.5 kcal/mol) at 298 K⁵⁹, one obtains a value for the heat of formation of O_2H^+ ion of 1108 or 1098 kJ/mol (264.9 or 262.4 kcal/mol) at 0 or 298 K, respectively, leading to a value for the proton affinity of oxygen of 420 kJ/mol (100.5 kcal/mol). This result is in good agreement with a value of 422 kJ/mol (100.9 kcal/mol) which has been cited⁶⁰ for this proton affinity, based on the appearance potential of O_2H^+ in H_2O_2 .

The entropy change associated with the half reaction ($\text{O}_2 \rightarrow \text{O}_2\text{H}^+$) is taken to be 27 J/mol K (6.5 cal/mol K)¹¹. The gas basicity of O_2 is taken as 397 kJ/mol (95.0 kcal/mol).

3.3. Summary of Thermochemical Data on Comparison Standards for Proton Affinity Scale

Standard(M)	$\Delta_f H(M)(298\text{ K})$		$\Delta_f H(MH^+)^b$			Proton Affinity ^c		
	kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol	
NH ₃	-11.02	-46.11	⁵⁹	151.0	632.	¹³	203.6	852.
				153.6	643.	^{13,14}	202.0	845.
						¹⁵	205.0	858.
						¹⁶	210.0	879.
							204.0±3*	853.5±12*
CH ₂ =C=O	-11.4	-47.7	¹⁸	156.0	653.	¹⁷	198.3	830.
	-8.7	-36.4	¹⁹				198.8	832.
							198.0±2*	828.±8*
From thermochemical ladder, relative to isobutene:								
(CH ₃) ₂ C=CH ₂ -4.04		-16.9	³⁴	<166.2	<695.	²¹	>195.5	>818.
				166.5	697.	^{22,23}	195.2	817.
				165.8	694.	^{22,31,32,33}	195.9±1.5*	819.5±6*
CH ₃ CH=CH ₂	4.83	20.2	³⁴	190.8	798.	³⁷	179.7	752.
				190.9	799.	³⁸	179.6	752.
				191.8	802.5	²¹	178.7	748.
							179.7	752.
				191.0	799.		179.5±0.8*	751.±3*
From thermochemical ladder, relative to ethylene:								
H ₂ O						⁴⁷	164.8	689.
							166.3	695.8
From thermochemical ladder, relative to ethylene:								
	-57.8	-241.8	⁵⁹	141.4	592.	⁴⁶	166.5±2*	697.±8*
CH ₂ =CH ₂	12.48	52.2	³⁴	215.3	901	³⁷	162.9	681.5
				215.9	903	³⁸	162.3	679.
				215.6	902		162.6±1*	680.±4*
CO	-26.4	110.5	⁵⁹	196.9	824.	⁵¹	142.4	596.
				197.6	827.	⁵²	141.7	593.
				199.6	835.	^{27,53}	139.7	584.5
							141.9±2*	594.±8*
CO ₂	-94.05	-393.5	⁵⁹	140.8	589.	⁵⁴	130.9*	548.*
O				306.3	1281.5	⁵⁶	118.9	498.
				309.3	1294.	⁵⁷	115.9	485.
	59.56	249.2	⁵⁴	308.96	1293.	⁵⁵	116.3±1*	491.±4*
O ₂	0.00	0.00		262.4	1098.	^{58,59}	100.5	420.
				264.8	1108.	⁶⁰	100.9±0.5*	422.±2*

*Selected value.

^aSee Sec. 3.2 for detailed discussion and error limits.^bHeats of formation using "stationary electron convention" (described in Sec. 5.2.1)^c298 K heat of formation of H⁺ (Eq. 1) in "stationary electron convention" = 365.7 kcal/mol, 1530.0 kJ/mol.

4. Experimental Error Limits

As shown in Sec. 3.3, the heats of formation of the MH^+ ions used for establishing absolute values for the proton affinity scale have error limits which are larger than ± 4 kJ/mol (± 1 kcal/mol), sometimes considerably larger (e.g., NH_4^+). In general, absolute values assigned to the proton affinities of species determined through equilibrium measurements can be considered to be known to within ± 8 kJ/mol (± 2 kcal/mol) if the proton affinity lies in the region of the scale between water and ammonia (i.e. if the proton affinity is in the range 700–850 kJ/mol). In the region of the scale below water (proton affinities less than about 700 kJ/mol), the absolute values of proton affinities can be assumed to be known to within 4–6 kJ/mol for species determined through equilibrium measurements. In the region of the scale above ammonia (proton affinities greater than 850 kJ/mol), the absence of reliable absolute standards makes it impossible to check on the reliability of assigned proton affinity values. In particular, if the temperatures at which measurements were made were not well known, the thermochemical ladders generated from equilibrium constant determinations could be too long or too short, causing proton affinity values at the top end of the scale to vary considerably from their correct absolute values. Some confidence in the assigned values is gained from the general agreement observed in the reported thermochemical ladders generated in different laboratories. However, the lack of reference proton affinities in the upper part of the scale necessitates the assignment of relatively large error limits in this region, ± 16 kJ/mol. Throughout the scale, relative values derived from equilibrium constant measurements can be considered to be known to within ± 1 kJ/mol (0.2 kcal/mol) or less when ΔG is small (≤ 4 –6 kJ/mol). Since the experimentally measured free energy changes are given in Table 1, users of this compilation can identify for themselves species for which larger or smaller error limits on relative gas basicities are warranted. Proton affinities derived from bracketing measurements have error limits which are determined by the size of the bracket (given under the appropriate literature reference).

In Table 2, the values given for the heats of formation of the positive ions include, of course, the error limits associated with the assignment of absolute

values to the proton affinity scale (described above in Sec. 3.2) as well as the error limits associated with the heat of formation of the neutral molecule.

Nearly all of the data compiled here were originally reported in units of kcal/mol; the cited values in these units are given to tenths of a kcal/mol, in order to preserve the originally-reported information about relative ordering. On the other hand, in order to emphasize the large absolute uncertainties in cited proton affinity values, the columns giving data in kJ/mol show values only to within the nearest kilojoule per mole.

5. Explanation of the Tables

5.1. Explanation of Table 1

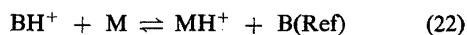
Table 1 presents a summary of the data from the literature, an evaluation of those data, and evaluated values for the gas basicities at 300 K and the proton affinities, in units of both kcal/mol and kJ/mol (where one kcal/mol is defined as 4.184 kJ/mol). The data in Table 1 are presented in order of descending proton affinity. Data from proton transfer equilibrium measurements are given in considerable detail, with (as described below) reference bases and temperatures specifically noted in most cases. The bracketing experiments, denoted by (br), have been related to the evaluated thermodynamic ladders from equilibrium constant measurements. Details of bracketing and hydride transfer experiments are provided in annotations under the appropriate literature references, as is information about equilibrium measurements related to reference bases not included in the standard list given below in Sec. 5.1.1; these latter experiments are identified in Table 1 by the notation (Key) appearing in the column usually used to identify the reference base.

In order to understand the structure of Table 1 (as well as the thought processes which went into the evaluation) it is best to consider specific examples of data sets, and illustrate how they are presented in Table 1. Table A represents a small portion of an extensive set of data originally published in J. F. Wolf, R. H. Staley, I. Koppel, M. Taagepera, R. T. McIver, Jr., J. L. Beauchamp, and R. W. Taft, *J. Am. Chem. Soc.* **99**, 5417 (1977) (denoted by 77WOL/STA in Table 1).

TABLE A.

	ΔG kcal/mol	Reference
NH ₃	0.0	77WOL/STA
CH ₃ COCH ₃	-7.2	77WOL/STA
iso-C ₄ H ₈	-8.6	77WOL/STA
H ₂ S	-27.8	77WOL/STA
H ₂ O	-31.4	77WOL/STA

In Table A, ΔG represents the free energy changes (in kcal/mol) derived for the process $(MH^+ + NH_3 \rightarrow NH_4^+ + M)$ from interlocking series of equilibrium constant determinations at a temperature assumed to be 300 K. As one can find in the comments under 77WOL/STA in the annotated references, the authors of this paper subsequently found that the operating temperature at which the original measurements were made was actually approximately 320 K, and therefore, from Eq. (3), the experimentally determined values of ΔG were increased by a factor of 320/300. These data then would appear in Table 1 as shown in Table B with the temperature-corrected values of ΔG (these are surrounded by parentheses) in the column labelled "Relative gas basicity, kcal/mol". The "Relative gas basicity" is the negative of the Gibbs free energy change associated with the reaction:



where B(Ref) is the reference base corresponding to the one- or two- alphabetic character denoter given in the list in Sec. 5.1.1.

TABLE B.

	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Reference
NH ₃	A	(0.0)	195.6	77WOL/STA
CH ₃ COCH ₃	A	(-7.9)	187.9	77WOL/STA
iso-C ₄ H ₈	A	(-9.2)	186.4	77WOL/STA
H ₂ S	A	(-29.6)	167.0	77WOL/STA
H ₂ O	A	(-33.5)	163.1	77WOL/STA

The entry "A" in Table B in the column labelled "Reference base" identifies ammonia as the species to which the original authors referred their basicity data. The "Gas basicity" values displayed here next to the corresponding "Relative gas basicity" are taken relative to the absolute gas basicity of ammonia derived in Sec. 3.2.1 and obtained using the relative ΔG values cited here.

Since, as described in Secs. 3.2.3 and 3.2.6, the gas basicities of iso-C₄H₈ and water are taken to be, respectively, 187.3 and 159.0 kcal/mol, it appears that in this data set cumulative errors exist in the overall thermochemical ladder over the span of more than 30 kcal/mol. Therefore, the data from this paper are related to several local standards. In the example set shown here, iso-C₄H₈ and H₂O happen to be species for which well-established values for the gas basicity and proton affinity can be assigned (Sec. 3.2), and therefore, these are chosen as local standards. In Table 1, data from any given reference are always reproduced in their original form (or, as here, their original form corrected for temperature) in addition to appearing in their evaluated form (if the two differ); original thermochemical ladders have been preserved in the presentation so that users of the compilation can examine the information as it appeared in the literature and re-evaluate the data for themselves if future experimental results bring about changes in values assigned to particular reference gas basicities or proton affinities. Thus, any given piece of data from 77WOL/STA could appear twice, once as given above and once with the cited gas basicity value as derived in relation to the appropriate local standard. The example data would be given in Table 1 as shown in Table C.

TABLE C.

	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Reference
NH ₃	A	(0.0)	195.6	77WOL/STA
	H		196.6	
CH ₃ COCH ₃	A	(-7.9)	187.9	77WOL/STA
	H		188.9	
iso-C ₄ H ₈	A	(-9.2)	186.4	77WOL/STA
	H	(0.0)	187.3	
H ₂ S	A	(-29.6)	167.0	77WOL/STA
	N		162.9	
H ₂ O	A	(-33.5)	163.1	77WOL/STA
	N	(0.0)	159.0	

In Table C (see the list in Sec. 5.1.1) the symbols H and N identify iso-C₄H₈ and H₂O as respective local

standards; the gas basicity values taken relative to these standards are shown in the appropriate column. In Table 1, when two entries appear together without a repetition of the "Reference" identifier, the second entry represents the same data item as the first but, as in the example given here, the "Gas basicity" in the second instance is that relative to a local standard not used by the original authors. In cases where such dual entries occur, only the second "evaluated" gas basicities were taken into account in the final evaluation of the gas basicity of the particular compound. The local standards used for particular data sets are identified in the list in Sec. 5.1.1 and also can be located in Table 1 by the appearance of the entry "(0.0)" in the "Relative gas basicity" column.

The evaluation of the gas basicity and proton affinity of a particular compound and its reconstruction from the entries in Table 1, can be illustrated by considering a single molecule from our example data set, H_2S . All the entries from Table 1 relating to H_2S are reproduced in Table D. However, for clarity of presentation we want the gas basicity values taken into account in the final evaluation to stand out and therefore, all the entries of "unevaluated" gas basicities (e.g. the basicity of H_2S relative to that of ammonia in the example given above) are here replaced by a row of dots.

TABLE D.

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Reference
H_2S				
340	Y	(0.0)	162.8	78FRE/HAR(2)
296	N	(4.6)	163.6	73HOP/BON
298	N	(3.8)	162.8	78TAN/MAC
320	A	(-29.6)	77WOL/STA
320	N		162.9	
550	N	(3.9)	162.7	77MAU/FIE
600	A	(-31.8)	79LAU
600	N		162.2	

In Table D, the absence of entries in the "Reference" column indicates that the particular experimental result comes from the reference cited immediately above; such a result has been related to a local standard (water, N, in this case) which is different from the comparison standard used by the original authors. In all such cases, it is the second entry which is taken into account in the final evaluation of the absolute gas basicity of the compound. Other entries shown here (73HOP/BON, 78TAN/MAC, 77MAU/FIE)

are clearly results derived from studies in which the authors related their experimentally-determined thermochemical ladders to water. The first entry, with a Y in

the "Reference base" column and (0.0) under "Relative gas basicity" shows that in the paper 78FRE/HAR(2), the authors have related the gas basicities of some other compound(s) to that of H_2S (symbol, Y); the gas basicity given opposite this entry is the evaluated value, 162.8 kcal/mol, derived from the five determinations relative to an H_2O standard.

A final example will illustrate how results from bracketing experiments are evaluated and listed in Table 1. In J. J. Solomon and R. F. Porter, J. Am. Chem. Soc. **94**, 1443 (1972) (72SOL/POR) it was seen that H_3O^+ transfers a proton to B_5H_9 , but H_3S^+ does not. Therefore, the gas basicity of B_5H_9 is assumed to lie between those of water and hydrogen sulfide. The entry for B_5H_9 is shown in Table E. The gas basicity given is the average of that of the two bracketing compounds; these are listed in the comments under the appropriate reference.

TABLE E.

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Reference
B_5H_9	(br)		161	72SOL/POR

In many instances, investigators have determined proton transfer equilibrium constants of pairs which do not include any of the standards listed above in Secs. 3.2 and 3.3., nor even any of the more commonly-used secondary standards listed in Sec. 5.1.1. In such cases, the data have been tied in to the scale in the usual way by relating to evaluated thermochemical ladders, and the identities of the particular reference compounds are given in comments under the appropriate reference at the end of Table 1. Such cases are identified by the designation (Key) appearing in the "Reference base" column.

In some cases, "proton affinities" have been reported in the literature without any information about the details of the experiment leading to the cited value; such data are usually reported here as they originally appeared since evaluation is impossible. Unevaluated data are denoted by a double asterisk (**). Other papers have presented revisions by original authors of data published earlier; in such cases the details of the original experiments are available, but the nature of the revisions are not; these data also have not been evaluated, and are denoted by a single asterisk (*).

The entries in the column labelled "Gas basicity" are actual gas phase basicities only for measurements made at temperatures close to 300 K. The gas basicities listed for experiments at higher temperatures have been normalized to those at 300 K in order to make

intercomparison easier; that is, in the defining reaction, that part of the term $T\Delta S$ which is associated with the proton is taken as 32.6 kJ/mol (7.8 kcal/mol)⁶¹ in all cases, so that actual differences in the free energy change of the defining reaction due to entropy changes associated with the half reaction ($M \rightarrow MH^+$) may be discerned.

Literature references, identified by the eight-character identifier at the end of a data-containing line, are given at the end of the Table 1, sorted alphabetically according to the characters appearing in the identifying tag. The individual references contain annotations giving any special remarks pertaining to the particular paper.

5.1.1. Explanation of Symbols Used in Table 1

Symbol used in Table 1	Reference base	Evaluated gas basicity (300 K)		Evaluated proton affinity	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol
A	NH ₃	195.6	818.	204.0	853.5
B	CH ₃ NH ₂	205.7	861.	214.1	896.
C	n-C ₃ H ₇ NH ₂	210.1	879.	217.9	912.
D	Pyridine	213.1	892.	220.8	924.
E	(CH ₃) ₃ N	217.3	909.	225.1	942.
F	C ₆ H ₅ NH ₂	202.5	847.	209.5	876.5
G	C ₂ H ₅ NH ₂	208.5	872.	217.0	908.
H	(CH ₃) ₂ C=CH ₂	187.3	784.	195.9	820.
I	C ₆ H ₅ N(CH ₃) ₂	215.4	901.	223.4	935.
J	C ₆ H ₆	174.6	730.5	181.3	758.5
K	C ₂ H ₄	155.6	651.	162.6	680.
L	CH ₂ (CN) ₂	167.4	700.	175.6	735.
M	H ₂ CO	164.3	687.	171.7	718.
N	H ₂ O	159.0	665.	166.5	697.
O	O	110.7	463.	116.3	487.
O'	O ₂	95.0	397.	100.9	422.
P	(C ₂ H ₅) ₂ O	192.4	805.	200.2	838.
Q	HCO ₂ C ₂ H ₅	185.3	775.	193.1	808.
R	1,2-C ₆ H ₄ (CH ₃) ₂	186.1	779.	193.3	809.
S	CO ₂	124.4	520.	130.9	548.
T	CH ₃ CHO	178.6	747.	186.6	781.
U	(CH ₃) ₂ CO	188.9	790.	196.7	823.
V	CH ₃ COOCH ₃	190.0	795.	197.8	828.
W	C ₆ H ₅ CH ₃	182.0	761.	189.8	794.
X	CH ₃ COOC ₂ H ₅	192.9	807.	200.7	840.
Y	H ₂ S	162.8	681.	170.2	712.
Z	CO	134.4	562.	141.9	593.
AA	CH ₃ CH=CH ₂	171.7	718.	179.5	751.
BB	H ₂	94.6	396.	101.3	424.
EE	(CH ₃) ₃ P	219.3	917.5	227.1	950.
XX	(n-C ₃ H ₇) ₂ O	194.5	814.	202.3	846.
ZZ	(CH ₃) ₆ C ₆	200.0	837.	207.3	867.

(Key) — Identity of reference base or other experimental details (in the case of hydride transfer equilibrium constant determinations, for instance) are given in a comment under the appropriate literature citation. (In the case of proton transfer equilibria, usually used for small data sets which were experimentally related to unusual reference bases.)

(br) — Bracketing result.

* — Asterisk appearing after gas basicity value indicates that original authors reevaluated the original data in a subsequent publication, but without giving

sufficient information for relating the reevaluation to the present scale. Data as cited are the reevaluated values of the authors.

** — Double asterisks appearing after gas basicity or proton affinity values indicate that insufficient information is available to evaluate the data. Proton affinity values are given as reported by the original authors, and are not necessarily internally consistent with the evaluated proton affinity scale.

RN — CAS Registry Number.

5.2. Explanation of Table 2

Table 2 presents the evaluations from Table 1 sorted according to the empirical formula of the appropriate compound. The empirical formulas are given at the beginning of each new data item enclosed in square brackets; component atoms are listed alphabetically, with no concessions to chemical meaning (except that lower case letters are given a lower priority than upper case letters), and the compounds are given simply according to an alphabetical ordering of these empirical formulas. Table 2 also lists the heat of formation at 298 K of the compound, and a 298 K heat of formation of the protonated molecule derived from the evaluated proton affinity, using the "stationary electron convention".

5.2.1. Conventions Used

As mentioned above, the heats of formation of protonated molecules given in Table 2 are given according to the so-called "stationary electron" convention¹². According to this convention, the integrated heat capacity of the electron in eqs. (5) and (6) is taken as zero at all temperatures, and the thermal energy of the electron is not taken into account in calculating the heat of formation of the ion at temperatures other than zero kelvin. Obviously, since ionic species occur on both sides of equation (1), the absolute value for the proton affinity does not change with the convention, but one must take care to use consistent data in calculating a proton affinity from ionic heats of formation. If the heats of formation of ions given here are to be used in conjunction with heats of formation of ions taken from a compilation of thermodynamic data (such as "The NBS Tables of Chemical Thermodynamic Properties: Selected Values for Inorganic and C₁ and C₂ Organic Substances in SI Units"⁵⁹ or the "JANAF Thermochemical Tables"⁶¹) in which the electron is treated as a conventional chemical element, the 298 K values given here must be increased by 1.48 kcal/mol or 6.2 kJ/mol in order to achieve consistency. The existence of two different conventions does not matter for neutral species. For the defining eq. (1), the 298 K heat of formation of the proton using this convention is 1530.0 kJ/mol or 365.7 kcal/mol. Relevant heats of formation of reference ions used to standardize the scale were taken at 298 K in all cases where sufficient information was available that this was possible.

5.2.2. Heats of Formation of Neutral Molecules

Heats of formation of the neutral molecule, M, in eq. (1) are listed in Table 2. The experimental data in Table 1 lead to a value for the proton affinity of M, and if the heat of formation of M is known or, as described below, can be estimated, one can thereby obtain a value

for the heat of formation of the protonated molecule MH⁺. In a few cases, a reliable value for the heat of formation of MH⁺ was available from other sources, and the proton affinity determination led to a determination of the heat of formation of the neutral molecule or radical. No distinction is made between these cases in Table 2, but in the latter case, the cited reference will be to the paper describing the proton affinity determination.

Values for the heats of formation of neutral molecules were taken from the experimental literature wherever possible. If data for a particular compound could be obtained from an evaluated data compilation, this value was selected for inclusion. Such compilations included, for organic compounds:

(1) J. B. Pedley and J. Rylance, "Sussex-N. P. L. Computer Analysed Thermochemical Data: Organic and Organometallic Compounds," University of Sussex (1977)³⁴. The numerous data from this evaluated compilation of 300 K heats of formation of organic compounds are identified as [77PED/RYL]. Where condensed phase data from this reference have been used in conjunction with experimental or estimated heats of vaporization or sublimation, the source of the heat of formation is identified simply by an asterisk (*) in the first space of the reference identification.

and for inorganic compounds (in order of preference):

(2) D. D. Wagman, W. H. Evans, V. B. Parker, R. H. Schumm, I. Halow, S. M. Bailey, K. L. Churney, and R. L. Nuttall, "The NBS Tables of Chemical Thermodynamic Properties: Selected Values for Inorganic and C₁ and C₂ Organic Substances in SI Units," J. Phys. Chem. Ref. Data, Vol. 11, Suppl. 2 (1982), hereafter referred to as 82/TN270 from the original publication of this compilation as a series of NBS Technical Notes called the 270-series⁵⁹.

(3) (a) D. R. Stull and H. Prophet, "JANAF Thermochemical Tables," NSRDS-NBS 37 (1971); (b) M. W. Chase, J. L. Curnutt, H. Prophet, R. A. McDonald, and A. N. Syverud, "JANAF Thermochemical Tables, 1975 Supplement," J. Phys. Chem. Ref. Data 4, 1 (1975); (c) M. W. Chase, Jr., J. L. Curnutt, J. R. Downey, Jr., R. A. McDonald, A. N. Syverud, and E. A. Valenzuela, J. Chem. Phys. Ref. Data 11, 695 (1982). Information from these sources is hereafter referred to as JANAF⁶¹.

The preferential use of these compilations as sources of experimental data recognizes that these data have been evaluated for internal consistency insofar as possible. Experimental information about heats of formation of compounds not included in these sources was obtained

from scientific articles presenting relevant thermochemical determinations; when more than one such article was available for a given piece of data, the most recent value was usually arbitrarily chosen in the possibly naive hope that improved instrumentation and compound purity, as well as a greater awareness of the problems inherent in particular measurements or of the thermochemistry of particular classes of compounds, should contribute to improved values. In such cases, it should be noted, the cited references are not always the primary references for the determination, but may represent a choice by a subsequent author. Heats of formation of deuterated or partially deuterated molecules were assumed to be the same as the heats of formation of the non-deuterated analogues. Also, gas phase heats of formation of racemic mixtures are the same as the heats of formation of the optically pure enantiomers.

Implicitly, in assigning gas phase heats of formation to the neutral species, the compounds are assumed to be ideal gases at STP. Few of the species are, in fact, gases under these conditions — most are liquids or solids. Numerous methods exist for measuring and interpreting experimental heats of sublimation [$\Delta H(\text{solid} \rightarrow \text{gas})$] and of vaporization [$\Delta H(\text{liquid} \rightarrow \text{gas})$], and, so need not be discussed here. Where available, such measurements were used to translate condensed phase data into gas phase information. However, such measurements do not exist for many compounds of interest here, either because of lack of adequate volatility, purity, or thermal stability, or even, seemingly because of a lack of interest in the species in the gas phase. Estimation methods for heats of vaporization, and occasionally sublimation, have been described in the literature. Some require other experimental data (e.g., critical constants), while others are limited to well-defined, however large, classes of compounds. Some of these methods have been used to generate values of gas phase molecular heats of formation from corresponding liquid or solid phase data; although such heats of formation are based on an estimation of one parameter, comparisons indicate that the resulting gas phase data are usually within 4 kJ/mol (1 kcal/mol) of accepted values, at least for heats of vaporization. Therefore, these entries are not labelled "Est" for "Estimate" (see below), but are rather identified by a two-part reference, the first segment giving the source of the liquid or solid phase heat of formation of the compound (abbreviated to an asterisk (*) when that source is the often-referred to [77PED/RYL]), and the second segment, listing a reference describing the technique of determining the heat of vaporization or sublimation.

Heats of formation of compounds for which no experimental data were available were estimated, and are labelled "Est". Several estimation approaches were utilized. Only brief descriptions of these will be given here; a detailed discussion will be presented elsewhere⁶².

One approach utilizes experimental information

about isomerization processes for the formation of two or more isomeric species in a particular reaction. The assumption is made that $\Delta\Delta_f H = \Delta\Delta_f G$ for a pair of isomers (ΔS is assumed negligible) in the experimentally observed process, so that a heat of formation of an unknown species can be predicted from thermochemical information about isomeric molecules. This approach, which is not well characterized and so is less trustworthy than other approaches used here, utilizes various types of information including direct equilibration studies on isomers and the more casual observation that two isomers are formed in comparable quantities in a particular process, and so have comparable Gibbs free energies and enthalpies of formation.

A related approach to estimation is based on the assumption that $\Delta\Delta_f H$ can be equated with ΔE_{tot} for two isomers, where ΔE_{tot} is the total energy of the molecule calculated quantum chemically. For this assumption, as well as all other estimation approaches employing quantum chemical studies, only ab initio calculations were considered as opposed to any of the plethora of semi-empirical studies at a wide variety of levels and approximations to the correct Hamiltonian and wavefunction. In all cases, care was taken to compare isomeric species with the same basis set and degree of geometry optimization. Built into this last assumption that $\Delta\Delta_f H = \Delta E_{\text{tot}}$ are the requirements that the zero point energy and enthalpy function $H^0 - H_0^0$ are essentially equal for an arbitrary pair of isomers, and that conformational isomerism and relative energies hardly affect heats of formation. None of these assumptions is rigorously true, but experience indicates that they represent a good first guess as to molecular energetics.

There remain three related approaches which were employed to estimate heats of formation of molecules here. The first and best characterized is Benson's "group increment" approach^{63,64} in which the molecule of interest is defined as a collection of groups, and a "group", in turn, is defined as a polyvalent atom (ligancy ≥ 2) with all its ligands in a molecule. The heat of formation of the molecule is obtained by summing statistically-determined contributions from the heats of formation of the various "groups", and correcting for various higher order interaction and other "correction" terms. These corrections include such things as the presence of gauche configurations in substituted alkanes, and the presence of heterocyclic and/or non-six membered rings.

A related approach consists of formulating the molecule of interest as a substituted derivative of a well-characterized species. The unknown heat of formation is taken to be the sum of the known heat of formation and a suitable correction term associated with the exchange of the substituent and parent components. For example, the heat of formation of an arbitrary aliphatic azoxy compound may be determined⁶⁵ from the heat of formation of the related (E)-olefin by:

$$\Delta_f H(\text{R}-\text{NN}(\text{O})-\text{R}') = \Delta_f H(\text{R}-\text{CH}=\text{CH}-\text{R}') + 14.6 \text{ kcal/mol}, \quad (23)$$

where the 14.6 kcal/mol for the $[\Delta_f H(-\text{NN}(\text{O})-) - \Delta_f H(-\text{CH}=\text{CH}-)]$ correction term was obtained by averaging the difference of the heats of formation of $\text{R}-\text{NN}(\text{O})-\text{R}'$ and $(\text{E})\text{R}-\text{CH}=\text{CH}-\text{R}'$ for $\text{R}=\text{R}'=\text{tert-butyl}$ and $n\text{-propyl}$.

The final estimation approach, termed "macro-incrementation reactions"⁶⁶, assumes that "if for each of two sets of molecules the total number of bonds, atoms and structural types is the same, then the total heat of formation of each set of molecules is the same. Then, if all but one of the heats are available, the remaining one can be estimated by simple arithmetic. It deals with ring strain, resonance energy and other interactions much more explicitly but less universally by embedding these corrections in the heat of formation of the individual molecules which among them possess the desired bonding characteristics. Further, it maximizes the direct use of available experimental data."⁶⁶

Quite clearly, the last three approaches described here are interrelatable, and "were they flawless, they would agree with each other and with experiment"⁶⁷. Furthermore, macroincrementation considerations, when coupled with the earlier assumption, $\Delta\Delta_f H = \Delta E_{\text{tot}}$, result in "isodesmic reactions"⁶⁸ and "group separation reactions"⁶⁹, widely employed here and elsewhere.

5.2.3. References

The reference citations given in Table 2 refer to the sources of data on the heats of formation of the neutral molecules. The references having to do with proton affinity determinations are given at the end of Table 1. In the column labelled "Reference" in Table 2, the appropriate literature reference is given as an eight-character identifier showing the year the paper appeared and the first three letters of the names of the first two authors. In addition, as discussed above in part, the following specialized notations and abbreviations are used:

Notation	Definition
82/TN270	D. D. Wagman, W. H. Evans, V. B. Parker, R. H. Schumm, I. Halow, S. M. Bailey, K. L. Churney, and R. L. Nuttall, "The NBS Tables of Chemical Thermodynamic Properties: Selected Values for Inorganic and C ₁ and C ₂ Organic Substances in SI Units," <i>J. Phys. Chem. Ref. Data</i> 11 , Suppl. 2 (1982).
JANAF	(a) D. R. Stull and H. Prophet, "JANAF Thermochemical Tables," NSRDS-NBS 37 (1971). (b) M. W. Chase, J. L. Curnutt, H. Prophet, R. A. McDonald, and A. N. Syverud, "JANAF Thermochemical Tables," 1975 Supplement, <i>J. Phys. Chem. Ref. Data</i> 4 , 1 (1975); (c) M. W. Chase, Jr., J. L. Curnutt, J. R. Downey, Jr., R. A. McDonald, A. N. Syverud, and E. A. Valenzuela, <i>J. Phys. Chem. Ref. Data</i> 11 , 695 (1982).

Notation	Definition
*00ABC/DEF	A condensed phase heat of formation was obtained from 77PED/RYL (J. B. Pedley and J. Rylance, "Sussex-N. P. L. Computer Analysed Thermochemical Data: Organic and Organometallic Compounds," University of Sussex (1977)) and translated into a gas phase heat of formation using heat of vaporization or sublimation data from the reference 00ABC/DEF.
Est	The gas phase heat of formation was estimated using the approaches described in the previous section.
DEF	A defined heat of formation.
(E) or (Z)	In the absence of other information, the most stable isomeric configuration (usually E) has been assumed.

6. Literature Coverage

Data from literature which appeared before June 1983, are included. A number of researchers who were aware that this compilation was in progress made available to the authors preprints of work which was in press or in preparation, and these more recent data are also included. Older data, which have been superseded by newer, better, measurements have been omitted. These include primarily bracketing measurements which have been replaced by results from equilibrium studies. In some cases where an author has repeated his own measurements in a later study and obtained different results, both sets of data are included, with preference generally given to the more recent set in carrying out the evaluation. Exceptions to this policy are specifically mentioned under the description of the evaluation of a paper in the annotated reference. Particularly in laboratories which have been very active in determinations of proton affinities from ion-molecule equilibrium constant measurements, one sees that closely similar or identical results on a particular molecule or set of molecules often appear in various publications from the laboratory without specific reference to earlier appearances of the data in print. In these cases, we do not presume to try to judge who the "original" author was, but simply present all the reincarnations of the data set.

In a few cases, authors have bracketed the gas phase basicities of molecules between limits which are so widely spaced (i.e., ammonia and water) that the result is difficult to present in the more quantitative scale given here; in such cases, the paper is listed in the bibliography, but the result is not included in the Tables 1 and 2. Other authors have given only an upper or lower limit to a gas basicity; these results have not been included.

Acknowledgements

As specifically noted, Mahnaz Motevalli-Aliabadi collaborated in the preparation of Table 2, carrying out

estimates of the heats of formation of neutral molecules. In addition, the authors gratefully acknowledge the work of Carol Martin and Clairemarie Lanthier who critically proofread the Tables, and Ricardo Metz, Nathan Seidenman, Carlo Messina, and Dr. Thomas J. Buckley who carried out the computer processing leading to the production of the final manuscript.

We would also like to acknowledge those colleagues, notably Dr. Michael Mautner, who made available to the authors experimental data which was in various stages of preparation, so that the compilation could be as complete and up-to-date as possible.

One of us (J. F. L.) acknowledges partial support from the U. S. Department of Energy, Office of Health and Environmental Research.

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Table 1. Gas phase basicities and proton affinities

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₁₄ H ₁₈ N ₂]	N,N,N',N'-Tetramethyl-1,8- naphthalenediamine	RN 20734-58-1		234.8	982.	241.8	1012	
320	ZZ	(34.9)	234.9					83TAF
600	A	(39.2)	234.8					78LAU/SAL
[C ₈ H ₂₀ N ₂]	(CH ₃) ₂ N(CH ₂) ₄ N(CH ₃) ₂	RN 111-51-3		232.6	973.	240.4	1006.	
320	A	(37.9)	233.5					TAFT
300			231.7**					79AUE/BOW
[C ₁₄ H ₂₇ N]	1-Methyl-2,6-t-butylpiperidine	RN xxxxx		231.4	968.	239.2	1001.	
320	ZZ	(31.4)	231.4					83TAF
[C ₇ H ₁₈ N ₂]	(CH ₃) ₂ N(CH ₂) ₃ N(CH ₃) ₂	RN 110-95-2		231.0	967.	238.8	999.	
320	A	(36.2)	231.8					TAFT
300			230.2**					79AUE/BOW
[C ₅ H ₁₄ N ₂]	1,5-Diaminopentane	RN 462-94-2		223.0	933.	238.1	996.	
298	A	(30.4)	225.0					73YAM/KEB
300	C	(12.9)	223.0					73AUE/WEB
300			221.9*					79AUE/BOW
600	A	(26.2)	221.3					78LAU/SAL
[C ₇ H ₁₈ N ₂]	1,7-Diaminoheptane	RN 646-19-5		224.2	938.	238.	996.	
298	E	(6.9)	224.2					73YAM/KEB
[C ₁₀ H ₂₄ N ₂]	(CH ₃) ₂ N(CH ₂) ₆ N(CH ₃) ₂	RN 111-18-2		230.1	963.	237.9	995.	
320	A	(35.3)	230.9					TAFT
300			229.3**					79AUE/BOW
[C ₆ H ₁₆ N ₂]	1,6-Diaminohexane	RN 124-09-4		223.0	933.	237.7	994.5	
300	C	(12.9)	223.0					73AUE/WEB
300			221.9**					79AUE/BOW
[C ₄ H ₁₂ N ₂]	1,4-Diaminobutane	RN 110-60-1		225.0	941.	237.6	994.	
330	D	(12.2)	225.3					80MAU/HAM
300	C	(14.6)	224.7					73AUE/WEB
300			223.9*					79AUE/BOW
[C ₅ H ₁₄ N ₂]	(CH ₃) ₂ N(CH ₂) ₃ NH ₂	RN 109-55-7		229.4	959.	237.2	992.	
320	A	(33.8)	229.4					TAFT
300			228.0**					79AUE/BOW
[C ₆ H ₁₃ N]	(CH ₃) ₂ NC(CH ₃)=CHCH ₃	RN 52113-79-8		~229	~958	~237	~991	
	(br)		~229					81ELL/DIX
[C ₆ H ₁₆ N ₂]	(CH ₃) ₂ N(CH ₂) ₂ N(CH ₃) ₂	RN 110-18-9		228.6	956.5	236.4	989.	
320	A	(33.0)	228.6					75TAF
300			227.2**					79AUE/BOW
[C ₇ H ₁₅ N]	(CH ₃) ₂ NC(C ₂ H ₅)=CHCH ₃	RN 78733-73-0		228.6	956.	236.4	989.	
	(br)		228.6					81ELL/DIX

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₇ H ₁₀ N ₂]	N,N-Dimethyl-4-pyridinamine RN 1122-58-3			228.4	956.	236.2	988.	
320	A	(33.3)	228.9					TAFT
300			227.9*					76AUE/WEB (2)
[C ₁₃ H ₁₆ N ₂]	N,N,N'-Trimethyl-1,8-naphtha- lenediamine RN 20723-57-0			227.8	953.	235.6	986.	
600	A	(32.2)	227.8					78LAU/SAL
[C ₈ H ₁₉ N]	(i-C ₃ H ₇) ₂ (C ₂ H ₅)N RN 7087-68-5			227.5	952.	235.3	984.	
320	A	(31.9)	227.5					TAFT
[C ₁₀ H ₁₆ N ₂]	1,2-(N(CH ₃) ₂) ₂ C ₆ H ₄ RN 704-01-8			227.4	951.	235.2	984.	
600	A	(31.8)	227.4					78LAU/SAL
[C ₉ H ₂₁ N]	(t-C ₄ H ₉)C(CH ₃) ₂ N(CH ₃) ₂ RN 3733-36-6			227.3	951.	235.1	984.	
320	A	(31.7)	227.3					78SHE/GOB
[C ₁₂ H ₂₇ N]	(n-C ₄ H ₉) ₃ N RN 102-82-9			227.0*	950.*	234.8*	982.*	
300	B	(21.3)	227.0*					79AUE/BOW
[C ₁₃ H ₂₅ N]	2,6-Di-t-butylpiperidine RN xxxxx			226.5	948.	234.3	980.	
320	ZZ	(26.5)	226.5					83TAF
[C ₈ H ₁₅ N]	1,4,4-Trimethyl-1,2,3,4-tetrahydropyridine RN 35079-50-6					234.2**	980.**	
								80HOU/VOG
[C ₃ H ₁₀ N ₂]	1,3-Diaminopropane RN 109-76-2			222.0	929.	234.1	979.	
298	A	(30.1)	225.7					73YAM/KEB
330	D	(8.8)	221.9					80MAU/HAM
300	C	(12.0)	222.1					73AUE/WEB
300			220.9*					79AUE/BOW
[C ₉ H ₂₁ N]	(n-C ₃ H ₇) ₃ N RN 102-69-2			226.2	946.	234.0	979.	
320	ZZ	(27.1)	227.1					83TAF
320	A	(31.3)	226.9					TAFT
320	A	(30.6)	226.2					75TAF-75ARN
300	B	(19.9)	225.6*					79AUE/BOW
300	B	(20.5)	226.2					72AUE/WEB
[C ₄ H ₁₁ NO]	NH ₂ (CH ₂) ₄ OH RN 13325-10-5			220.7	923.	233.8	978.	
	D	(7.6)	220.7					80MAU/HAM
[C ₁₃ H ₂₁ N]	2,6-Di-t-butylpyridine RN 585-48-4			224.7	940.	233.4	976.	
320	ZZ	(24.4)	224.4					83TAF
320	A	(28.8)	224.4					TAFT
320	A	(29.8)	225.4					75TAF-75ARN
320	A	(28.2)	223.8					75WOL/HAR
300			223*					76AUE/WEB (2)
425	D	(13.6)	226.7					83MAU/SIE
[C ₈ H ₁₉ N]	(t-C ₄ H ₉) ₂ NH RN 21981-37-3			225.4	943.	233.2	976.	
320	A	(29.8)	225.4					TAFT

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₉ H ₁₇ N]	1-Cyclopentylpyrrolidine		RN 18707-33-0			233.1**	975.**	79AUE/BOW
[C ₁₅ H ₁₈]	1,4-Dimethyl-7-isopropylazulene			225.	941.	233.	975.	
	RN 489-84-9							
	320 ZZ	(25.)	225.					77WOL/ABB
[C ₁₁ H ₁₇ N]	2,6-Diisopropylpyridine			225.1	942.	232.9	974.	
	RN 6832-21-9							
	425 D	(12.0)	225.1					83MAU/SIE
[C ₉ H ₁₉ N]	N-Isobutylpiperidine		RN 10315-89-6			232.9**	974.**	80HOU/VOG
[C ₉ H ₂₁ N]	(t-C ₅ H ₁₁)(t-C ₄ H ₉)NH		RN 58471-09-3	224.7*	940.	232.5*	973.	
	300 B	(19.0)	224.7*					79AUE/BOW
[C ₉ H ₁₈ N ₂]	1,5-Diazabicyclo[3.3.3]-undecane		RN 283-58-9		224.6	940.	232.4	972.
	320 A	(29.0)	224.6					81ALD/ARR
[C ₆ H ₁₅ N]	(C ₂ H ₅) ₃ N		RN 121-44-8		224.5	939.	232.3	972.
	320 A	(29.3)	224.9					83TAF
	320 A	(28.9)	224.5					72ARN/JON
	320 A	(28.5)	223.9					77STA/TAA-75TAF
	320 A	(28.8)	224.5					83MCI
	320 E	(7.3)	225.1					74STA/BEA (2)
	300 B	(17.7)	223.4*					79AUE/BOW
	300 B	(18.2)	223.9					72AUE/WEB
	550 C	(14.7)	224.8					79MAU
[C ₇ H ₁₃ N]	1-Azabicyclo[2.2.2]octane (Quinuclidine)		RN 100-76-5		224.3	938.	232.1	971.
	320 ZZ	(25.5)	225.5					83TAF
	320 A	(28.9)	224.5					77STA/TAA-75TAF
	300 B	(17.9)	223.6*					79AUE/BOW
	320 E	(7.7)	225.5					74STA/BEA (2)
						232.1**	971.**	80HOU/VOG
[C ₆ H ₁₅ N]	(CH ₃) ₂ (t-C ₄ H ₉)N		RN 918-02-5	224.2	938.	232.0	971.	
	320 A	(28.6)	224.2					TAFT
[C ₁₁ H ₁₃ N]	1,4-Dihydro-1,4-ethanoquinoline		RN 4363-25-1	224.2	938.	232.0	971.	
	320 A	(28.6)	224.2					TAFT
[C ₇ H ₁₇ N]	(C ₂ H ₅) ₂ (n-C ₃ H ₇)N		RN 4458-31-5	224.2**	938.**	232.0**	971.**	
	300 B	(18.5)	224.2*					79AUE/BOW
[C ₉ H ₁₇ NO ₂]	3,3-Dimethoxy-1-azabicyclo[2.2.2]octane		RN xxxxx	224**	937**	232**	971**	
	300		224**					79AUE/BOW
[C ₁₃ H ₉ N]	Acridine		RN 260-94-6	224.1	938.	231.9	970.	
	550 C	(14.0)	224.1					79MAU

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₆ H ₉ N ₃ O ₂]		L-Histidine	RN xxxxx		224.1	938.	231.9	970.	
		A	(28.5)	224.1					83MCI
[C ₇ H ₁₉ NSi]		(CH ₃) ₃ Si(CH ₂) ₂ N(CH ₃) ₂			224.0	937.	231.8	970.	
		RN 23138-94-5							
	320	A	(28.4)	224.0					78SHE/GOB
[C ₈ H ₂₁ NSi]		(CH ₃) ₃ Si(CH ₂) ₃ N(CH ₃) ₂			224.0	937.	231.8	970.	
		RN 28247-29-2							
	320	A	(28.4)	224.0					78SHE/GOB
[C ₆ H ₁₉ N]		(sec-C ₄ H ₉) ₂ NH	RN 626-23-3		223.6	935.5	231.8	970.	
		B	(17.0)	223.6					72AUE/WEB
[C ₇ H ₁₄ N ₂]		3-Amino-1-azabicyclo[2.2.2]octane			224.0**	937.**	231.8**	970.**	
		RN 6238-14-8							
	300			224.0**					79AUE/BOW
[C ₈ H ₁₅ N]		3-Methyl-1-azabicyclo[2.2.2]octane			223.9**	937.**	231.7**	969.**	
		RN 695-88-5							
	300			223.9**					79AUE/BOW
[C ₆ H ₁₅ P]		(C ₂ H ₅) ₃ P	RN 554-70-1		223.9**	937.**	231.7**	969.**	
	300			223.9**					79AUE/BOW
[C ₉ H ₁₉ N]		2,2,6,6-Tetramethyl- piperidine	RN 768-66-1		223.9**	937.**	231.7**	969.**	
	300			223.9**					79AUE/BOW
[C ₆ H ₁₇ NSi]		(CH ₃) ₃ SiCH ₂ N(CH ₃) ₂	RN 18182-40-6		223.6	936.	231.5	968.	
	320	A	(28.0)	223.6					TAFT
	320	A	(27.9)	223.5					78SHE/GOB
[C ₁₃ H ₂₁ N]		Pyridine,2,4-di- t-butyl	RN 29939-31-9		223.6*	935.5**	231.4**	968.**	
	300			223.6*					76AUE/WEB(2)
[C ₉ H ₁₃ N]		2,6-Diethylpyridine	RN 935-28-4		223.3	934.	231.1	967.	
	425	D	(10.2)	223.3					83MAU/SIE
[C ₈ H ₁₃ N]		1-Azabicyclo[2.2.2]- oct-2-ene,3-methyl	RN xxxxx		223.2**	934.**	231.0**	966.5**	
	300			223.2**					79AUE/BOW
[C ₆ H ₁₅ NO]		NH ₂ (CH ₂) ₆ OH	RN 4048-33-3		216.0**	904.**	231.0**	966.5**	
	300			216.0**					79AUE/BOW
[C ₈ H ₁₇ N]		1,4,4-Trimethylpiperidine	RN 1003-84-5				230.8**	966.**	
									80HOU/VOG
[C ₁₂ H ₂₁ N]		(CH ₂ =C(CH ₃)CH ₂) ₃ N	RN xxxxx		222.9**	932.6**	230.7**	965.**	
	300			222.9**					79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol		Proton affinity kJ/mol	Reference
[C ₉ H ₁₇ N]		c-C ₅ H ₁₀ NCH=C(CH ₃) ₂	RN 673-33-6		222.9**	932.6**	230.7**	965.**
	300			222.9**				79AUE/BOW
[C ₈ H ₁₉ N]		(CH ₃) ₃ C(CH ₂) ₂ N(CH ₃) ₂	RN 15673-04-8		222.6	931.	230.4	964.
	320	A	(27.0)	222.6				78SHE/GOB
[C ₁₃ H ₁₃ P]		(C ₆ H ₅) ₂ (CH ₃)P	RN 1486-28-8		222.5	931.	230.3	963.5
	320	EE	(3.2)	222.5				82IKU/KEB
[C ₆ H ₁₄ N ₂ O ₂]		L-Lysine	RN 56-87-1		222.5	931.	230.3	963.5
		A	(26.9)	222.5				83MCI
[C ₆ H ₁₅ N]		(i-C ₃ H ₇) ₂ NH	RN 108-18-9		222.0	929.	230.2	963.
	320	A	(26.4)	222.0				TAFT
	320	A	(25.5)	221.1				75ARN
		A	(26.4)	222.0				83MCI
	300	B	(15.0)	220.7*				79AUE/BOW
	300	B	(16.2)	221.9				72AUE/WEB
[C ₁₀ H ₁₉ N]		1-Azabicyclo[3.3.3]undecane (Manxine)	RN 31023-92-4		222.3	930.	230.1	963.
	300	B	(16.6)	222.3				75AUE/WEB
	300			223**				79AUE/BOW
[C ₈ H ₁₃ N]		1-Azabicyclo[2.2.2]- octane,3-methylene	RN 22207-84-7		222.3**	930.**	230.1**	963.**
	300			222.3**				79AUE/BOW
[C ₅ H ₁₃ N]		(CH ₃)(C ₂ H ₅) ₂ N	RN 616-39-7		222.2	930.	230.0	962.
	320	A	(26.9)	222.5				TAFT
	320	A	(26.3)	221.9				75TAF-75ARN
[C ₁₂ H ₂₄ O ₆]		1,4,7,10,13,16-Hexaoxa- cyclooctadecane (18-Crown-6)	RN 17455-13-9		216.0	904.	230.0	962.
	300	(Key)		211.3				83MAU
	300	A	(20.4)	216.0				84SHA/BLA
[C ₁₂ H ₁₄ N ₂]		N,N'-Dimethyl-1,8-naphthalene- diamine	RN 20734-56-9		223.0	933.	230.0	962.
	600	A	(27.9)	223.0				78LAU/SAL
[C ₉ H ₁₅ N]		(CH ₂ =CHCH ₂) ₃ N	RN 102-70-5		222.2	930.	230.0	962.
	320	A	(27.0)	222.6				TAFT
	320	A	(26.4)	222.0				75TAF-75ARN
	300			221.5**				79AUE/BOW
[C ₅ H ₆ N ₂]		4-Pyridinamine	RN 504-24-5		222*	929*	230*	962*
	300			222*				76AUE/WEB (2)
[C ₁₈ H ₁₅ P]		(C ₆ H ₅) ₃ P	RN 603-35-0		222.5	931.	230.	962.
	320	EE	(3.2)	222.5				82IKU/KEB
[C ₇ H ₁₇ N]		(CH ₃) ₂ (neo-C ₅ H ₁₁)N	RN 10076-31-0		222.1	929.	229.9	962.
	320	A	(26.6)	222.1				TAFT
	320	A	(26.5)	222.0				78SHE/GOB

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	kJ/mol	Proton affinity kcal/mol	kJ/mol	Reference
[C ₇ H ₁₀ N ₂]		N,N-Dimethyl-3-pyridinamine			222.1**	929.**	229.9**	962.**	
		RN 18437-57-5							
	300			222.1**					76AUE/WEB (2)
[C ₅ H ₁₃ N]		(CH ₃) ₂ (i-C ₃ H ₇)N		RN 996-35-0	222.0	929.	229.8	961.	
	320	A	(26.9)	222.5					TAFT
	300	B	(15.4)	221.1*					79AUE/BOW
[C ₈ H ₂₁ NSi]		(CH ₃) ₂ (t-C ₄ H ₉)SiN(CH ₃) ₂		RN 66365-05-7	221.9	928.	229.7	961.	
	320	A	(26.3)	221.9					78SHE/GOB
[C ₆ H ₁₃ N]		1-Methylpiperidine		RN 626-67-5	221.9	928.	229.7	961.	
	320	A	(26.8)	222.4					TAFT
	320	A	(27.4)	223.0					75ARN
	300	B	(15.3)	221.0					76AUE/WEB
[C ₈ H ₁₁ P]		C ₆ H ₅ P(CH ₃) ₂		RN 672-66-2	221.8	928.	229.6	961.	
	320	EE	(2.5)	221.8					82IKU/KEB
[C ₅ H ₁₁ N]		(CH ₃) ₂ C=NC ₂ H ₅		RN 15673-04-8	221.7**	927.5**	229.5**	960.**	
	300			221.7**					79AUE/BOW
[C ₆ H ₁₃ N]		(CH ₃) ₂ C=CHN(CH ₃) ₂		RN xxxxx	221.7	928.	229.5	960.	
		(br)		221.7					81ELL/DIX
[C ₅ H ₁₁ N]		CH ₃ CH=CHN(CH ₃) ₂		RN 6163-56-0	221.7	928.	229.4	960.	
		(br)		221.7					81ELL/DIX
[C ₁₂ H ₁₉ N]		C ₆ H ₄ N(CH ₃) ₂ ,2-t-C ₄ H ₉		RN 22025-87-2	221.5	927.	229.3	959.	
	320	A	(25.9)	221.5					TAFT
[C ₇ H ₁₀ N ₂]		N,N-Dimethyl-2-pyridinamine		RN 5683-33-0	221.4*	926.*	229.2*	959.*	
	300			221.4*					76AUE/WEB (2)
[C ₆ H ₁₂ N ₂]		1,4-Diazabicyclo[2.2.2]octane		RN 280-57-9	221.2	925.5	229.0	958.	
	320	A	(25.1)	220.6					75ARN
	320	E	(3.8)	221.6					74STA/BEA (2)
	300	B	(15.0)	220.7					76AUE/WEB
	300			220**					79AUE/BOW
[CH ₂ O]		HCOH		RN xxxxx	221.	925.	229.	958.	
		(br)		221.					82PAU/HEH (2)
[C ₄ H ₆ N ₂]		1-Methylimidazole		RN 616-47-7	221.1	925.	228.9	958.	
	320	A	(25.1)	220.7					TAFT
		(br)		221.4					81ELL/DIX
	600	(Key)		220.					83MAU
[C ₁₁ H ₁₇ N]		2-C ₆ H ₁₃ (c-C ₅ H ₄ N)		RN 1129-69-7	221.1	925.	228.9	958.	
	425	D	(8.0)	221.1					83MAU/SIE

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₁₁ H ₁₇ N]	3-CH ₃ C ₆ H ₄ N(C ₂ H ₅) ₂	RN 91-67-8		220.9	924.	228.9	958.	
320	I	(5.5)	220.9					LIA/JAC
[C ₅ H ₁₁ N]	N-Methylpyrrolidine	RN 120-94-5		220.9	924.	228.7	957.	
320	A	(25.7)	221.3					75TAF-75ARN
300	B	(14.3)	220.0					76AUE/WEB
	D	(8.3)	221.4					73TAF/TAA-78TAA/WOL
[C ₃ H ₉ NO]	NH ₂ (CH ₂) ₃ OH	RN 156-87-6		217.3	909.	228.6	956.5	
330	D	(4.2)	217.3 216.0**					80MAU/HAM 79AUE/BOW
[C ₈ H ₁₉ N]	(i-C ₄ H ₉) ₂ NH	RN 110-96-3		220.4	922.	228.6	956.	
300	B	(14.7)	220.4*					72AUE/WEB
320	A	(25.2)	220.4					75ARN
[C ₁₁ H ₁₇ N]	4-CH ₃ C ₆ H ₄ N(C ₂ H ₅) ₂	RN 613-48-9		220.6	923.	228.6	956.	
	I	(5.2)	220.6					LIA/JAC
[C ₁₂ H ₁₉ N]	C ₆ H ₅ N(C ₃ H ₇) ₂	RN 2217-07-4		220.6	923.	228.6	956.	
320	I	(5.2)	220.6					LIA/JAC
[C ₇ H ₁₁ N]	1-Azabicyclo[2.2.2]oct-2-ene	RN 13929-94-7		220.7**	923.**	228.5**	956.**	
300			220.7**			229.3**	959.**	79AUE/BOW 80HOU/VOG
[C ₈ H ₁₉ N]	(n-C ₄ H ₉) ₂ NH	RN 111-92-2		220.3	922.	228.4	956.	
300	B	(14.5)	220.2*					76AUE/WEB
300	B	(14.6)	220.3					72AUE/WEB
320	A	(24.6)	220.2					75ARN
[C ₇ H ₉ N]	2,6-Dimethylpyridine	RN 108-48-5		220.4	922.	228.2	955.	
300			219.2*					76AUE/WEB (2)
320	A	(24.3)	219.9					75ARN
425	D	(7.8)	220.9					83MAU/SIE
[C ₆ H ₁₁ NO]	c-C ₅ H ₈ N(2-OCH ₃)	RN 53687-79-9		220.3*	922.*	228.1*	954.*	
300	B	(14.6)	220.3*					79AUE/BET
[C ₇ H ₁₂ N]	3-Fluoro-1-azabicyclo[3.2.1]octane	RN xxxxx		220.3**	922.**	228.1**	954.**	
300			220.3**					79AUE/BOW
[C ₉ H ₁₃ N]	C ₆ H ₅ CH ₂ N(CH ₃) ₂	RN 103-83-3		220.3**	922.**	228.1**	954.**	
300			220.3**					79AUE/BOW
[C ₄ H ₉ N]	(CH ₃) ₂ NCH=CH ₂	RN 5763-87-1		220.0	920.	227.8	953.	
	(br)		220.0					81ELL/DIX
[C ₉ H ₁₁ N]	2,3-Cyclohexenopyridine	RN 10500-57-9		219.9**	920.**	227.7**	953.**	
300			219.9**					79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₉ H ₁₁ N]	3,4-Cyclohexenopyridine RN 36566-06-6			219.9**	920.**	227.7**	953.**	
300			219.9**					79AUE/BOW
[C ₇ ClH ₁₄ N]	c-C ₅ H ₉ N,2-CH ₂ Cl,1-CH ₃ RN 49665-74-9			219.8**	920.**	227.6**	952.**	
300			219.8**					79AUE/BOW
[C ₁₀ H ₁₅ N]	C ₆ H ₅ N(C ₂ H ₅) ₂ RN 91-66-7			219.6	919.	227.6	952.	
600	A	(26.9)	222.0					73YAM/KEB
325	I	(4.2)	219.6					LIA/JAC
[C ₆ H ₇ NO]	4-Methoxypyridine RN 620-08-6			219.8	920.	227.6	952.	
320	D	(8.7)	221.7					72TAA/HEN
320	A	(24.8)	220.4					81TAA/SUM
320	A	(24.2)	219.8					75TAF-75ARN
300			218.8*					76AUE/WEB (2)
[C ₄ H ₁₁ N]	(CH ₃) ₂ (C ₂ H ₅)N RN 598-56-1			219.7	919.	227.5	952.	
320	A	(24.5)	220.1					TAF ^T
320	A	(23.9)	219.5					75TAF-75ARN
300	B	(13.1)	218.8*					76AUE/WEB
[C ₆ H ₁₅ N]	(n-C ₃ H ₇) ₂ NH RN 142-84-7			219.7	919.	227.5	952.	
320	A	(24.3)	219.9					75TAF-83TAF
300	B	(13.5)	219.2*					79AUE/BOW
300	B	(13.8)	219.5					72AUE/WEB
535	C	(9.6)	219.7					79MAU
[C ₉ H ₁₃ N]	2-t-Butylpyridine RN 5944-41-2			219.6**	919.**	227.4**	951.**	
300			218.6**					79AUE/BOW
425	D	(7.4)	220.5					83MAU/SIE
[C ₅ H ₁₃ N]	(C ₂ H ₅)(i-C ₃ H ₇)NH RN 19961-27-4			219.4	918.	227.4	951.	
320	A	(23.8)	219.4					TAF ^T
[C ₇ H ₉ N]	2,4-Dimethylpyridine RN 108-47-4			219.5*	918.*	227.3*	951.*	
300			219.5*					76AUE/WEB (2)
[C ₈ H ₁₁ N]	2-Isopropylpyridine RN 75981-47-4			219.4	918.	227.2	951.	
425	D	(6.3)	219.4					83MAU/SIE
[C ₁₀ H ₂₂ O ₅]	CH ₃ (OCH ₂ CH ₂) ₄ OCH ₃ RN 143-24-8			213.2	892.	227.2	951.	
300	A	(17.6)	213.2					84SHA/KEB
[C ₃ H ₉ P]	(CH ₃) ₃ P RN 594-09-2			219.3	917.5	227.1	950.	
320	EE	(0.0)	219.3					82IKU/KEB
320	E	(1.6)	219.3					74STA/BEA
320	A	(23.4)	219.0					75TAF
[BrC ₇ H ₁₂ N]	3-Bromo-1-azabicyclo[2.2.2]- octane RN xxxxx			219.3**	917.5**	227.1**	950.**	
300			219.3**					79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	kJ/mol	Proton affinity kcal/mol	kJ/mol	Reference
[C ₉ H ₁₃ N]	C ₆ H ₅ N(CH ₃)(C ₂ H ₅)	RN 613-97-8		219.3	917.5	227.1	950.	
600	A	(24.2)	219.3					73YAM/KEB-78LAU/SAL
[C ₁₀ H ₁₅ N]	3,5-(CH ₃) ₂ C ₆ H ₃ N(CH ₃) ₂	RN 4913-13-7		219.0	916.	227.0	950.	
320	I	(3.6)	219.0					LIA/JAC
[C ₈ H ₉ N]	3,4-Cyclopentenopyridine	RN xxxxx		219.0**	916.**	226.8**	949.**	
300			219.0**					79AUE/BOW
[C ₉ H ₇ N]	Quinoline	RN 91-22-5		218.7	915.	226.5	948.	
425	(Key)					225.8		81MCL/CAM
535	C	(8.6)	218.7					79MAU
[C ₃ H ₈ Si]	(CH ₃) ₂ Si=CH ₂	RN 4112-23-6		218.0	912.	226.4	947.	
320	A	(22.4)	218.0					82PIE/HEH
320	(br)		~219.0					79PIE/POL
[C ₅ H ₁₅ NSi]	(CH ₃) ₃ SiN(CH ₃) ₂	RN 18135-05-2				226.4	947.	
	(br)							83HEN/FRE
[C ₅ H ₁₁ N]	Piperidine	RN 110-89-4		218.2	913.	226.4	947.	
320	A	(22.3)	217.9					75TAF-75ARN-83TAF
300	E	(1.0)	218.8					71BOW/AUE
300	C	(8.3)	218.4					73AUE/WEB
300	B	(11.9)	217.4					75AUE/WEB (2)
300	B	(11.5)	217.2*					76AUE/WEB
600	A	(23.1)	218.2					78LAU/SAL
600	A	(24.5)	219.6					73YAM/KEB
[C ₃ H ₇ N]	CH ₂ =C(CH ₃)NH ₂	RN 4427-28-5		218.5	914.	226.3	947.	
	(Key)		218.5					81LELL/DIX
[C ₇ H ₉ N]	2-Ethylpyridine	RN 100-71-0		218.4	914.	226.2	946.	
300			217.1*					76AUE/WEB (2)
425	D	(5.3)	218.4					83MAU/SIE
[C ₇ H ₉ N]	2,3-Dimethylpyridine	RN 583-61-9		218.4*	914.*	226.2*	946.*	
300			218.4*					76AUE/WEB (2)
[C ₇ H ₉ N]	3,4-Dimethylpyridine	RN 583-58-4		218.4*	914.*	226.2*	946.*	
300			218.4*					76AUE/WEB (2)
[C ₆ H ₁₃ O ₃ P]	cis,cis-2-Methoxy-4,6-dimethyl-1,3,2-dioxaphosphorinane	RN 7735-82-2		218.4	914.	226.2	946.	
320	(Key)		218.4					80HOD/HOU
[C ₁₂ H ₂₄ N ₂]	1,6-Diazabicyclo[4.4.4]-tetradecane	RN 71058-67-8		218.9	916.	226.0	946.	
320	A	(23.3)	218.9					81ALD/ARR
[C ₇ H ₉ N]	2,5-Dimethylpyridine	RN 589-93-5		218.2*	913.*	226.0*	946.*	
300			218.2*					76AUE/WEB (2)

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₇ H ₉ NO]	Pyridine-2-methoxymethyl RN 23579-92-2			218.2**	913.**	226.0**	945.5**	
300			218.2**					79AUE/BOW
[C ₅ H ₉ NO]	c-C ₄ H ₆ N(2-OCH ₃) RN 5264-35-7			218.1	912.5	225.9	945.	
300	B	(12.4)	218.1					79AUE/BET
[C ₂ H ₈ N ₂]	1,2-Diaminoethane RN 107-15-3			219.2	917.	225.9	945.	
298	A	(24.1)	219.2					73YAM/KEB
330	D	(3.4)	216.6					80MAU/HAM
300	C	(6.5)	216.6					73AUE/WEB
[C ₄ H ₁₁ N]	(C ₂ H ₅) ₂ NH RN 109-89-7			217.7	911.	225.9	945.	
320	A	(22.0)	217.6					83TAF
320	A	(21.6)	217.2					75TAF-75ARN
320	A	(22.0)	217.6					72ARN/JON
300	B	(12.0)	217.7					72AUE/WEB
300	B	(11.7)	217.4					75AUE/WEB (2)
300	B	(11.2)	216.9*					76AUE/WEB
550	C	(7.6)	217.7					79MAU
[C ₉ H ₇ N]	Isoquinoline RN 119-65-3			218.1	912.	225.9	945.	
535	C	(8.0)	218.1					79MAU
[C ₉ H ₁₃ N]	4-t-Butylpyridine RN 3978-81-2			218.1	913.	225.9	945.	
300			218.1*					76AUE/WEB (2)
[C ₇ H ₇ N]	3,4-Cyclobutenopyridine RN xxxxx			218.1**	912.**	225.9**	945.**	
300			218.1**					79AUE/BOW
[C ₁₁ H ₁₅ N]	1-Phenylpiperidine RN 4096-2-2			219.5	918.	225.8	945.	
320	A	(23.9)	219.5					TAFT
[C ₈ H ₉ N]	2,3-Cyclopentenopyridine RN xxxxx			218.0**	912.**	225.8**	945.**	
300			218.0**					79AUE/BOW
[C ₇ ClH ₁₂ N]	3-Chloro-1-azabicyclo[2.2.2]- octane RN 42332-45-6			218.0**	912.**	225.8**	945.**	
300			218.0**					79AUE/BOW
[C ₉ H ₁₃ N]	4-CH ₃ C ₆ H ₄ N(CH ₃) ₂ RN 99-97-8			217.6	910.	225.6	944.	
320	I	(2.2)	217.6					LIA/JAC
[C ₁₀ ClH ₁₄ N]	4-ClC ₆ H ₄ N(C ₂ H ₅) ₂ RN 2873-89-4			217.8	911.	225.6	944.	
320	I	(-2.4)	213.0					LIA/JAC
[C ₇ H ₉ N]	3,5-Dimethylpyridine RN 591-22-0			217.7*	911.*	225.5*	943.*	
300	B	(12.0)	217.7*					76AUE/WEB
[C ₆ H ₇ NS]	4-(Methylthio)-pyridine RN 22581-72-2			217.7**	911.**	225.5**	943.**	
300			217.7**					79AUE/BOW
[C ₁₁ H ₁₂ N ₂ O ₂]	L-Tryptophan RN 54-12-6			217.6	910.	225.4	943.	
	A	(22.0)	217.6					83MCI

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol		kJ/mol	Proton affinity kcal/mol		kJ/mol	Reference	
[C ₆ H ₁₃ N]		n-C ₃ H ₇ CH=NC ₂ H ₅	RN 1611-12-7		217.5**	910.**		225.3**	943.**			
	300			217.5**							79AUE/BOW	
[C ₆ H ₇ N]		4-Methylpyridine	RN 108-89-4		217.4	909.		225.2	942.			
	320	D	(+5.4)	218.4							72TAA/HEN	
	320	A	(21.1)	216.7							83TAF	
	320	A	(21.4)	217.0							75TAF-75ARN	
[C ₄ H ₉ N]		Pyrrolidine	RN 123-75-1		217.3	909.		225.2	942.			
	320	A	(21.3)	216.9							83TAF-81TAA/SUM	
	320	A	(21.4)	217.0							75TAF-75ARN	
	300	B	(10.4)	216.1*							76AUE/WEB	
	320	E	(-0.3)	217.5							71BOW/AUE	
[C ₃ H ₉ N]		(CH ₃) ₃ N	RN 75-50-3		217.3	909.		225.1	942.			
	320	E	(0.0)	217.3							74STA/BEA	
	320	E	(0.0)	217.3							74STA/BEA (2)	
	300	E	(0.0)	217.3							71BOW/AUE	
	300	B	(11.5)	217.2							72AUE/WEB	
	300	B	(10.7)	216.4							75AUE/WEB (2)	
	300	B	(10.8)	216.5*							76AUE/WEB	
	320	A	(21.9)	217.4							TAFT	
	320	A	(21.4)	216.9							75TAF-75ARN	
	320	A	(21.8)	217.3							72HEN/TAA-72ARN/JON	
		A	(22.0)	217.6							83MCI-83TAF	
	330	D	(4.4)	217.5							80MAU/HAM	
	600	A	(23.3)	218.4							72BRI/YAM	
[C ₆ H ₇ N]		2-Methylpyridine	RN 109-06-8		217.2	909.		225.0	942.			
	320	A	(21.5)	217.1							TAFT	
				215.9*							76AUE/WEB (2)	
	425	D	(4.1)	217.2							83MAU/SIE	
[C ₆ H ₁₃ O ₃ P]		trans-2-Methoxy-cis,cis-4,6-dimethyl-1,3,2-dioxaphosphorinane	RN 41821-91-4		216	904		225	941			
	320	A		216							80HOD/HOU	
[C ₁₀ H ₁₃ N]		N-Phenylpyrrolidine	RN 4096-21-3		216.9	907.		224.7	940.			
	320	A	(21.3)	216.9							TAFT	
[C ₆ H ₁₁ N]		(CH ₂ =CHCH ₂) ₂ NH	RN 124-02-7		216.9	907.		224.7	940.			
	320	A	(21.3)	216.9							TAFT	
[C ₉ H ₇ NO]		Quinoline-1-oxide	RN 1613-37-2		216.8	907.		224.6	940.			
	526	C	(6.7)	216.8							79MAU	
[C ₇ H ₉ N]		4-Ethylpyridine	RN 536-75-4		216.8**	907.**		224.6**	940.**			
	300			216.8*							76AUE/WEB (2)	
[C ₉ H ₁₃ N]		3-CH ₃ C ₆ H ₄ N(CH ₃) ₂	RN 121-72-2		216.7	907.		224.5	939.			
	320	F	(14.3)	216.7							77POL/DEV	
[C ₆ H ₁₁ NO ₃]		CH ₃ CONHCH(CH ₃)COOCH ₃ (N-Acetyl L-alanine methyl ester)	RN xxxxx (Key)		211.	883.		224.5	939.			
					211.						83MAU	
[C ₄ H ₆ N ₂]		4-Methylimidazole	RN 822-36-6		216.6	906.		224.4	939.			
	600	(Key)		216.6							83MAU	

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₄ H ₁₀ N ₂]	Piperazine	RN 110-85-0		216.4 905.	224.2 938.	
300	C	(6.3)	216.4			73AUE/WEB
[C ₆ H ₇ N]	3-Methylpyridine	RN 108-99-6		216.2 905.	224.1 938.	
320	A	(20.6)	216.2			TAFT
300	B	(9.3)	215.0*			76AUE/WEB
			215.0*			76AUE/WEB (2)
600	D	(4.2)	217.3			83MAU
[C ₈ H ₁₈ O ₄]	CH ₃ (OCH ₂ CH ₂) ₃ OCH ₃	RN 112-49-2		210.8 882.	224.1 938.	
300	(Key)		~212			83MAU
300	A	(15.2)	210.8			84SHA/BLA
[C ₇ ClH ₁₀ N]	3-Chloro-1-azabicyclo[2.2.2]-oct-2-ene	RN xxxxx		216.2** 904.5**	224.0** 937.**	
			216.2**			79AUE/BOW
[C ₁₀ H ₁₇ NO]	cis-3-Amino-2-twistanol	RN xxxxx		216.2 904.5	224.0 937.	
	(Key)		216.2			83HOU/RUF
[C ₈ H ₁₅ NO]	cis-3-Aminobicyclo[2.2.2]octan-2-ol	RN 17997-65-8		216.1 904.	223.9 937.	
	(Key)		216.1			83HOU/RUF
[C ₃ H ₈ Pb]	(CH ₃) ₂ Pb=CH ₂	RN 82065-01-8		216.1 904.	223.9 937.	
320	A	(20.5)	216.1			82PIE/HEH
[C ₇ H ₉ N]	3-Ethylpyridine	RN 536-78-7		216.1* 904.*	223.9* 937.*	
			216.1*			76AUE/WEB (2)
[C ₁₀ H ₁₀ N ₂]	1,8-Diaminonaphthalene	RN 479-27-6		216.2 904.5	223.8 936.	
600	A	(21.1)	216.2			78LAU/SAL
[C ₄ H ₅ N ₃ O]	Cytosine	RN 71-30-7		216.0 904.	223.8 936.	
535	C (br)	(5.9)	216.0 ~215			79MAU 75WIL/MCC
[C ₅ H ₆ N ₂]	2-Pyridinamine	RN 504-29-0		216.0* 904.*	223.8* 936.*	
			216.0*			79AUE/BOW
[C ₁₂ H ₈ N ₂]	Phenazine	RN 92-82-0		216.6 906.	223.7 936.	
514	C	(6.5)	216.6			79MAU
[C ₁₀ H ₂₀ O ₅]	1,4,7,10,13-Pentaoxacyclopentadecane (15-Crown-5)	RN 33100-27-5		212.5 889.	223.6 935.	
300	(Key)		212.			83MAU
300	A		212.5			84SHA/BLA
[C ₆ H ₇ NO]	3-Methoxypyridine	RN 7295-76-3		215.7 902.	223.6 935.	
320	A	(20.9)	216.5 214.7*			TAFT 76AUE/WEB (2)
[C ₅ H ₅ N ₅]	Adenine	RN 73-24-5		215.7 902.	223.5 935.	
550	C (br)	(5.6)	215.7 ~215			79MAU 75WIL/MCC

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₃ H ₇ N]	Azetidine	RN 503-29-7		215.7 902.	223.5 935.	
	300 B	(9.1)	214.8			75AUE/WEB (2)
	300 B	(8.8)	214.5*			76AUE/WEB
	300 E	(-2.1)	215.3			71BOW/AUE
[C ₈ H ₁₁ N]	C ₆ H ₅ N(CH ₃) ₂	RN 121-69-7		215.4 901.	223.4 935.	
	320 I	(0.0)	215.4			LIA/JAC
	320 A	(19.9)	215.5			75TAF-83TAF
		(19.8)	215.4			83MCI
	320 F	(12.5)	215.4			77POL/DEV
	600 A	(21.7)	216.8			73YAM/KEB-78LAU/SAL
[C ₇ H ₇ N]	2,3-Cyclobutenopyridine	RN xxxxx		215.5** 902.**	223.3** 934.**	
	300		215.5**			79AUE/BOW
[C ₃ H ₉ NO]	CH ₃ OCH ₂ CH ₂ NH ₂	RN 109-85-3		212.3* 888.*	223.3* 934.*	
	300 C	(2.2)	212.3*			73AUE/WEB
[C ₇ H ₇ N]	4-Vinylpyridine	RN 100-43-6		215.4** 901.**	223.2** 934.**	
			215.4**			79AUE/BOW
[C ₈ H ₆ N ₂]	Cinnoline	RN 253-66-7		215.4 901.	223.2 934.	
	535 C	(5.3)	215.4			79MAU
[C ₁₀ H ₁₀ Ni]	Ni(C ₅ H ₅) ₂	RN 1271-28-9		216. 904.	223. 933.	
	320 (Key)		216.			76COR/BEA
	320 A	(19.1)	214.7			81STE/BEA
[C ₅ H ₅ N ₅ O]	Guanine	RN 73-40-5		~215 ~899	~223 ~933	
	(br)		~215			75WIL/MCC
[C ₆ H ₆ N ₄]	6-Methylpurine	RN 2004-03-7		~215 ~899	~223 ~933	
	(br)		~215			75WIL/MCC
[C ₃ H ₉ N]	(CH ₃)(C ₂ H ₅)NH	RN 624-78-2		215.1 900.	222.8 932.	
	320 A	(19.8)	215.3			TAFT
	320 A	(19.1)	214.7			75TAF-75ARN
	300 B	(9.1)	214.8*			76AUE/WEB
[C ₄ H ₉ N]	CH ₃ CH=NC ₂ H ₅	RN 1190-79-0		214.9 899.	222.7 932.	
	300 B	(9.2)	214.9			75AUE/WEB (2)
[C ₆ H ₈ N ₂]	1,3-C ₆ H ₄ (NH ₂) ₂	RN 108-45-2		214.7 898.	222.4 930.5	
	600 F	(12.2)	214.7			81LAU/NIS
	600 A	(19.7)	214.8			78LAU/SAL
[C ₉ H ₁₁ NO ₃]	L-Tyrosine	RN xxxxx		214.5 897.	222.3 930.	
	A	(18.9)	214.5			83MCI
[C ₅ H ₁₃ N]	t-C ₅ H ₁₁ NH ₂	RN 594-39-8		213.9* 895.*	222.3* 930.*	
	300 B	(8.2)	213.9*			76AUE/WEB

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₁₀ H ₁₉ NO]		4-Aminodecahydro-3-naphthalenol RN xxxxx			214.3 897.	222.1 929.	
		(Key)		214.3			83HOU/RUF
[C ₆ H ₇ NS]		2-(Methylthio)-pyridine RN 18438-38-5			214.2 896.	222.0 929.	
	300	B (8.5)		214.2 214.2*			79AUE/BET 76AUE/WEB (2)
[C ₇ H ₆ O]		4-Methylene-2,5-cyclohexadiene-1-one (br)			RN 502-87-4	-222** -929**	77DIT/NIB
[C ₇ H ₁₁ NO]		1-Azabicyclo[2.2.2]octan-3-one RN 3731-38-2			214.1** 896.**	221.9** 928.**	
				214.1**			79AUE/BOW
[C ₆ H ₇ NO]		2-Methoxypyridine RN 1628-89-3			214.1 896.	221.9 928.	
	320	A (18.8)		214.4			76COO/KAT
	300	B (7.8)		213.5 213.5*			79AUE/BET 76AUE/WEB (2)
[C ₆ H ₁₄ O ₂]		CH ₃ O(CH ₂) ₄ OCH ₃ RN 13179-96-9			209.0 874.	221.8 928.	
	300	A (13.4)		209.0			84SHA/BLA
[C ₇ H ₁₆ O ₂]		CH ₃ O(CH ₂) ₅ OCH ₃ RN 111-89-7			208.8 874.	221.8 928.	
	300	A (13.2)		208.8			84SHA/BLA
[C ₈ H ₁₁ N]		C ₆ H ₅ NHC ₂ H ₅ RN 103-69-5			214.0 895.	221.8 928.	
	600	A (18.4)		214.0			73YAM/KEB-78LAU/SAL
	425	(Key)		218.8			81MCL/CAM
[C ₇ F ₂ H ₁₁ N]		3,3-Difluoro-1-azabicyclo- [2.2.2]octane RN xxxxx			214.0** 895.**	221.8** 928.**	
				214.0**			79AUE/BOW
[C ₇ H ₁₃ N]		Bicyclo[2.2.1]heptan-2-amine, exo (2-Aminonorbornane) RN 7242-92-4			213.3** 892.**	221.7** 927.**	
				213.3**			79AUE/BOW
[C ₇ H ₁₃ N]		Bicyclo[2.2.1]heptan-2-amine, endo (2-Aminonorbornane) RN 31002-73-0			213.3** 892.**	221.7** 927.**	
				213.3**			79AUE/BOW
[C ₈ H ₁₆ O ₄]		1,4,7,10-Tetraoxa- cyclododecane (12-Crown-4) RN 294-93-9			211.3 884.	221.6 927.	
	300	(Key)		211.0			83MAU
	300	A		211.3			84SHA/BLA
[C ₃ H ₇ N]		N-Methylaziridine RN 1072-44-2			213.8 894.	221.6 927.	
	300	B (9.1)		213.8			75AUE/WEB (2)
[C ₁₀ H ₁₇ NO]		trans-3-Amino-2-twistanol (Key)			213.7 894.	221.5 927.	
				213.7			83HOU/RUF

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₅ H ₁₁ NO ₂ S]	L-Methionine	RN 59-51-8		213.6	894.	221.4	926.	
	A	(18.0)	213.6					83MCI
[C ₂ H ₇ NO]	NH ₂ (CH ₂) ₂ OH	RN 141-43-5		213.4	893.	221.3	926.	
330	D	(0.3)	213.4					80MAU/HAM
[C ₆ H ₁₃ N]	c-C ₆ H ₁₁ NH ₂	RN 108-91-8		213.4	893.	221.2	925.5	
320	A	(17.5)	213.1					75TAF-75ARN-83TAF
300	B	(7.2)	212.9*					76AUE/WEB
600	A	(19.5)	214.6					73YAM/KEB
[C ₅ H ₆ N ₂]	3-Aminopyridine	RN 462-08-8		213.2*	892.*	221.0*	925.*	
			213.2*					76AUE/WEB (2)
[C ₆ H ₁₄ OSi]	CH ₂ =C(CH ₃)OSi(CH ₃) ₃	RN 1833-53-0		213.	891.	221.	925.	
	(br)		213.					82HEN/WEI
[C ₄ H ₁₁ N]	t-C ₄ H ₉ NH ₂	RN 75-64-9		213.0	891.	220.8	924.	
320	A	(17.1)	213.0					75TAF-75ARN
320	A	(17.6)	213.2					72HEN/TAA-72ARN/JON
	A	(17.8)	213.4					83MCI-83TAF
300	B	(7.2)	212.9*					76AUE/WEB
300	B	(7.0)	212.7					72AUE/WEB
514	C	(3.3)	213.4					79MAU
550	A	(16.9)	212.1					80MAU
[C ₅ H ₅ N]	Pyridine	RN 110-86-1		213.1	892.	220.8	924.	
425	D	(0.0)	213.1					83MAU/SIE
	D	(0.0)	213.1					80MAU/HAM
	D	(0.0)	213.1					83MAU
320	A	(17.7)	213.3					83TAF
320	A	(17.0)	212.6					75TAF-75ARN
300	B	(6.4)	212.1					75AUE/WEB (2)
300	B	(6.9)	212.6*					76AUE/WEB
520	C	(3.0)	213.1					79MAU
550	A	(17.3)	212.5					80MAU
600	A	(17.6)	212.7					78LAU/SAL
600	A	(18.6)	213.7					72BRI/YAM
[C ₃ H ₇ N]	(CH ₃) ₂ C=NH	RN 38697-07-3		212.9	891	220.7	923	
	(br)		212.9					81ELL/DIX
[C ₁₀ H ₂₃ N]	n-(C ₁₀ H ₂₁)NH ₂	RN 2016-57-1		212.3**	888.**	220.7**	923.**	
			212.3**					79AUE/BOW
[C ₃ H ₉ O ₃ P]	P(OCH ₃) ₃	RN 121-45-9		213.0	891.	220.6	923.	
300	(Key)		213.0					80HOD/MCD
[C ₉ H ₁₂ O ₃]	1,3,5-C ₆ H ₃ (OCH ₃) ₃	RN 621-23-8		213.1	892.	220.6	923.	
320	A	(17.5)	213.1					TAFT
[C ₈ H ₁₅ NO]	trans-3-Aminobicyclo[2.2.2]- octan-2-ol	RN 40335-14-6		212.8	890.	220.6	923.	
	(Key)		212.8					83HOU/RUF

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₂ H ₇ N]	(CH ₃) ₂ NH	RN 124-40-3		212.8	890.	220.6	923.	
320	A	(17.1)	212.7					83TAF
320	A	(16.8)	212.4					75TAF-75ARN
320	A	(16.8)	212.4					72HEN/TAA-72ARN/JON
	A	(16.8)	212.4					83MCI
300	B	(6.8)	212.5					72AUE/WEB
300	B	(6.5)	212.1					75AUE/WEB (2)
300	B	(6.6)	212.3*					76AUE/WEB
600	A	(18.3)	213.4					72BRI/YAM
	D	(0.1)	213.2					80MAU/HAM
[C ₄ H ₁₁ N]	sec-C ₄ H ₉ NH ₂	RN 13952-84-6		211.7	886.	220.5	922.	
300	B	(6.4)	212.1*					76AUE/WEB
300	B	(6.0)	211.7					72AUE/WEB
[C ₈ H ₁₉ N]	n-(C ₈ H ₁₇)NH ₂	RN 111-86-4		212.0**	887.**	220.4**	922.**	
			212.0**					79AUE/BOW
[C ₅ H ₅ NO]	Pyridine-N-oxide	RN 694-59-7		213.3	892.	220.3	922.	
550	C	(3.2)	213.3					79MAU
[C ₉ H ₉ N]	(HCCCH ₂) ₃ N	RN 6921-29-5		212.4	889.	220.2	921.	
320	A	(16.6)	212.2					TAFT
320	A	(16.0)	211.6					75TAF-75ARN
[C ₆ H ₇ NO]	1-Methyl-2-pyridinone	RN 694-85-9		212.2	888.	220.2	921.	
300	B	(6.5)	212.2					79AUE/BET
320	D	(-1.0)	212.6					76COO/KAT
[C ₅ H ₉ NO ₂]	c-C ₄ H ₇ NH(2-COOH)	(L-Proline) RN 609-36-9		212.4	889.	220.2	921.	
370	A	(20.1)	215.7					83MCI
600	A	(11.0)	212.4					73YAM/KEB
[C ₁₀ H ₁₇ NO]	trans-3-Amino-2-twistanol	RN xxxxx		212.2	888.	220.0	920.	
	(Key)		212.2					83HOU/RUF
[C ₁₂ H ₂₁ NO]	3-Amino-tricyclo[7.3.0.0 ^{4,8}] dodecan-2-ol	RN xxxxx		212.2	888.	220.0	920.	
	(Key)		212.2					83HOU/RUF
[C ₁₀ H ₈]	Azulene	RN 275-51-4		212.5	889.	220.	921.	
320	A	(23.7)	219.3					TAFT
320	A	(22.6)	218.2					75WOL/HAR
320	ZZ	(18.1)	218.1					77WOL/ABB
550	A	(17.3)	212.5	(1983, value reconfirmed)				80MAU
[C ₂ H ₈ N ₂]	(CH ₃) ₂ NNH ₂	RN 57-14-7		212.1	887.	219.9	920.	
320	A	(16.8)	212.4					TAFT
320	A	(16.2)	211.8					75TAF-75ARN
[C ₄ H ₈ N ₂ O ₃]	L-Asparagine	RN 3130-87-8		212.0	887.	219.8	920.	
	A	(16.4)	212.0					83MCI

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₃ H ₄ N ₂]	Imidazole	RN 288-32-4		212.0	887.	219.8	920.	
600	D	(-1.1)	212.0					83MAU
[C ₄ H ₉ O ₃ P]	2-Methoxy-1,3,2-dioxaphosphorinane	RN 31121-06-9		211.7	886.	219.4	918.	
320	(Key)		211.7					80HOD/HOU
[C ₄ H ₉ NO]	Morpholine	RN 110-91-8		211.6	885.	219.4	918.	
300	C	(1.5)	211.6					73AUE/WEB
[C ₆ H ₁₄ O ₃]	CH ₃ (OCH ₂ CH ₂) ₂ OCH ₃	RN 111-96-6		207.4	868.	219.4	918.	
300	(Key)		207.4					83MAU
300	A	(10.9)	206.5					84SHA/BLA
[C ₅ H ₁₃ N]	neo-C ₅ H ₁₁ NH ₂	RN 5813-64-9		211.7	886.	219.3	917.5	
320	A	(16.1)	211.6					TAFT
300	B	(6.1)	211.8					76AUE/WEB
[C ₅ H ₄ N ₄]	Purine	RN 120-73-0		211.5	885.	219.3	917.5	
515	C	(+1.4)	211.5					79MAU
	(br)		209					75WIL/MCC
[C ₆ H ₁₁ NO]	c-C ₅ H ₈ N(2-O)1-CH ₃	RN 931-20-4		211.5	885.	219.3	917.5	
300	B	(5.8)	211.5					79AUE/BET
320	A	(15.6)	211.2					76COO/KAT
			209.1**					79AUE/BOW
[C ₃ H ₇ N]	2-Methylaziridine	RN 75-55-8		211.4**	884.**	219.2**	917.**	
			211.4**					79AUE/BOW
[C ₂ H ₅ N]	CH ₂ =CHNH ₂	RN 593-67-9		211.3	884.	219.1	917.	
	(Key)		211.3					81LELL/DIX
[C ₇ H ₁₇ N]	n-C ₇ H ₁₅ NH ₂	RN 111-68-2		211.2	884.	219.0	916.	
300	C	(1.1)	211.2					73AUE/WEB
[C ₆ H ₆ N]	C ₆ H ₅ NH radical	RN xxxxx		211	883	219	916	
	(br)		211					82MAU
[C ₆ ClH ₆ N]	2-Chloro-6-methylpyridine	RN 18368-63-3		211**	883*	219**	916**	
			211**					79AUE/BOW
[Mg ₂]	Mg ₂	RN 29904-79-8				219+7	916.	
	(Key)							77PO/POR

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
	[C ₇ H ₆ O]	2,4,6-Cycloheptatriene-1-one (Tropone) RN 539-80-0 (br) (br)		212	887	219	918	
			212			226**	945**	77DIT/NIB 83CAS/FRE
	[C ₅ H ₁₃ N]	n-C ₅ H ₁₁ NH ₂ RN 110-58-7		211.1	883.	218.9	916.	
		300 B (5.6)	211.3					79AUE/BET
		535 C (+0.8)	210.9					79MAU
	[C ₆ H ₁₃ NO ₂]	L-Isoleucine RN 73-32-5		211.1	883.	218.9	916.	
		A (15.6)	211.1					83MCI
	[C ₆ H ₁₅ N]	n-C ₆ H ₁₃ NH ₂ RN 111-26-2		211.1	883.	218.9	916.	
		300 C (1.0)	211.1					73AUE/WEB
			211.7**					79AUE/BOW
	[C ₄ H ₁₁ N]	i-C ₄ H ₉ NH ₂ RN 78-81-9		211.1*	883.*	218.8*	915.*	
		300 B (5.4)	211.1*					76AUE/WEB
		300 B (4.8)	210.5					72AUE/WEB
		320 A (17.1)	212.7					75ARN
	[C ₃ H ₇ NO ₂]	Sarcosine RN xxxxx		210.9	882.	218.7	915.	
		A (15.3)	210.9					83MCI
	[C ₃ H ₉ N]	i-C ₃ H ₇ NH ₂ RN 75-31-0		211.0	883.	218.6	915.	
		320 A (15.4)	211.0					75TAF-72HEN/TAA-72ARN/JON-83TAF
		300 B (5.3)	211.0*					76AUE/WEB
		300 B (4.9)	210.6					72AUE/WEB
		600 D (-3.5)	209.6					83MAU
	[C ₄ H ₉ NO ₃]	L-Threonine RN xxxxx		210.8	882.	218.6	915.	
		A (15.2)	210.8					83MCI
	[C ₆ ClH ₆ N]	2-Chloro-4-methylpyridine RN 3678-62-4		210.8**	882.**	218.6**	915.**	
			210.8**					79AUE/BOW
	[C ₅ H ₁₀ N ₂ O ₃]	L-Glutamine RN 585-21-7		210.6	881.	218.4	914.	
		A (15.0)	210.6					83MCI
	[C ₄ H ₁₁ N]	n-C ₄ H ₉ NH ₂ RN 109-73-9		210.6	881.	218.4	914.	
		320 A (14.6)	210.2					83TAF
		320 A (14.5)	210.1					75TAF-75ARN
		320 A (14.8)	210.4					83MCI
		300 B (4.9)	210.6*					76AUE/WEB
		300 B (4.5)	210.2					72AUE/WEB
		515 C (0.5)	210.6					79MAU
	[C ₆ H ₉ N]	2,5-Dimethylpyrrole RN 625-84-3		210.6	881.	218.4	914.	
		600 D (-2.5)	210.6					83MAU

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₄ H ₉ N]	CH ₂ =C(CH ₃)CH ₂ NH ₂	RN 2878-14-0	209.6**	209.6** 877.**	218.2** 913.**	79AUE/BOW
[C ₇ H ₉ N]	C ₆ H ₅ NHCH ₃	RN 100-61-8	210.3	210.3 880.	218.1 912.5	72BRI/YAM-78LAU/SAL
	A	(15.1)	210.3			
[C ₆ H ₁₃ NO ₂]	(CH ₃) ₂ CHCH ₂ CH(NH ₂)COOH (L-Leucine)	RN 61-90-5	210.3	210.3 879.	218.1 912.5	
	500 F	(6.0)	208.5			79MAU/HUN
	A	(14.7)	210.3			83MCI
[C ₁₀ H ₁₀ Ru]	(C ₅ H ₅) ₂ Ru	RN 1287-13-4			218** 912**	
	320 A	(14±2)				81STE/BEA
[BrC ₅ H ₄ N]	Pyridine,4-Br	RN 1120-87-2	210.1*	210.1* 879.*	217.9* 912.*	
			210.1*			76AUE/WEB (2)
[C ₃ H ₉ N]	n-C ₃ H ₇ NH ₂	RN 107-10-8	210.1	210.1 879.	217.9 912.	
	535 C	(0.0)	210.1			79MAU
	300 C	(0.0)	210.1			73AUE/WEB
	320 A	(14.1)	209.7			83TAF
	320 A	(13.9)	209.5			75TAF-75ARN
	A	(14.4)	210.0			83MCI
	300 B	(4.4)	210.1*			76AUE/WEB
	300 B	(3.9)	209.6			72AUE/WEB
[C ₅ ClH ₄ N]	4-Chloropyridine	RN 626-61-9	210.0	210.0 879.	217.8 911.	
	320 A	(14.4)	210.0			TAFT
	320 A	(14.1)	209.7			75TAF-75ARN
			210.0*			76AUE/WEB (2)
[C ₁₈ H ₁₂]	Tetracene	RN 92-24-0	210.4	210.4 880.	217.8 911.	
	550 A	(14.8)	210.4			80MAU
[C ₃ FH ₈ N]	FCH ₂ CH ₂ CH ₂ NH ₂	RN 462-41-9	210.3*	210.3* 880.*	217.8* 911.*	
			210.3**			79AUE/BOW
[C ₆ ClH ₆ NO]	6-Chloro-1-methyl-2(1H)pyridinone	RN 17228-63-6	210.0	210.0 879.	217.8 911.	
	300 B	(4.3)	210.0			79AUE/BET
[C ₅ H ₉ NO ₃]	CH ₃ CONHCH ₂ COOCH ₃	RN xxxxx	205.8	205.8 861.	217.7 911.	
	(Key)		205.8			83MAU
[C ₇ H ₉ NO]	3-CH ₃ OC ₆ H ₄ NH ₂	RN 536-90-3	209.8	209.8 878.	217.6 910.	
	600 F	(7.3)	209.8			81LAU/NIS
[C ₁₀ H ₁₂ O]	4-CH ₃ OC ₆ H ₄ (CCH ₃ CH ₂)	RN 1712-69-2	209.6	209.6 877.	217.4 910.	
	320 A	(14.0)	209.6			TAFT
[C ₇ H ₇ NO]	1-(4-Pyridinyl)-ethanone	RN 1122-54-9	209.6	209.6 877.	217.4 910.	
	320 A	(14.0)	209.6			TAFT
			209.6**			79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol		kJ/mol	Proton affinity kcal/mol		kJ/mol	Reference
[C ₇ H ₇ NO]		1-(3-Pyridinyl)-ethanone RN 350-03-8			209.4	876.		217.2	909.		
	320	A	(13.8)	209.4							TAFT
[C ₆ H ₁₅ O ₄ P]		OP(OC ₂ H ₅) ₃ RN 78-40-0			~209.5	~877		~217	~910		
	300	(Key)		~209.5							80HOD/MCD
[C ₅ H ₁₁ NO ₂]		(CH ₃) ₂ CHCH(NH ₂)COOH (L-Valine) RN 72-18-4			209.2	875.		217.0	908.		
	500	F	(5.4)	207.9							79MAU/HUN
		A	(13.6)	209.2							83MCI
[C ₂ H ₇ N]		C ₂ H ₅ NH ₂ RN 75-04-7			208.5	872.		217.0	908.		
	320	A	(12.8)	208.4							72HEN/TAA-72ARN/JON
	320	A	(12.7)	208.3							75TAF-75ARN-83TAF
		A	(12.7)	208.3							83MCI
	300	B	(2.2)	207.9							72AUE/WEB
	300	B	(2.7)	208.4							75AUE/WEB (2)
	300	B	(3.0)	208.7*							76AUE/WEB
	550	A	(12.0)	207.2							80MAU
	320	G	(0.0)	208.5							74STA/BEA
	535	G	(0.0)	207.6							79MAU
[C ₄ H ₄ N ₂ S ₂]		Dithiouracil RN 2001-93-6			~209	~874		~217	~907		
		(br)		~209							75WIL/MCC
[C ₅ H ₄ N ₄ O]		Hypoxanthine RN 68-94-0			~209	~874		~217	~907		
		(br)		~209							75WIL/MCC
[C ₁₀ H ₉ N]		1-Naphthylenamine RN 134-32-7			209.1	875.		216.9	907.5		
	600	A	(14.0)	209.1							78LAU/SAL
[C ₄ H ₉ NO]		Dimethylacetamide RN 127-19-5			209.0	874.		216.8	907.		
	320	A	(13.4)	209.0							75TAF
	320	A	(12.8)	208.4							83TAF
[C ₇ H ₉ N]		C ₆ H ₅ CH ₂ NH ₂ RN 100-46-9			209.0	874.		216.8	907.		
	320	A	(13.2)	208.8							TAFT
				211.3**							79AUE/BOW
[C ₅ H ₉ NO]		1-Methyl-2-pyrrolidinone RN 872-50-4			209.0	874.		216.8	907.		
	300	B	(3.3)	209.0							79AUE/BET
[C ₃ H ₇ NO ₃]		L-Serine RN 302-84-1			209.0	874.		216.8	907.		
		A	(13.4)	209.0							83MCI
[C ₄ H ₇ NO ₄]		L-Aspartic Acid RN 617-45-8			208.9	874.		216.7	907.		
		A	(13.3)	208.9							83MCI
[C ₅ FH ₄ N]		4-Fluoropyridine RN 694-52-0			209.2	875.		216.6	906.		
	320	A	(13.6)	209.2							75TAF-81TAA/SUM
				209.1**							79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	kJ/mol	Proton affinity kcal/mol	kJ/mol	Reference
[C ₉ H ₁₁ NO ₂]	C ₆ H ₅ CH ₂ CH(NH ₂)COOH (L-Phenylalanine)	RN 150-30-1		208.7	873.	216.5	906.	
500	F	(6.2)	208.7					79MAU/HUN
	A	(16.7)	212.3					83MCI
[C ₈ H ₁₂]	(c-C ₃ H ₅) ₂ C=CH ₂	RN 822-93-5		208.7	873.	216.5	906.	
300	ZZ	(8.7)	208.7					77WOL/ABB
			209.5**					79AUE/BOW
[C ₅ H ₉ NO ₄]	L-Glutamic Acid	RN 617-65-2		208.7	873.	216.5	906.	
	A	(13.1)	208.7					83MCI
[C ₂ H ₇ P]	(CH ₃) ₂ PH	RN 676-59-5		208.6	873.	216.3	905.	
320	A	(12.9)	208.7					74STA/BEA
[C ₆ H ₇ N]	(HCCCH ₂) ₂ NH	RN 6921-28-4		208.3	872.	216.1	904.	
320	A	(12.7)	208.3					TAFT
[C ₁₀ H ₁₆]	1,5,5-Trimethyl-3-methylenecyclohexene	RN 16609-28-2		207.7**	869.**	216.1**	904.**	
			207.7**					79AUE/BOW
[C ₆ H ₈ N ₂]	1,4-C ₆ H ₄ (NH ₂) ₂	RN 106-50-3		208.1	870.	215.9	903.	
600	F	(5.6)	208.1					81LAU/NIS
[C ₆ ClH ₆ NO]	2-Chloro-6-methoxypyridine	RN 17228-64-7		208.1	870.	215.9	903.	
	B	(2.4)	208.1					79AUE/BET
[C ₃ H ₇ N]	H ₂ C=CHCH ₂ NH ₂	RN 107-11-9		207.9	870.	215.8	903.	
320	A	(12.3)	207.9					TAFT
320	A	(12.0)	207.6					75ARN
	B	(2.4)	208.1*					76AUE/WEB
[C ₃ H ₈ Sn]	(CH ₃) ₂ Sn=CH ₂	RN 82065-00-7		207.4	868.	215.8	903.	
320		(11.8)	207.4					82PIE/HEH
[C ₈ H ₈]	1,4-C ₆ H ₄ (=CH ₂) ₂	RN xxxxx				215.7	902.	
	(br)							81POL/RAI
[C ₂ H ₅ N]	Aziridine (Azirane)	RN 151-56-4		207.5	868.	215.7	902.	
320	A	(11.6)	207.1					TAFT
320	A	(11.9)	207.4					75TAF-75ARN
300	B	(1.7)	207.4					75AUE/WEB (2)
300	B	(1.9)	207.6*					76AUE/WEB
300	A	(11.9)	207.5					80AUE/WEB
[C ₄ H ₄ N ₂]	Pyridazine (1,2-Diazine)	RN 289-80-5		208.8	874.	215.6	902.	
320	A	(13.6)	209.2					TAFT
535	C	(-0.8)	208.4					79MAU

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₁₆ H ₁₆]	(4-CH ₃ C ₆ H ₄) ₂ C=CH ₂	RN xxxxx		207.6	868.	215.4	901.	
	ZZ	(7.6)	207.6					77WOL/ABB
[C ₆ H ₅ NO]	4-Pyridinecarboxaldehyde			207.4**	868.**	215.2**	900.**	
	RN 872-85-5		207.4**					79AUE/BOW
[BrC ₅ H ₄ N]	3-Bromopyridine	RN 626-55-1		207.3	867	215.1	900.	
	B	(1.6)	207.3					79AUE/BET
			208.5*					76AUE/WEB (2)
[C ₄ F ₃ H ₈ N]	CF ₃ CH ₂ N(CH ₃) ₂	RN 819-06-7		207.4	868.	215.0	900.	
	320 A	(11.8)	207.4					
			207.1**					79AUE/BOW
[C ₃ H ₇ N]	c-C ₃ H ₅ NH ₂	RN 765-30-0		206.6**	864.**	215.0**	899.**	
			206.6**					79AUE/BOW
[C ₄ H ₁₀ N ₂]	c-C(CH ₃)(C ₂ H ₅)NHNH	RN 4901-75-1		207.1**	867.**	214.9**	899.**	
			207.1**					79AUE/BOW
[C ₈ H ₈]	1,2-C ₆ H ₄ (=CH ₂) ₂	RN xxxxx		207.4	868.	214.8	899.	
	320 (br)		207.4					81POL/RAI
[C ₅ ClH ₄ N]	3-Chloropyridine	RN 626-60-8		207.0	866.	214.8	899.	
	320 A	(11.5)	207.1					75TAF
			207.1					83MCI
	300 A	(12.3)	207.9*					76AUE/WEB (2)
	550 A	(11.5)	206.7					80MAU
[C ₃ H ₇ NO ₂]	L-Alanine	RN 56-41-7		206.6	864.	214.8	899.	
	A	(11.0)	206.6					83MCI
	500 F	(6.2)	208.7					79MAU/HUN
[BrC ₅ H ₄ N]	2-Bromopyridine	RN 109-04-6		207.1	866.5	214.7	898.	
	B	(2.1)	207.8					79AUE/BET
			207.8*					76AUE/WEB (2)
	320 A	(11.5)	207.1					TAF
	320 A	(10.8)	206.5					75TAF-75ARN
[C ₇ H ₉ NO]	2-CH ₃ OC ₆ H ₄ NH ₂ (o-Anisidine)			206.9	866.	214.7	898.	
	RN 90-04-0							
	600 A	(15.1)	210.2					73YAM/KEB
	600 A	(11.8)	206.9					78LAU/SAL
[C ₇ H ₉ NS]	3-CH ₃ SC ₆ H ₄ NH ₂	RN 1783-81-9		206.7	865.	214.5	897.	
	600 F	(4.7)	206.7					81LAU/NIS
[C ₃ H ₉ O ₃ PS]	SP(OCH ₃) ₃	RN 29952-66-79		206.7	865.	214.5	897.	
	300 (Key)		206.7					80HOD/MCD
[C ₅ ClH ₄ N]	2-Chloropyridine	RN 109-09-1		206.6	864.	214.4	897.	
	320 A	(11.0)	206.6					TAF
	320 A	(10.8)	206.4					75TAF-75ARN
			207.0*					76AUE/WEB (2)
	500 A	(10.0)	205.1					84SHA/BLA
	546 G	(-1.0)	206.6					79MAU
	550 A	(11.0)	206.2					80MAU

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₈ H ₆ N ₂]	Quinoxaline	RN 91-19-0		207.3	867.	214.4	897.	
535	G	(-0.3)	207.3					79MAU
[C ₁₃ H ₂₅ N]	out-6H-1-Azabicyclo[4.4.4] tetradecane	RN xxxxx		206.5	864.	214.3	897.	
320	A	(10.9)	206.5					81ALD/ARR
[C ₄ F ₃ H ₈ N]	CF ₃ (CH ₂) ₃ NH ₂	RN 819-46-5		206.5	864.	214.3	897.	
320	A	(10.9)	206.5					TAFT
320	A	(10.7)	206.3					75TAF-75ARN
			206.9**					79AUE/BOW
[C ₇ H ₉ NO]	4-CH ₃ OC ₆ H ₄ NH ₂	RN 104-94-9		206.5	864.	214.3	897.	
320	A	(10.9)	206.5					77SUM/POL-81TAA/SUM
			206.6**					79AUE/BOW
[C ₅ FH ₄ N]	3-Fluoropyridine	RN 372-47-4		206.2	863.	214.3	897.	
320	A	(10.6)	206.2					TAFT
			207.0*					76AUE/WEB(2)
[C ₃ H ₇ NO ₂ S]	L-Cysteine	RN 3374-22-9		206.5	864.	214.3	897.	
	A	(10.9)	206.5					83MCI
[C ₆ H ₇ NO]	2-(OH)C ₆ H ₄ NH ₂	RN xxxxx		206.4	864.	214.2	896.	
600	F	(3.9)	206.4					81LAU/NIS
[C ₆ H ₇ NO]	3-(OH)C ₆ H ₄ NH ₂	RN 591-27-5		206.4	864.	214.2	896.	
600	F	(3.9)	206.4					81LAU/NIS
[CH ₅ N]	CH ₃ NH ₂	RN 74-89-5		205.7	861.	214.1	896.	
300	B	(0.0)	205.7					72AUE/WEB
300	B	(0.0)	205.7					75AUE/WEB
300	B	(0.0)	205.7*					76AUE/WEB
320	B	(0.0)	205.7					75HOD/BEA
320	A	(9.8)	205.4					75TAF-83TAF
320	A	(10.1)	205.7					72HEN/TAA-72ARN/JON
382	F	(2.65)	205.2					79LOC/HUN
	A	(9.8)	205.4					83MCI
600	A	(10.0)	205.1					78LAU/SAL
600	A	(10.8)	205.9					72BRI/YAM
[CH ₆ N ₂]	CH ₃ NHNH ₂	RN 60-34-4		206.3**	863**	214.1**	896.**	
			206.3**					79AUE/BOW
[C ₈ H ₁₁ N]	3-C ₂ H ₅ C ₆ H ₄ NH ₂	RN 587-02-0		206.2	863.	214.0	895.	
600	F	(3.7)	206.2					81LAU/NIS
[C ₆ H ₁₀]	1,3,3-Trimethylcyclopropene	RN 3664-56-0		206.**	862.**	214.**	895**	
			206.**					79AUE/BOW
[C ₂ H ₅ N]	CH ₃ CH-NH	RN 20729-41-3		206.1	862.	213.9	895.	
	(br)		206.1					81ELL/DIX
	(br)		206.1					79ELL/EAD

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₁₅ H ₁₂]		9-Methylanthracene	RN 779-02-2		206.1	862.	213.9	895.	
	550	A	(10.9)	206.1					80MAU
[C ₆ H ₉ O ₃ P]		2,8,9-Trioxa-1-phosphadamantane	RN 281-33-4		206.0	862.	213.8	894.	
		(Key)		206.0					80HOD/HOU
[C ₅ H ₁₂ O ₂]		CH ₃ O(CH ₂) ₃ OCH ₃	RN 17081-21-9		204.8	857.	213.8	894.	
	300	(Key)		204.8					83MAU
[C ₅ H ₁₁ NO ₂]		(CH ₃) ₂ NCOOC ₂ H ₅	RN 687-48-9		205.9	861.	213.7	894.	
	320	A	(10.3)	205.9					TAFT
[C ₇ H ₉ N]		4-CH ₃ C ₆ H ₄ NH ₂	RN 106-49-0		205.9	861.	213.7	894.	
	320	A	(10.1)	205.7					77SUM/POL-81TAA/SUM
	320	A	(10.7)	206.3					75ARN
[C ₈ H ₈ O ₂]		4-CH ₃ OC ₆ H ₄ CHO	RN 123-11-5		205.7	861.	213.5	893.	
	320	A	(9.9)	205.5					TAFT
	320	A	(10.4)	206.0					
[C ₇ H ₉ N]		3-CH ₃ C ₆ H ₄ NH ₂	RN 108-44-1		205.6	860.	213.4	893.	
	320	A	(10.1)	205.7					75TAF-77SUM/POL
	600	F	(3.0)	205.5					81LAU/NIS
[C ₉ H ₁₃ N]		3-CH ₃ C ₆ H ₄ N(CH ₃) ₂	RN 121-72-2		205.6	860.	213.4	893.	
	320	F	(3.0)	205.9					77POL/DEV
		A	(10.0)	205.6					83MCI
[AsC ₃ H ₉]		(CH ₃) ₃ As	RN 593-88-4		205.6	860.	213.4	893.	
	320	B	(0.1)	205.6					75HOD/BEA
[C ₃ H ₃ NS]		Thiazole	RN 288-47-1		205.4	859.	213.2	892.	
	600	(Key)		205.4					83MAU
[C ₆ H ₁₀ O ₂]		CH ₃ COCH ₂ CH ₂ COCH ₃	RN 110-13-4		201.5	843.	213.2	892.	
	300	(Key)		201.5					83MAU
[C ₇ H ₁₂]		(CH ₃) ₂ C=CHC(CH ₃)=CH ₂	RN xxxxx		204.3**	855.	213.1**	892.	
				204.3**					79AUE/BOW
[C ₆ H ₄]		o-Benzyne	RN xxxxx		205.7	861.	213.0	891.	
		(br)		205.7					80POL/HEH
[C ₆ F ₃ H ₄ N]		4-Trifluoromethylpyridine	RN 3796-24-5		205.0	858.	212.8	890.	
	320	D	(-11.7)	201.4					72TAA/HEN
	320	A	(9.4)	205.0					83TAF
	320	A	(9.3)	204.9					75TAF-75ARN
				205.2*					/6AUE/WEB(2)

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference	
					kcal/mol	kJ/mol	kcal/mol	kJ/mol		
[C ₆ H ₈ N ₂]	600	1,2-C ₆ H ₄ (NH ₂) ₂ F	RN 95-54-5 (3.9)	206.4	206.4	206.4	864.	212.8	890.	81LAU/NIS
[C ₈ H ₁₈ S]	320	(t-C ₄ H ₉) ₂ S A	RN 107-47-1 (9.9)	205.0	205.0	205.0	858.	212.8	890.	TAFT
[C ₃ H ₇ O ₃ P]	320	2-Methoxy-1,3,2-dioxaphos- pholane (Key)	RN 3741-36-4	204.9	204.9	204.9	857.	212.7	890.	80HOD/HOU
[C ₆ F ₃ H ₄ N]	320	3-Trifluoromethylpyridine A	RN 3796-23-4 (9.2)	204.8 205.0*	204.8	204.8	857.	212.6	889.	75TAF-75ARN 79AUE/BOW
[C ₂ FH ₆ N]	320	CH ₂ FCH ₂ NH ₂ A	RN 406-34-8 (8.7)	204.3 204.6**	204.3	204.3	856.	212.3	888.	75TAF-75ARN 79AUE/BOW
[C ₃ H ₉ O ₄ P]	300	OP(OCH ₃) ₃ (Key)	RN 512-56-1	204.2 202.0	204.2	204.2	854.	212.0	887.	80HOD/MCD 893. ***82PIE/HEH
[C ₄ H ₁₀ O ₂]		HO(CH ₂) ₄ OH 198**	RN 110-63-4	198**	198**	198**	828**	212**	887**	79AUE/BOW
[C ₃ H ₅ N]	300	1-Azabicyclo[1.1.0]butane T	RN 19540-05-7 (24.0)	202. 204**	202.	202.	853**	212**	887**	75AUE/WEB (2) 79AUE/BOW
[C ₆ H ₈]		1-Methyl-3-methylenecyclobutene 204**	RN 15082-13-0	204**	204**	204**	853**	212**	887**	79AUE/BOW
[C ₇ H ₅ O ₂ Rh]	320	(C ₅ H ₅)Rh(CO) ₂ A	RN 12192-97-1 (8+2)					212**	887.**	81STE/BEA
[C ₁₄ H ₁₂]	320 320 320	(C ₆ H ₅) ₂ C=CH ₂ A A ZZ	RN 530-48-3 (8.2) (8.5) (4.1)	203.8 204.0 204.1 204.0**	203.8 204.0 204.1 204.0**	203.8 204.0 204.1 204.0**	854.	211.9	887.	75TAF 75WOL/HAR 77WOL/ABB 79AUE/BOW
[C ₆ H ₈ O ₂]	300	1,3-Cyclohexanedione (Key)	RN 504-02-9	204.5	204.5	204.5	856.	211.9	886.	83MAU
[C ₂ H ₅ NO ₂]	382 500	NH ₂ CH ₂ COOH (Glycine) F A F	RN 56-40-6 (+1.2) (8.2) (0.0)	203.7 203.8 202.5	203.7 203.8 202.5	203.7 203.8 202.5	852.	211.6	885.	79LOC/HUN 83MCI 79MAU/HUN

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₆ F ₃ H ₄ N] 2-Trifluoromethylpyridine RN 368-48-9								
320	A	(8.0)	203.6	203.6	852.	211.5	885.	75TAF
[C ₂₀ H ₁₂] Perylene RN 198-55-0								
550	A	(+9.1)	204.3	204.3	855.	211.4	884.	80MAU
[C ₃ H ₇ NO] (CH ₃) ₂ NCHO RN 68-12-2								
320	A	(8.0)	203.6	203.6	852.	211.4	884.	75TAF
382	F	(0.95)	203.6	204.6**				79AUE/BOW 79LOC/HUN
[C ₂ H ₆ OS] (CH ₃) ₂ SO RN 67-68-5								
320	A	(7.7)	203.3	203.3	851.	211.3	884.	75TAF-83TAF
	(br)		177.					77MCA
600	A	(8.5)	203.6	203.6				79LAU
[C ₄ H ₈ N ₂] NCCH ₂ N(CH ₃) ₂ RN 926-64-7								
320	A	(7.7)	203.3	203.3	851.	211.1	883.	TAFT-75ARN
[C ₁₀ H ₁₂] 4-CH ₃ C ₆ H ₄ C(CH ₃)CH ₂ RN 1195-32-0								
320	A	(7.6)	203.2	203.2	850.	211.0	883.	TAFT
[C ₁₃ H ₁₀ O] (C ₆ H ₅) ₂ CO RN 119-61-9								
320	A	(7.5)	203.1	203.1	850.	210.9	882.	83TAF
[C ₃ H ₅ N] HCCCH ₂ NH ₂ RN 2450-71-7								
320	A	(7.4)	202.9	202.9	849.	210.8	882.	TAFT
			203.1**					79AUE/BOW
[C ₇ H ₁₀ O] (c-C ₃ H ₅) ₂ CO RN 1121-37-5								
320	ZZ	(2.9)	202.9	202.9	849.	210.7	881.5	83TAF
320	U	(9.8)	198.7	198.7				81BRO/ABB
[C ₃ F ₃ H ₆ N] CF ₃ CH ₂ CH ₂ NH ₂ RN 460-39-9								
320	A	(7.4)	202.9	202.9	849.	210.6	881.	TAFT
320	A	(7.2)	202.7	202.7				75TAF-75ARN
			203.4**					79AUE/BOW
[C ₅ FH ₄ N] 2-Fluoropyridine RN 372-40-5								
320	A	(7.4)	202.9	202.9	849.	210.6	881.	TAFT
320	A	(7.2)	202.7	202.7				75TAF-75ARN
	A	(7.3)	202.8	202.8				83MCI
			204.0	204.0				76AUE/WEB (2)
382	F	(0.35)	202.8	202.8				79LOC/HUN
500	A	(6.6)	201.7	201.7				84SHA/BLA
[C ₈ H ₁₄] (CH ₃) ₂ C=C(CH ₃)C(CH ₃)=CH ₂ RN xxxxx								
			201.8**	201.8**	844.**	210.6**	881.**	79AUE/BOW
[C ₄ H ₄ N ₂] Pyrimidine (1,3-Diazine) RN 289-95-2								
320	A	(8.0)	203.6	203.6	851.	210.5	881.	TAFT
510	G	(-4.2)	203.4	203.4				79MAU
[C ₁₅ H ₁₂] 2-Methylanthracene RN 613-12-7								
550	A	(+7.3)	202.5	202.5	847.	210.3	880.	80MAU

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₆ H ₄ N ₂]	4-Pyridinecarbonitrile	RN 100-48-1	202.5	851.	210.3	880.		
320	A	(6.5)	202.1 202.8*					75TAF-75ARN 76AUE/WEB (2)
[C ₅ H ₉ O ₃ P]	4-Methyl-3,6,7-trioxa-1-phospha- bicyclo[2.2.2]-octane	RN 1449-91-8	202.2	846.	210.0	879.		
320	(Key)		202.2					80HOD/HOU
[C ₄ H ₉ NO]	n-C ₃ H ₇ NHCHO	RN 6281-94-3		202.2**	846.	210.0**	879.	
				202.2**				79AUE/BOW
[C ₁₀ FeH ₁₀]	(C ₅ H ₅) ₂ Fe	RN 102-54-5		-202	-845.	-210	-879.	
	(br)			-202				75FOS/BEA
[C ₃ F ₃ H ₆ N]	CF ₃ CH ₂ NHCH ₃	RN 2730-67-8		202.2	846.	209.8	878.	
320	A	(6.3)	201.9 202.4**					75TAF 79AUE/BOW
[C ₃ H ₄ N ₂]	Pyrazole	RN 288-13-1		202.0	845.	209.8	878.	
600	(Key)		202.0					83MAU
[C ₆ H ₁₄ S]	(i-C ₃ H ₇) ₂ S	RN 625-80-9		201.8	844.	209.6	877.	
320	A	(6.2)	201.8 202.0**					TAFT 79AUE/BOW
[C ₆ H ₇ N]	C ₆ H ₅ NH ₂	RN 62-53-3		202.5	847.	209.5	876.	
320	F	(0.0)	202.8					77POL/DEV
382	F	(0.0)	202.5					79LOC/HUN
500	F	(0.0)	202.5					79MAU/HUN
600	F	(0.0)	202.5					81LAU/NIS
320	A	(7.2)	202.8					75TAF-75ARN
320	A	(6.9)	202.5					83TAF
550	G	(-5.2)	202.4					79MAU
550	A	(+5.8)	201.0					80MAU
600	A	(6.9)	202.0					78LAU/SAL
600	J	(25.6)	201.2					76LAU/KEB
600	A	(8.9)	203.5					72BRI/YAM
[C ₆ H ₄ N ₂]	3-Pyridinecarbonitrile	RN 100-54-9	201.5	843.	209.3	876.		
320	A	(5.6)	201.2					TAFT
320	A	(5.9)	201.5 201.7*					75TAF-75ARN 76AUE/WEB (2)
[C ₆ H ₈ O]	2,5-Dimethylfuran	RN 625-86-5		201.3	842.	209.1	875.	
600	(Key)		201.3					83MAU
[C ₄ H ₄ N ₂]	Pyrazine (1,4-Diazine)	RN 290-37-9	201.2	842.	209.0	874.		
320	A	(5.6)	201.2					TAFT
550	G	(-6.0)	201.2					79MAU
[C ₆ H ₁₀]	c-C ₃ H ₅ C(CH ₃)=CH ₂	RN 4663-22-3		201.2	842.	209.0	874.	
320	A	(5.6)	201.2 200.9**					TAFT 79AUE/BOW
[C ₈ H ₁₈ O]	(sec-C ₄ H ₉) ₂ O	RN 6863-58-7		201.2	842.	209.0	874.	
335	XX	(6.7)	201.2					82MAU

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₆ H ₆ IN]	3-IC ₆ H ₄ NH ₂ RN 626-01-7			201.1	841.	208.9	874.	
600	F	(-0.9)	201.1					81LAU/NIS
[C ₅ H ₆ N ₂ O ₂]	Thymine RN 65-71-4			201.0	841.	208.8	874.	
550	G (br)	(-6.6)	201.0 ~200					79MAU 75WIL/MCC
[C ₇ H ₁₆ O]	(i-C ₃ H ₇)O(t-C ₄ H ₉) RN 17348-59-3			201.0**	841.**	208.8**	874.**	
			201.0**					79AUE/BOW
[C ₉ H ₁₀ O]	(4-CH ₃)C ₆ H ₄ COCH ₃ RN xxxxx			200.9	840.5	208.7	873.	
320	A	(12.0)	200.9					81BRO/ABB
[C ₈ H ₁₈ S]	(n-C ₄ H ₉) ₂ S RN 544-40-1			200.9	840.5	208.7	873.	
300	(Key)		200.9					83MAU
[C ₆ ClH ₆ N]	4-ClC ₆ H ₄ NH ₂ RN 106-47-8			201.0	841.	208.6	873.	
320	A	(5.1)	200.7					TAPT
320	A	(5.4)	201.0					75TAF-75ARN-77SUM/POL
[C ₅ H ₄ N ₂ O ₂]	4-Nitropyridine RN 1122-61-8			200.7	840.	208.5	872.	
320	D	(-18.1)	194.9					72TAA/HEN
320	A	(5.0)	200.6					TAPT
320	A	(5.2)	200.8 201.7**					75TAF-75ARN 79AUE/BOW
[C ₂₂ H ₁₂]	1,12-Benzoperylene RN 191-24-2			201.1	841.	208.5	872.	
550	A	(5.9)	201.1					80MAU
[C ₃ H ₃ NO]	Oxazole RN 288-42-6			200.6	839.	208.4	872.	
600	(Key)		199.2					83MAU
[C ₄ H ₈ O]	C ₂ H ₅ OCH=CH ₂ RN 109-92-2			200.4	838.	208.2	871.	
600	(Key)		200.4					83MAU
[C ₆ H ₄ N ₂]	2-Pyridinecarbonitrile RN 100-70-9			200.3	838.	208.1	871.	
320	A	(4.3)	199.9 201.1*					TAPT
500	F	(-1.9)	200.6					76AUE/WEB(2) 79MAU/HUN
[BrC ₆ H ₆ N]	3-BrC ₆ H ₄ NH ₂ RN 591-19-5			200.3	838.	208.1	871.	
600	F	(-2.2)	200.3					81LAU/NIS
[C ₆ FH ₆ N]	4-FC ₆ H ₄ NH ₂ RN 371-40-4			200.3	838.	208.1	871.	
320	A	(4.6)	200.2					81TAA/SUM
320	A	(5.0)	200.6					75TAF-75ARN
[C ₁₀ H ₁₄ N ₂ O ₅]	Thymidine RN 50-89-5			~200	~837	~208	~870	
	(br)		~200					75WIL/MCC
[C ₉ H ₁₂ N ₂ O ₆]	Uridine RN 58-96-8			~200	~837	~208	~870	
	(br)		~200					75WIL/MCC
[C ₉ H ₁₄ N ₂ O ₆]	5,6-Dihydrouridine RN 5627-05-4			~200	~837	~208	~870	
	(br)		~200					75WIL/MCC

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol		Proton affinity kJ/mol	Reference
							kcal/mol	
[C ₁₂ H ₁₆ N ₂ O ₆]		2',3'-O-Isopropylideneuridine		~200	~837		~208	~870
		RN 362-43-6						
		(br)		~200				75WIL/MCC
[C ₄ H ₄ N ₂ O]		2(1H)-Pyrimidinone		RN 557-01-7	~200	~837	~208	~870
		(br)		~200				75WIL/MCC
[C ₄ H ₄ N ₂ O ₂]		Uracil		RN 66-22-8	~200	~837	~208	~870
		(br)		~200				75WIL/MCC
[C ₅ H ₃ ClN ₄]		6-Chloropurine		RN 87-42-3	~200	~837	~208	~870
		(br)		~200				75WIL/MCC
[C ₆ H ₁₀]		CH ₃ CH=CHC(CH ₃)=CH ₂		RN 1118-58-7	199.9**	836.**	207.9**	870.**
				199.9**				79AUE/BOW
[C ₅ H ₈ O ₂]		CH ₃ COCH=C(OH)CH ₃		RN 123-54-6	199.3	834.	207.8	869.
	320	A	(3.7)	199.3				TAFT
	300	(Key)		199.2				83MAU
				200.1**				79AUE/BOW
[C ₄ H ₅ N]		Pyrrole		RN 109-97-7	200.3	838.	207.6	868.
	550	G	(-6.4)	201.6**				79AUE/BOW
	600	A	(5.6)	200.8				79MAU
	600	A	(7.15)	200.3				79LAU
				201.8				73YAM/KEB
[C ₂ F ₂ H ₅ N]		CF ₂ HCH ₂ NH ₂		RN 430-67-1	199.8	836.	207.5	868.
	320	A	(4.0)	199.6				TAFT
	320	A	(4.2)	199.8				75TAF-75ARN
				200.0**				79AUE/BOW
[C ₃ H ₆ O]		CH ₂ =CHOCH ₃		RN 107-25-5	199.6	835.	207.4	868.
	600	(Key)		199.6				83MAU
[C ₁₂ H ₁₈]		(CH ₃) ₆ C ₆		RN 87-85-4	200.0	837.	207.3	867.
	320	ZZ	(0.0)	200.0				77WOL/ABB
	320	A	(4.2)	199.8				83TAF
	320	J	(25.1)	199.7				76WOL/DEV
	320	A	(4.8)	200.4				75WOL/HAR
[C ₆ ClH ₆ N]		3-Chlorobenzeneamine		RN 108-42-9	199.4	834.	207.2	867.
	320	A	(4.5)	200.0				75ARN
	320	A	(4.0)	199.6				77SUM/POL
	600	A	(3.6)	198.8				79LAU
	600	F	(-3.1)	199.4				81LAU/NIS
[C ₂ H ₇ O ₃ P]		(CH ₃ O) ₂ P=O		RN 868-85-9			207.2	867.
		(br)		(PA associated with P-protonation: 213.5 kcal/mol)				82PIE/HEH(2)
[C ₄ F ₂ H ₇ NO]		CF ₂ HCON(CH ₃) ₂		RN 667-50-5			207.2	867.
								**82PIE/HEH(2)
[C ₄ H ₇ O ₃ P]		2,6,7-Trioxa-1-phosphabicyclo-		RN 280-45-5	200.0	837.	207.1	866.5
		[2.2.2]octane		RN 280-45-5				
	320	(Key)		200.0				80HOD/HOU

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₈ H ₁₂]					199** 833**	207** 866**	79AUE/BOW
		RN 497-35-8		199**			
[C ₁₄ H ₁₀]					199.9 836.	207.0 866.	80MAU
		RN 120-12-7		199.9			
	550	A	(4.7)	199.9			
[B ₄ C ₂ H ₆]					199. 833.	207. 866.	80DIX
		RN 20693-67-8		199.			
		(br)					
[C ₆ FH ₆ N]					199.2 833.	207.0 866.	77SUM/POL 81LAU/NIS
		RN 372-19-0		199.2			
	320	A	(3.5)	199.1			
	600	F	(-3.2)	199.3			
[C ₉ H ₁₀]					199.2 833.	207.0 866.	75TAF-75WOL/HAR-78TAF/WOL 77WOL/ABB
		RN 98-83-9		199.2			
	320	A	(3.6)	199.2			
	320	ZZ	(-1.0)	199.0			
[C ₃ H ₆ N ₂]					198.1 829.	207.0 866.	TAFT 75ARN 79MAU 80MAU
		RN 151-18-8		198.1			
	320	A	(2.6)	198.2			
	320	A	(3.2)	198.7			
	550	G	(-9.6)	198.0			
	550	A	(+2.4)	197.6			
[C ₂ H ₆ N ₂]					199.1 833.	206.9 866.	74FOS/WIL 72FOS/BEA
		RN 4143-41-3		199.1			
		(Key)		199.1			
		(br)		200			
[C ₉ FH ₉]					199.0 833.	206.7 865.	TAFT
		RN 350-40-3		199.0			
	320	A	(3.4)	199.0			
[C ₆ H ₁₄ S]					198.7 831.	206.5 864.	TAFT
		RN 111-47-7		198.7			
	320	A	(3.1)	198.7			
[C ₉ H ₁₈ O]						206.5 864.	*82PIE/HEH (2)
		RN 815-24-7					
[C ₂ H ₅ NO]					198.4 830.	206.2 863.	73YAM/KEB
		RN 60-35-5		198.4			
	320	A	(3.3)	198.4			
[C ₁₆ H ₁₀]					199.8 836.	206.1 862.	80MAU
		RN 129-00-0		199.8			
	550	A	(4.6)	199.8			
[C ₃ H ₆ N ₂]					198.2 829.	206.0 862.	TAFT
		RN 5616-32-0		198.2			
	320	A	(2.6)	198.2			
[C ₆ H ₁₄ O]					198.4 830.	206.0 862.	81BRO/ABB 82MAU 80LIA/SHO 79AUE/BOW
		RN 108-20-3		198.4			
	320	U	(9.7)	198.6			
	335	XX	(3.9)	198.4			
	340	H	(10.9)	198.0			
				198.7**			
[C ₈ H ₁₂]						206. 862.	76SOL/FIE
		RN xxxxx					
		(Key)					

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₄ H ₆]	1-Methylcyclopropene	RN 3100-04-7	198**	828**	206**	862**		79AUE/BOW
[C ₆ H ₁₀]	CH ₂ =CH(CH ₃)C(CH ₂) ₂	RN 16906-27-7	198*	828*	206*	862*		79AUE/BOW
[C ₉ CrH ₈ O ₃]	(C ₅ H ₅)Cr(CO) ₃ CH ₃	RN 41311-89-1 320 A (2.0+2.0)			206**	862**		81STE/BEA
[C ₂ H ₅ NO]	HCONHCH ₃	RN 123-39-7 320 U (9.1)	198.0	828.	205.8	861.		81BRO/ABB
[C ₄ H ₉ NO ₂]	t-C ₄ H ₉ ONO	RN 540-80-7 (br)	197.9	828.	205.7	861.		78FAR/MCM
[C ₅ H ₆ S]	2-Methylthiophene	RN 554-14-3 600 (Key)	197.9	828.	205.7	861.		83MAU
[C ₆ H ₁₀]	CH ₃ CH=C(CH ₃)CH=CH ₂	RN 4549-74-0	197.3**	825.5**	205.7**	861.		79AUE/BOW
[C ₅ H ₆ O]	2-Methylfuran	RN 534-22-5 600 (Key)	197.8	828.	205.6	860.		83MAU
[C ₈ H ₈ O]	C ₆ H ₅ COCH ₃	RN 98-86-2 320 A (1.7) 320 U (8.9) 600 A (2.0) 600 F (-4.7)	197.3 197.8 197.1 197.8	826.	205.4	859.		83TAF 81BRO/ABB 79LAU 81LAU/NIS
[C ₆ H ₁₂ O]	2,2-Dimethyltetrahydrofuran	RN xxxx 320 U (8.7)	197.6	827.	205.4	859.		81BRO/ABB
[C ₆ H ₁₄ O]	C ₂ H ₅ O(t-C ₄ H ₉)	RN 637-92-3 320 A (1.9)	197.5	826.	205.3	859.		TAFT
[C ₁₀ H ₂₂ O]	(n-C ₅ H ₁₁) ₂ O	RN 693-65-2	197.9**	828.**	205.2**	859.**		79AUE/BOW
[C ₅ H ₈ O]	c-C ₃ H ₅ COCH ₃	RN 765-43-5 320 A (0.7) 320 U (8.4)	196.3 197.3	826.	205.1	858.		83TAF
[C ₄ H ₁₀ S]	(C ₂ H ₅) ₂ S	RN 352-93-2 320 A (1.6)	197.2 198.3**	825.	205.0	858.		TAFT 79AUE/BOW
[C ₉ ClH ₉]	4-ClC ₆ H ₄ C(CH ₃)=CH ₂	RN 1712-70-5 320 A (1.6)	197.2	825.	205.0	858.		TAFT
[C ₂₄ H ₁₂]	Coronene	RN 191-07-1 550 A (4.3)	199.9	836.	205.0	858.		80MAU
[CH ₂ N ₂]	CH ₂ N ₂	RN 334-88-3 (br)	197.	824.	205.	858.		72FOS/BEA

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₁₀ CrH ₇ O ₃]	(C ₆ H ₅ CH ₂)Cr(CO) ₃	RN 32984-97-7			205** 858**	
320	A	(1+2)				81STE/BEA
[C ₃ GeH ₈]	(CH ₃) ₂ Ge=CH ₂	RN 82064-99-1		195.6 818.	204.9 857.	
320	A	(0.9)	196.5			82PIE/HEH
[C ₄ H ₁₀ O ₂]	CH ₃ OCH ₂ CH ₂ OCH ₃	RN 110-71-4		195.8 819.	204.9 857.	
300	(Key)		195.8			83MAU
300	A	(0.2)	195.3			84SHA/BLA
[C ₇ H ₁₄ O]	(i-C ₃ H ₇) ₂ CO	RN 565-80-0		197.0 824.	204.9 857.	
320	ZZ	(-3.9)	196.1			83TAF
320	U	(8.1)	197.0			81BRO/ABB
[C ₆ H ₈ O ₂]	1,2-Cyclohexanedione	RN 765-87-7		197.4 825.	204.8 857.	
300	(Key)		197.4			83MAU
[C ₆ H ₅ NO]	Nitrosobenzene	RN 586-96-9		197.0 824.	204.8 857.	
	(br)		197.0			80REE/FRE
[H ₄ N ₂]	H ₂ NNH ₂	RN 302-01-2		196.7 823.	204.7 856.	
320	A	(4.0)	199.6			75ARN
600	(Key)		196.7			83MAU
[C ₁₄ H ₁₈]	1,2,3,4,5,6,7,8-Octahydro- phenanthrene	RN 5325-97-3		195.1 816.	204.7 856.	
550	A	(-0.1)	195.1			80MAU
[C ₇ F ₃ H ₆ N]	3-CF ₃ C ₆ H ₄ NH ₂	RN 98-16-8		196.4 822.	204.2 854.	
600	F	(-6.1)	196.4			81LAU/NIS
[CH ₅ P]	CH ₃ PH ₂	RN 593-54-4		196.3 821.	204.1 854.	
320	A	(-0.3)	195.3			74STA/BEA
320	H		196.3			
[C ₆ H ₅ O]	C ₆ H ₅ O radical	RN xxxxx		~196 ~820	~204 ~853	
	(br)		~196			80DEF/MCI
[H ₃ N]	NH ₃	RN 7664-41-7		195.6 818.	204.0 853.5	
	Threshold Value				203.6	79CEY/TIE
320	A	(0.0)	195.6			TAF ^T
320	A	(0.0)	195.6			83TAF
320	A	(0.0)	195.6			77WOL/STA
320	A	(0.0)	195.6			75TAF
320	A	(0.0)	195.6			72HEN/TAA
320	A	(0.0)	195.6			72ARN/JON
320	A	(0.0)	195.6			82PIE/HEH
	A	(0.0)	195.6			83MCI
550	A	(0.0)	195.2			80MAU
600	A	(0.0)	195.1			79LAU
600	A	(0.0)	195.1			73YAM/KEB
600	A	(0.0)	195.1			78LAU/SAL
340	H	(8.3)	195.6			80LIA/SHO
320	H		196.6			77WOL/STA
[C ₇ CoH ₅ O ₂]	(C ₅ H ₅)Co(CO) ₂	RN 12078-25-0			~204** 853**	
320	A	(0+2)				81STE/BEA

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol		Proton affinity kcal/mol		Reference
					kJ/mol		kJ/mol	
[C ₈ H ₁₈ O]	(n-C ₄ H ₉) ₂ O	RN 142-96-1		195.9	820.	203.7	852.	
	335 XX	(1.4)	196.6** 195.9					79AUE/BOW 82MAU
[C ₈ H ₈ O]	4-(CH ₃) ₆ H ₄ CHO	RN 104-87-0		195.9	820.	203.7	852.	
	320 A	(0.3)	195.9					TAFT
[C ₈ H ₁₄ O ₂]	c-C ₆ H ₁₁ COOCH ₃	RN 4630-82-4		195.9	820.	203.7	852.	
	320 A	(-0.7)	194.9					83TAF
	320 H		195.9					
[C ₈ H ₈ O ₂]	C ₆ H ₅ CO ₂ CH ₃	RN 95-58-3		195.9	820.	203.7	852.	
	320 U	(7.1)	196.0					81BRO/ABB
	320 H		195.9					
[C ₆ F ₃ H ₁₀ NO]	CF ₃ CONH(n-C ₄ H ₉)	RN 400-59-9		195.8	819.	203.6	852.	
	320 A	(0.2)	195.8					TAFT
[C ₅ H ₁₀ O]	c-C ₄ H ₇ O(2-CH ₃)	RN 96-47-9		195.8	819.	203.6	852.	
	320 U	(6.9)	195.8					81BRO/ABB
[C ₁₂ H ₁₀]	Acenaphthene	RN 83-32-9		196.4	822.	203.5	851.	
	550 A	(1.2)	196.4					80MAU
[C ₃ H ₈ S]	CH ₃ SC ₂ H ₅	RN 624-89-5		195.7	819.	203.5	851.	
	320 A	(-0.9)	194.7					TAFT
	320 H		195.7					79AUE/BOW
			195.8**					
[C ₅ H ₁₂ O]	C ₂ H ₅ O(i-C ₃ H ₇)	RN 625-54-7		195.7	819.	203.5	851.	
	320 A	(-0.9)	194.7					77WOL/STA
	320 H		195.7					81BRO/ABB
	320 U	(7.3)	196.2					
[C ₂₂ H ₁₄]	Picene	RN 213-46-7		196.3	821.	203.4	851.	
	550 A	(1.1)	196.3					80MAU
[C ₁₂ H ₈]	Biphenylene	RN 259-79-0		196.3	821.	203.4	851.	
	550 A	(1.1)	196.3					80MAU
[C ₆ H ₁₈ OSi ₂]	((CH ₃) ₃ Si) ₂ O	RN 107-46-0		~195	~816	~203	~849	
	(br)		~195±3					75PIT/BUR
[C ₄ H ₁₄ OSi ₂]	((CH ₃) ₂ SiH) ₂ O	RN 3277-26-7		~195	~816	~203	~849	
	(br)		~195±3					75PIT/BUR
[C ₄ H ₁₂ OSi]	(CH ₃) ₃ SiOCH ₃	RN 1825-61-2		~195	~816	~203	~849	
	(br)		~195±3					75PIT/BUR
[C ₅ H ₈]	3,3-Dimethylcyclopropene	RN 3907-06-0	196*	196*	820.	203*	849*	
			196*					76AUE/DAV

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	kJ/mol	Proton affinity kcal/mol	kJ/mol	Reference
[C ₅ H ₈ O ₂]	c-C ₃ H ₅ COOCH ₃	RN 2868-37-3		195.1	816.	202.9	849.	
320	A	(-1.5)	194.1					83TAF
320	H		195.1					
320	U	(6.2)	195.1					81BRO/ABB
[C ₆ H ₁₂ O ₂]	t-C ₄ H ₉ COOCH ₃	RN 598-98-1		195.0	816.	202.8	848.5	
320	A	(-1.6)	194.0					83TAF
320	H		195.0					
[C ₄ H ₈ O ₃]	C ₂ H ₅ COOCH ₃	RN 623-53-0		194.9	815.	202.7	848.	
320	A	(-1.7)	193.9					TAFT
320	H		194.9					
[C ₁₄ H ₁₈]	1,2,3,4,5,6,7,8-Octahydro- anthracene	RN 1079-71-6		194.8	815.	202.5	847.	
550	A	(-0.4)	194.8					80MAU
[C ₂ F ₃ H ₄ N]	CF ₃ CH ₂ NH ₂	RN 753-90-2		194.7	815.	202.5	847.	
320	A	(-1.9)	193.7					77STA/TAA-83TAF
320	H		194.7					
320	A	(-1.5)	194.1					75TAF-75ARN
320	H		195.1					
			194.9**					79AUE/BOW
[C ₉ H ₁₁]	C ₆ H ₅ C(CH ₃) ₂ radical	RN xxxxx		194.6	814.	202.4	847.	
	(br)		194.6					82MAU
[C ₈ H ₁₄ O]	c-C ₆ H ₁₁ COCH ₃	RN 823-76-7		194.6	814.	202.4	847.	
320	A	(-2.0)	193.6					83TAF
320	H		194.6					
320	U	(6.4)	195.1					81BRO/ABB
[C ₃ H ₃ NO]	Isooxazole	RN 288-14-2		194.5	814.	202.3	846.	
600	(Key)		194.5					83MAU
[C ₆ H ₁₂ O]	t-C ₄ H ₉ COCH ₃	RN 75-97-8		194.5	814.	202.3	846.	
320	A	(-2.1)	193.5					83TAF
320	H		194.5					
320	U	(5.8)	194.7					81BRO/ABB
[C ₆ H ₁₄ O]	(n-C ₃ H ₇) ₂ O	RN 111-43-3		194.5	814.	202.3	846.	
335	XX	(0.0)	194.5					82MAU
320	U	(5.5)	194.4					81BRO/ABB
320	H		194.7					
340	H	(+7.0)	194.3					80LIA/SHO
			195.6**					79AUE/BOW
[C ₅ H ₁₂ O]	t-C ₄ H ₉ OCH ₃	RN 1634-04-4		194.4	813.	202.2	846.	
320	A	(-2.2)	193.4					TAFT
320	H		194.4					
335	XX	(+0.2)	194.7					82MAU
			196.0**					79AUE/BOW
	(br)		~195+3					75PIT/BUR
[C ₆ H ₁₀]	CH ₂ =C(CH ₃)C(CH ₃)=CH ₂	RN 513-81-5		194.1**	812.**	202.1**	846.**	
			194.1**					79AUE/BOW
[C ₉ H ₁₁]	C ₆ H ₅ (CHC ₂ H ₅) radical	RN xxxxx		~194	~812	~202	~845	
	(br)		~194					82MAU

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	KJ/mol	kcal/mol	KJ/mol	
[C ₈ H ₈]		C ₆ H ₅ CH=CH ₂	RN 100-42-5		194.2	812.5	202.0	845.	
	320	A	(-2.4)	193.2					75WOL/HAR
	320	H		194.2					
[C ₆ H ₁₂ O]		c-C ₆ H ₁₂ O (Oxepane)	RN 592-90-5		195	816	202	845	
	300	(Key)		195					83MAU
[C ₅ FeO ₅]		(CO) ₅ Fe	RN 13463-40-6		~194	~812	~202	~845	
	320	A (br)	(-3+3)	192.4					75FOS/BEA (2) 75FOS/BEA (3)
				194					
[C ₃ H ₇ NO ₂]		i-C ₃ H ₇ ONO	RN 541-42-4		194.1	812.	201.9	845.	
		(br)		194.1					78FAR/MCM
[C ₅ H ₈]		(E)-1,3-Pentadiene	RN 2004-70-8		193.4**	809.**	201.8**	844.**	
				193.4**					79AUE/BOW
[C ₁₈ H ₁₂]		Chrysene	RN 218-01-9		193.8	811.	201.6	843.	
	550	A	(-1.4)	193.8					80MAU
[C ₅ H ₁₀ O ₂]		i-C ₃ H ₇ COOCH ₃	RN 547-63-7		193.8	811.	201.6	843.	
	320	A	(-2.8)	192.8					83TAF
	320	H		193.8					
[C ₅ H ₁₀ O]		(C ₂ H ₅) ₂ CO	RN 96-22-0		193.5	810.	201.4	843.	
	320	A	(-2.8)	192.8					TAFT
	320	H		193.8					
	340	H	(+6.0)	193.3					80LIA/SHO
[C ₆ H ₁₀ O]		Cyclohexanone	RN 108-94-1		194.0	812.	201.4	843.	
	300	(Key)		196.4					83MAU
	320	U	(4.8)	193.7					
	560	(Key)		194.4					79SAL/KEB
[C ₃ H ₆ S]		Thietane	RN 287-27-4		194.0**	812.**	201.3**	842.**	
				194.0**					79AUE/BOW
[C ₅ H ₁₀ O]		(i-C ₃ H ₇)COCH ₃	RN 563-80-4		193.3	809.	201.1	841.	
	320	A	(-3.1)	192.3					83TAF
	320	H		193.3					
[C ₃ H ₃ N ₃]		1,3,5-Triazine	RN 290-87-9		194.5	814.	201.1	841.	
	550	G	(-13.0)	194.5					79MAU
[C ₄ H ₈ O ₂ S]		C ₂ H ₅ S(OCH ₃)CO	RN 38103-96-7		193.2	808.	201.0	841.	
	320	A	(-3.4)	192.2					TAFT
	320	H		193.2					
[C ₈ H ₉]		C ₆ H ₅ CHCH ₃ radical	RN xxxxx		~193	~807	~201	~841	
		(br)		~193					82MAU
[C ₆ H ₁₀]		1,2-Dimethylcyclobutene	RN 1501-58-2		194*	812*	201*	841*	
				194*					76AUE/DAV
[C ₅ H ₈]		1-Methylcyclobutene	RN xxxxx		194*	812*	201*	841*	
				194*					76AUE/DAV

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₅ Hmno ₅]	(CO) ₅ MnH	RN 16972-33-1				201**	841**	
320	A	(-3.8)						81STE/BEA
[C ₆ H ₁₀]	c-C ₅ H ₈ =CH ₂	RN 1528-30-9		193.4	809.	200.8	840.	
320	A	(-3.2)	192.4					77POL/WOL
320	H		193.4					
[C ₉ H ₁₂]	Mesitylene	RN 108-67-8		193.6	810.	200.7	840.	
320	J	(18.0)	192.6					76DEV/WOL
550	A	(-1.4)	194.6					80MAU
[C ₇ H ₆ N ₂]	m-NCC ₆ H ₄ NH ₂	RN 2237-30-1		192.9	807.	200.7	840.	
600	F	(-9.6)	192.9					01LAU/NIS
[C ₁₁ H ₁₀]	1-Methylnaphthalene	RN 90-12-0		192.9	807.	200.7	840.	
550	A	(-2.3)	192.9					80MAU
[C ₄ H ₈ O ₂]	CH ₃ COOC ₂ H ₅	RN 141-78-6		192.9	807	200.7	840.	
373	X	(0.0)	192.9					79VAJ/HAR
600	X	(0.0)	192.9					76KEB/YAM
320	A	(-3.7)	191.9					77WOL/STA
320	H		192.9					
320	U	(+4.3)	193.0					81BRO/ABB
			193.5**					79AUE/BOW
340	H	(+5.3)	192.6					80LIA/SHO
370	U	(+2.0)	190.7					76HAR/LIN
600	A	(-2.7)	192.5					76YAM/KEB
[C ₂ H ₆ S]	(CH ₃) ₂ S	RN 75-18-3		192.8	807.	200.6	839.	
320	A	(-3.7)	191.9					77WOL/STA-83TAF
320	H		192.8					79AUE/BOW
			193.4**					
[C ₃ H ₆ S]	2-Methylthiirane	RN 1072-43-1		192.8**	807.**	200.6**	839.**	
				192.8**				79AUE/BOW
[C ₉ H ₇ MnO ₃]	(CH ₃ C ₅ H ₄)Mn(CO) ₃	RN 12108-13-3				200.6**	839.**	
320	A	(-4.2)						81STE/BEA
[C ₅ H ₁₀ O ₂]	CH ₃ COOC ₃ H ₇	RN 109-60-4		192.8	807.	200.6	839.	
550	A	(-2.3)	192.8					80MAU
600	A	(-2.3)	192.8					79TAF
[C ₇ H ₁₀]	Bicyclo[2.2.1]hept-2-ene (Norbornene)	RN 498-66-8		193.1	808.	200.4	838.	
320	P (Key)	(0.5)	193.1					77STA/WIE
			191.2			198.5	830.5	76SOL/FIE
			193.8**					79AUE/BOW
560	(Key)		192.6					79SAL/KEB
[C ₆ H ₁₀ O]	(CH ₂ =CHCH ₂) ₂ O	RN 557-40-4		192.6	806.	200.4	838.	
320	A	(-4.0)	191.6					TAF
320	H		192.6					

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
				192.0** 803.**	200.4** 838.**	
			192.0**			79AUE/BOW
				192.5 806.	200.3 838.	
						84SHA/BLA
						76LAU/KEB
						81LAU/NIS
						79LAU
				192.4 805.	200.2 838.	
						77STA/WIE
						77WOL/STA-83TAF
						81BRO/ABB
						79AUE/BOW
						80LIA/SHO
						76HAR/LIN
						84SHA/BLA
						78DAV/LAU
						81LAU/NIS
				192.4 805.	200.2 838.	
						83TAF
						81BRO/ABB
						80MAU
						76LAU/KEB
						79LAU
				192.4 805.	200.2 838.	
						77WOL/STA
				192.4 805.	200.2 838.	
						83TAF
				192.4 805.	200.2 838.	
						TAFT
				192.4 805.	200.2 838.	
						79VAJ/HAR
				192.6 806.	200.1 837.	
						79MAU
						80MAU
					200.1** 837.**	
						81STE/BEA
				192.9 807.	200.0 837.	
						80MAU

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₁₁ H ₁₀]	2-Methylnaphthalene	RN 91-57-6		192.2	804.	200.0	837.	
550	A	(-3.0)	192.2					80MAU
[CCl ₂]	CCl ₂	RN 1605-72-7		~192.2	~804.	~200.0	~837.	
	(br)		~192.2					78AUS/LIA (2)
[C ₇ FH ₆]	3-FC ₆ H ₄ CH ₂ radical	RN xxxxx		~192	~803	~200	~837	
	(br)		~192					82MAU
[C ₄ H ₈ O]	CH ₃ COC ₂ H ₅	RN 78-93-3		192.0	803.	199.8	836.	
320	A	(-4.7)	190.7					83TAF
320	H		191.7					
340	H	(+4.4)	191.6					80LIA/SHO
600	X	(-0.1)	192.8					76KEB/YAM
[C ₅ H ₁₀ O]	c-C ₅ H ₁₀ O	RN 142-68-7		191.9	803.	199.7	835.5	
320	A	(-4.2)	191.4					77WOL/STA
320	H		192.4					
320	U	(+3.4)	192.3					81BRO/ABB
600	U	(+2.8)	191.5					83MAU
[C ₄ H ₆ O]	CH ₃ CH=CHCHO	RN 4170-30-3		191.9	803.	199.7	835.5	
373	(Key)		191.9					79VAJ/HAR
[C ₁₀ F ₃ H ₉]	4-CF ₃ C ₆ H ₄ C(CH ₃)CH ₂	RN 55186-75-9		191.9	803.	199.7	835.5	
320	A	(-4.7)	190.9					TAFT
320	H		191.9					
[C ₅ H ₆]	o-C ₅ H ₆	RN 542-92-7		192.5	805.	199.6	835.	
	(br)		182.8					81HOU/SCH
550	A	(-2.7)	192.5					80MAU
			192.2**					79AUE/BOW
	Threshold Value					197.8	828.	75LOS/TRA
[C ₇ H ₇]	c-C ₇ H ₇ radical	RN 3551-27-7		192.4	805.	199.4	834.	
	(br)		192.4					80DEF/MCI
[C ₃ H ₄ O]	CH ₃ CH=CO	RN 6004-44-0		191.6	802.	199.4	834.	
320	V	(+1.6)	191.6					80ARM/HIG
[C ₁₆ H ₁₀]	Fluoranthene	RN 206-44-0				199.3	834.	
550	A	(-3.0)	192.2					80MAU
[C ₇ FH ₅ O]	4-FC ₆ H ₄ CHO	RN 459-57-4		191.4	801.	199.2	833.	
320	A	(-5.1)	190.5					TAFT
320	H		191.5					
[C ₇ H ₇]	C ₆ H ₅ CH ₂	RN 2154-56-5		191.7	802.	199.1	833.	
	(br)		192.9					82MAU
	(br)		191.7					80DEF/MCI
	Threshold Value		191.3			198.7	832.	78HOU/BEA

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol		Proton affinity kJ/mol	Reference	
[CH ₄ N]		CH ₂ NH ₂	RN 54088-53-8		191	799	199	833	
		(br)		191					81MCA/NIC
[C ₆ H ₁₂]		(CH ₃) ₂ C=C(CH ₃) ₂	RN 563-79-1				199.0	833.	
		(Key)							76GOR/MUN-75SOL/FIE
[C ₅ H ₅]		c-C ₅ H ₅ radical	RN xxxxx		~191	~799	~199	~833	
		(br)		~191					80DEF/MCI
[C ₄ H ₈ O ₂]		1,3-Dioxane	RN 505-22-6		191.2	800.	198.8	832.	
		(Key)		191.2					83MAU
[C ₄ H ₈ O]		c-C ₄ H ₈ O (Tetrahydrofuran)	RN 109-99-9		191.4	801.	198.8	831.	
		320	A	(-4.9)	190.7				77WOL/STA-83TAF
		320	H		191.6				
		320	U	(+2.7)	191.6				81BRO/ABB
		340	H	(+4.0)	191.3				80LIA/SHO
					192.3**				79AUE/BOW
		500	A	(-4.7)	190.4				84SHA/BLA
[C ₅ H ₈ O]		Cyclopentanone	RN 120-92-3		191.5	801.	198.8	832.	
		320	U	(+2.6)	191.5				81BRO/ABB
					192.5**				79AUE/BOW
[C ₇ H ₁₂]		1-Methylcyclohexene	RN 591-49-1		191.0	799.	198.8	832.	
		320	A	(-5.6)	190.0				77POL/WOL
		320	H		191.0				
[C ₇ H ₁₂]		c-C ₅ H ₆ -1,2-(CH ₃) ₂	RN 765-47-9		191.0	799.	198.8	832.	
		320	A	(-5.8)	190.0				77POL/WOL
		320	H		191.0				
[C ₁₄ H ₁₀]		Phenanthrene	RN 85-01-8		191.6	802.	198.7	831.	
		550	A	(-3.6)	191.6				80MAU
[C ₁₈ H ₁₂]		Triphenylene	RN 217-59-4		191.4	001.	190.5	830.5	
		550	A	(-3.8)	191.4				80MAU
[CH ₃ NO]		HCONH ₂	RN 75-12-7		190.6	797.	198.4	830.	
		600	U	(1.9)	190.6				83MAU
[C ₆ H ₁₂]		CH ₃ CH=C(CH ₃)C ₂ H ₅	RN 922-61-2				198.2	829.	
		(Key)							76GOR/MUN-75SOL/FIE
[C ₇ H ₆ O ₂]		C ₆ H ₅ COOH	RN 65-85-0		189.6	793.	198.2	829.	
		600	A	(-5.5)	189.6				79LAU
[C ₄ H ₇ O ₃ P]		Methyltrioxaphospha- bicycloheptane	RN 61580-09-4		190.3	796.	198.1	829.	
		325	(Key)		190.3				80HOD/HOU

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₅ H ₈]	(CH ₃) ₂ CHCCH	RN 598-23-2	190**	190**	795**	198**	828**	79AUE/BOW
[C ₂ H ₂ O]	CH ₂ C=O	RN 463-51-4		189.5	793.	198.0	828.	
	(Key)		189.7					79LIA
	600 A	(-6.3)	189.3					78DAV/LAU
	Threshold Value		189.0			197.3		82TRA/MCL
[C ₃ H ₄]	Cyclopropene	RN 2781-85-3	190*	190*	795*	198*	828*	76AUE/DAV
[C ₆ H ₁₂]	(CH ₃) ₂ C=CHCH ₂ CH ₃	RN 625-27-4	190.1	190.1	795.	197.9	828.	
	340 H	(+2.8)	190.1					78AUS/LIA
	(Key)					197.4		75SOL/FIE
	(Key)					198.0		76GOR/MUN
[C ₃ H ₆ O ₂]	CH ₃ COOCH ₃	RN 79-20-9		190.0	795.	197.8	828.	
	320 V	(0.0)	190.0					80ARM/HIG
	320 A	(-6.4)	189.2					77WOL/STA-83TAF
	320 H		190.2					
	320 U	(+1.2)	190.1					81BRO/ABB
			190.5**					79AUE/BOW
	340 H	(+2.7)	190.0					80LIA/SHO
	340 H	(+2.7)	190.0					78AUS/LIA
	370 U	(+0.2)	188.9					76HAR/LIN
	600 A	(-5.7)	189.4					79LAU
	600 X	(-2.9)	190.0					76KEB/YAM
[C ₅ H ₈]	c-C ₃ H ₅ CH=CH ₂	RN 693-86-7		189.9	794.5	197.7	827.	
	320 A	(-6.7)	188.9					77WOL/STA
	320 H		189.9					
[C ₂ H ₄ N ₂]	NCCH ₂ NH ₂	RN xxxxx		189.6	793.	197.4	826.	
	320 G	(-18.9)	189.6					83TAF
[C ₂ H ₅ NO ₂]	C ₂ H ₅ ONO	RN 109-95-5		189.5	793.	197.3	825.5	
	(br)		189.5					78FAR/MCM
[C ₆ H ₁₀]	1-Methylcyclopentene	RN 693-89-0		189.2	792.	197.0	824.	
	320 A	(-7.6)	188.2					77POL/WOL
	320 H		189.2					79AUE/BOW
			190.4**					
[C ₄ H ₁₀ S]	t-C ₄ H ₉ SH	RN 75-66-1		189.2	792.	197.0	824.	
	320 A	(-7.6)	188.2					TAFT
	320 H		189.2					
[C ₃ H ₆ O]	c-C ₃ H ₆ O (Oxetane)	RN 503-30-0		189.6*	793.*	196.9*	824.*	
			189.6*					79AUE/BOW
[C ₇ C _x H ₅ NO ₃]	(C ₅ H ₅)C _x (CO) ₂ NO	RN 36312-04-6				196.9**	824.**	
	320 A	(-7.7)						81STE/BEA

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₃ H ₆ O]	(CH ₃) ₂ CO	RN 67-64-1		188.9	790.	196.7	823.	
	320 U	(0.0)	188.9					81BRO/ABB
	600 U	(0.0)	188.9					83MAU
	370 U	(0.0)	188.9					76HAR/LIN
	320 A	(-7.9)	187.9					77WOL/STA
	320 H		188.9					
			189.9**					79AUE/BOW
	340 H	(+1.6)	188.9					80LIA/SHO
	340 H	(+1.5)	188.8					78AUS/LIA
	550 A	(-6.9)	188.3					80MAU
	600 A	(-6.5)	188.6					78DAV/LAU
	600 X	(-3.5)	189.4					76KEB/YAM
[C ₄ H ₄ S]	c-C ₄ H ₄ S	Thiophene	RN 110-02-1	189.5	793.	196.5	822.	
	320 A		189.5					TAFT
		(br)	183					81HOU/SCH
	600 (Key)		185.0					83MAU
[C ₇ FH ₅ O]	3-FC ₆ H ₄ CHO	RN 456-48-4		188.7	788.	196.5	822.	
	320 A	(-8.1)	187.7					TAFT
	320 H		188.7					
[C ₅ H ₁₀]	(CH ₃) ₂ C=CHCH ₃	RN 513-35-9		188.6	789.	196.4	822.	
	340 H	(+1.3)	188.6					78AUS/LIA
			189.5**					79AUE/BOW
	(Key)					197.8		76GOR/MUN
	(Key)					198.3		75SOL/FIE
[C ₃ H ₈ O]	CH ₃ OC ₂ H ₅	RN 540-67-0		188.6	789.	196.4	822.	
	320 A	(-8.2)	187.6					77WOL/STA
	320 H		188.6					
			189.2**					79AUE/BOW
[C ₆ H ₆ O]	C ₆ H ₅ OH	RN 108-95-2		188.5	789.	196.3	821.	
	550 A	(-6.3)	188.9					80MAU
	600 J	(+11.3)	186.9					76LAU/KEB
	600 A	(-7.1)	188.1					79LAU
								77DEF/MCI
								For protonation on O atom: PA -175+4 kcal/mol (754 kJ/mol)
[C ₁₂ H ₁₀]	Biphenyl	RN 92-52-4		188.3	788.	196.1	820.	
	550 A	(-6.9)	188.3					80MAU
[C ₇ H ₁₄]	(CH ₃) ₂ C=CHCH(CH ₃) ₂	RN xxxxx				196.1	820.	
	(Key)							76MAU/SOL
[C ₄ H ₈ O ₂]	HCOOCH(CH ₃) ₂	RN 625-55-8		188.2	787.	196.0	820.	
	370 U	(-0.5)	188.2					76HAR/LIN
[C ₂ H ₆ S ₂]	CH ₃ SSCH ₃	RN 624-92-0		~188	~787	~196	~820	
	(br)		~188					81KIM/BON
[C ₅ H ₈]	C ₂ H ₅ CCCH ₃	RN 627-21-4		188**	787**	196**	820**	
			188**					79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₂ H ₃ NS]		CH ₃ SCN	RN 556-64-9		188.1	787.	195.9	820.	
		(br)		188.1					74MCA
[C ₂ H ₃ NS]		CH ₃ NCS	RN 556-61-6		188.1	787.	195.9	820.	
		(br)		188.1					74MCA
[C ₄ H ₈]		(CH ₃) ₂ C=CH ₂	RN 115-11-7		187.3	784.	195.9	820.	
		Threshold Value		187.3					
		320 H	(0.0)	187.3					77WOL/STA
		340 H	(0.0)	187.3					80LIA/SHO
				188.3**					79AUE/BOW
		320 A	(-9.2)	186.4					77WOL/STA
		600 A	(-8.1)	187.0					78DAV/LAU
		600 X	(-5.1)	187.8					76KEB/YAM
[C ₇ H ₅ N]		C ₆ H ₅ CN	RN 100-47-0		188.1	787.	195.9	820.	
		600 J	(+11.4)	187.0					76LAU/KEB
		600 A	(-7.0)	188.1					79LAU
[C ₁₆ H ₁₈]		C ₆ H ₅ (CH ₂) ₄ C ₆ H ₅	RN 1083-56-3		188.1	787.	195.9	820.	
		350 R	(1.8)	188.1					80MAU/HUN
[C ₈ H ₁₀]		m-Xylene	RN 108-38-3		188.1	787.	195.9	820.	
		320 J	(13.5)	188.1					76DEV/WOL
		(Key)		184.2					72CHO/FRA (2)
[C ₄ H ₅ N]		c-C ₃ H ₅ CN	RN 5500-21-0		187.6	785.	195.4	817.5	
		320 A	(-8.9)	186.5					76STA/KLE
		320 H		187.6					
		320 U	(-1.2)	187.7					81BRO/ABB
[C ₄ H ₆ O]		CH ₂ =C(CH ₃)CHO	RN 78-85-3		187.4	784.	195.2	817.	
		373 (Key)		187.4					79VAJ/HAR
[C ₅ H ₁₀ O ₂]		HCO ₂ (n-C ₄ H ₉)	RN 592-84-7		186.9	824.	194.8	815.	
		320 A	(-9.6)	185.9					77WOL/STA
		320 H		187.0					
		340 H	(-0.4)	186.9					80LIA/SHO
				188.0**					79AUE/BOW
[C ₄ H ₆ O ₂]		CH ₃ COCOCH ₃	RN 431-03-8		186.2	779.	194.8	815.	
		300 U	(-2.7)	186.2					83MAU
[C ₁₀ H ₁₂]		1,2,3,4-Tetrahydro- naphthalene	RN 119-64-2		187.7	785.	194.7	815.	
		550 A	(-7.5)	187.7					80MAU
[C ₁₀ H ₈]		Naphthalene	RN 91-20-3		187.8	785.5	194.7	815.	
		550 A	(-7.6)	187.6					80MAU
		600 A	(-7.2)	187.9					78LAU/SAL
[C ₃ H ₆ O]		2-Methyloxirane	RN 75-56-9		186.9	782.	194.7	815.	
				186.9**					79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₁₄ H ₁₄]		C ₆ H ₅ (CH ₂) ₂ C ₆ H ₅	RN 103-29-7		187.3 784.	194.6 814.	
	350	R	(+0.9)	187.3			80MAU/HUN
[C ₂ H ₄ S]		c-C ₂ H ₄ S (Thiirane)	RN 420-12-2		187.3 784.	194.6 814.	
		(Key)		187.3 188.5**			80AUE/WEB 79AUE/BOW
[C ₆ O ₆ V]		(CO) ₆ V	RN 20644-87-5			194.5** 814.**	
	320	A	(-10.3)				81STE/BEA
[C ₄ H ₇ N]		i-C ₃ H ₇ CN	RN 78-82-0		186.4 780.	194.3 813.	
	320	A	(-9.8)	185.7			77WOL/STA
	320	H		186.5			
	320	Q	(+1.0)	186.3			76STA/KLE
[C ₄ H ₈ O ₂]		HCO ₂ (n-C ₃ H ₇)	RN 110-74-7		186.4 780.	194.2 812.5	
	320	A	(-9.8)	187.4** 185.9			79AUE/BOW 77WOL/STA
	320	H		186.7			
	340	U	(-0.6)	186.7			80LIA/SHO
	370	U	(-2.0)	186.8			76HAR/LIN
	600	A	(-9.5)	185.6			79LAU
	600	X	(-6.6)	186.3			76KEB/YAM
[C ₃ H ₈ S]		i-C ₃ H ₇ SH	RN 75-33-2		186.3 779.	194.1 812.	
	320	A	(-10.1)	185.5			Taft
	320	H		186.3			
[B ₃ H ₆ N ₃]		Borazine	RN 6569-51-3		186.3 779.	194.1 812.	
	298	(Key)		186.3			79DOI/GRE
[C ₅ H ₉ N]		n-C ₄ H ₉ CN	RN 110-59-8		186.2 779.	194.0 812.	
	320	A	(-10.2)	185.4			76STA/KLE
	320	H		186.2			
[C ₃ H ₅ O ₃ P]		2,6,7-Trioxa-1-phospha- bicyclo[2.2.1]heptane	RN 279-53-8		186.1 779.	194.0 812.	
	320	(Key)		186.1			80HOD/HOU
[C ₃ H ₄ O]		CH ₂ =CHCHO	RN 107-02-8		186.1 779.	193.9 811.	
	373	Q	(+0.8)	186.1			79VAJ/HAR
[C ₄ H ₆ O]		C-C ₄ H ₆ (=O)	RN xxxxx		186.4 780.	193.8 811.	
	320	U	(-2.5)	186.4			81BRO/ABB
[C ₄ H ₈ O ₂]		1,4-Dioxane	RN 123-91-1		186.0 778.	193.8 811.	
	320	A	(-10.4)	184.8			77WOL/STA
	320	H		186.0			
	320	U	(-2.8)	186.1			81BRO/ABB
	500	A	(-10.5)	184.6			84SHA/BLA
[C ₃ F ₃ H ₆ N]		CF ₃ N(CH ₃) ₂	RN 677-41-8		186. 778.	193.8 811.	
	320	A	(-10.4)	185.2			77STA/TAA
	320	H		186.0 187.0**			79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₄ H ₇ O ₂]	1,4-Dioxyl radical	RN 4598-47-4		186.0	778.	193.8	811.	
340	(Key)		186.0					83AUS/LUT
[C ₄ H ₁₀ O]	t-C ₄ H ₉ OH	RN 75-65-0		185.9	778.	193.7	810.	
600	A (Key)	(-9.2)	187** 185.9 183.2					76AUE/BOW 79LAU 77HIR/KEB
[C ₄ H ₇ N]	n-C ₃ H ₇ CN	RN 109-74-0		185.7	777.	193.7	810.	
320	A	(-10.7)	184.9					76STA/KLE
320	H		185.7 186.0**					79AUE/BOW
[B ₃ H ₅ N ₃]	B-Borazinyl radical	RN xxxxx		185.8	777.	193.6	810.	
	(br)		185.8					76DES/POR
[C ₆ H ₅ NO ₂]	C ₆ H ₅ NO ₂	RN 98-95-3		185.6	776.5	193.4	809.	
600	J	(+8.9)	184.5					76LAU/KEB
600	A	(-9.5)	185.6					79LAU
[C ₈ H ₁₀]	o-Xylene	RN 95-47-6		186.1	779.	193.3	809.	
350	R	(0.0)	186.4					80MAU/HUN
320	A	(-10.6)	185.0					74HEH/MCI
320	H		185.8					
320	J	(9.6)	184.2 184.2					76DEV/WOL 72CHO/FRA(2)
550	A	(-8.8)	186.4					80MAU
[C ₃ H ₆ O ₂]	HCO ₂ C ₂ H ₅	RN 109-94-4		185.3	775.	193.1	808.	
320	Q	(0.0)	185.3 186.4**					76STA/KLE 79AUE/BOW
320	A	(-11.0)	184.6					77WOL/STA
320	H		185.4					
320	U	(-3.3)	185.6					81BRO/ABB
340	H	(-2.4)	184.9					80LIA/SHO
370	U	(-2.0)	186.7					76HAR/LIN
600	A	(-10.0)	185.6					78DAV/LAU
[C ₄ H ₆]	CH ₂ =CHCH=CH ₂	RN 106-99-0		185**	774**	193**	807.5**	
			185**					79AUE/BOW
[C ₁₀ H ₁₄]	t-C ₄ H ₉ C ₆ H ₅	RN 98-06-6		185.2	775.	193.0	807.	
320	A	(-11.2)	184.4					74HEH/MCI
320	H		185.2					
[C ₃ H ₅ N]	C ₂ H ₅ CN	RN 107-12-0		184.1	770.	192.6	806.	
320	A	(-12.0)	183.6					76STA/KLE
320	H		184.4					
340	H	(-3.4)	183.9 185.0**					80LIA/SHO 79AUE/BOW
[C ₄ H ₈ O]	i-C ₃ H ₇ CHO	RN 78-84-2		184.8	773.	192.6	806.	
320	A	(-11.6)	184.0					77WOL/STA
320	H		184.8 185.5**					79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₅ H ₁₀ O]	n-C ₄ H ₉ CHO	RN 110-62-3		184.8	773.	192.6	806.	
320	A	(-11.6)	184.0					77WOL/STA
320	H		184.8					79AUE/BOW
			185.5**					
[CH ₃ NO ₂]	CH ₃ ONO	RN 624-91-9		184.7	773.	192.5	805.	
	(br)		184.7					78FAR/MCM
	(br)		184.7					76MCA/PIT
[C ₉ H ₁₂]	n-C ₃ H ₇ C ₆ H ₅	RN 103-65-1		184.6	772.	192.4	805.	
320	A	(-11.8)	183.8					74HEH/MCI
320	H		184.6					
600	J	(+7.5)	183.1					76YAM/KEB
[C ₄ H ₄ O]	Furan	RN 110-00-9		185.0	774.0	192.2	804.	
600	(Key)		185.0					83MAU
	(br)		183					81HOU/SCH
[C ₉ H ₁₂]	i-C ₃ H ₇ C ₆ H ₅	RN 98-82-8		184.3	771.	192.1	804.	
320	A	(-11.3)	184.3					74HEH/MCI
320	H		185.1					
600	J	(+7.9)	183.5					76YAM/KEB
[C ₁₀ H ₁₄]	n-C ₄ H ₉ C ₆ H ₅	RN 104-51-8		184.3	771.	192.1	804.	
320	A	(-11.4)	184.2					74HEH/MCI
320	H		185.0					
600	J	(+7.9)	183.5					76YAM/KEB
[C ₂ H ₆ O]	(CH ₃) ₂ O	RN 115-10-6		184.3	771.	192.1	804.	
320	A	(-11.7)	183.9					77WOL/STA-83TAF
320	H		184.7					
320	U	(-4.0)	184.9					81BRO/ABB
340	H	(-3.2)	184.1					80LIA/SHO
370	T	(+4.4)	182.8					76HAR/LIN
373	T	(+4.5)	183.1					75SOL/HAR
500	A	(-11.0)	184.1					84SHA/BLA
600	A	(-11.1)	184.0					76YAM/KEB
600	X	(-7.7)	185.2					76KEB/YAM
			185.8**					79AUE/BOW
[C ₃ FH ₅ O]	CH ₃ COCH ₂ F	RN 430-51-3		184.2	771.	192.0	803.	
298	(Key)		184.2					82DRU/MCM
[C ₈ H ₁₀]	p-Xylene	RN 106-42-3		184.6	772.	192.0	803.	
320	A	(-11.3)	184.3					74HEH/MCI
320	H		185.1					
320	J	(+8.9)	184.5					76DEV/WOL
	(Key)		184.1					72CHO/FRA (2)
[C ₃ H ₆ O ₂]	C ₂ H ₅ COOH	RN 79-09-4		184.0	770.	191.8	802.	
600	A	(-11.8)	183.3					76YAM/KEB
600	X	(-8.2)	184.7					76KEB/YAM
[C ₃ H ₈ S]	n-C ₃ H ₇ SH	RN 107-03-9		183.8	769.	191.6	802.	
320	A	(-12.6)	183.0					TAFT
320	H		183.8					

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₈ H ₁₀]	C ₂ H ₅ C ₆ H ₅	RN 100-41-4		183.8	769.	191.6	802.	
	320 A	(-12.7)	182.9					77WOL/STA, 74HEH/MCI
	320 H		183.7					
	600 J	(+7.3)	182.9					76LAU/KEB
	600 A	(-11.1)	184.0					79LAU
[C ₄ F ₉ H ₂ N]	(CF ₃) ₃ CNH ₂	RN 2809-92-9		183.1**	766.**	191.5**	801.**	
			183.1**					79AUE/BOW
[C ₄ H ₈ O]	n-C ₃ H ₇ CHO	RN 123-72-8		183.7	769.	191.5	801.	
	320 A	(-12.6)	183.0					77WOL/STA
	320 H		183.8					
	340 H	(-3.7)	183.6					80LIA/SHO
			185.8**					79AUE/BOW
[C ₂ H ₅ P]	c-C ₂ H ₄ PH (Phosphirane)	RN 6569-82-0		184.2	771.	191.4	801.	
	(Key)		184.2					80AUE/WEB
			187.6**					79AUE/BOW
[C ₃ H ₈ O]	i-C ₃ H ₇ OH	RN 67-63-0		183.4	767.	191.2	800.	
	600 A	(-11.7)	183.4					79LAU
[C ₄ H ₁₀ O]	n-C ₄ H ₉ OH	RN 71-36-3		183.3	767.	191.1	799.5	
	320 A	(-13.1)	182.5					TAFT
	320 H		183.3					
[C ₈ F ₃ H ₅ O]	p-CF ₃ C ₆ H ₄ CHO	RN 455-19-6		183.2	766.5	191.0	799.	
	320 A	(-13.2)	182.4					TAFT
	320 H		183.2					
[C ₄ H ₆]	Cyclobutene	RN 822-35-5		183**	766**	191**	799**	
			183**					79AUE/BOW
[C ₂ H ₆ S]	C ₂ H ₅ SH	RN 75-08-1		182.9	765.	190.8	798.	
	320 A	(-13.8)	181.8					TAFT
	320 H		182.6					
	340 H	(-4.0)	183.3					80LIA/SHO
[C ₆ H ₆ O]	(HCCCH ₂) ₂ O	RN 6921-27-3		183.0	766.	190.8	798.	
	320 A	(-13.4)	182.2					TAFT
	320 H		183.0					
[C ₃ H ₈ O]	n-C ₃ H ₇ OH	RN 71-23-8		183.0	766.	190.8	798.	
	320 U	(-5.9)	183.0					81BRO/ABB
			183.6**					79AUE/BOW
[C ₂ D ₆ O]	(CD ₃) ₂ O	RN 17222-37-6		182.8	765.	190.6	797.	
	370 T	(+4.2)	182.8					76HAR/LIN
[C ₈ FeH ₈ O ₂]	(C ₅ H ₅)Fe(CO) ₂ CH ₃	RN 12080-06-7				190.6**	797.**	
	320 A	(-14.2)						81STE/BEA
[CHN]	HNC	RN xxxxx		182.4	763.	190.2	796.	
	(br)		182.4					82PAU/HEH

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	kJ/mol	Proton affinity kcal/mol	kJ/mol	Reference
[C ₂ H ₄ O ₂]	CH ₃ COOH	RN 64-19-7		181.7	760.	190.2	796.	
	320 A	(-14.8)	180.8					77WOL/STA
	320 H		181.6					
	600 A	(-13.3)	181.8					76YAM/KEB
[C ₇ H ₈]	C ₆ H ₅ CH ₃	RN 108-88-3		182.0	761.	189.8	794.	
	320 A	(-13.7)	181.9					74HEH/MCI
	320 H		182.7					
	320 J	(+6.6)	181.2					76DEV/WOL
	340 (Key)		182.1					77AUS/LIA
	(Key)		184.1					72CHO/FRA (2)
	478 W	(0.0)	182.0					82STO/SPL
	550 A	(-11.0)	184.2					80MAU
	600 J	(+6.3)	181.9					76LAU/KEB
	600 A	(-12.1)	183.0					79LAU
[C ₇ D ₃ H ₅]	C ₆ H ₅ CD ₃	RN 1124-18-1		182.0	761.	189.8	794.	
	340 (Key)		182.1					77AUS/LIA
[C ₃ H ₃ N]	CH ₂ =CHCN	RN 107-13-1		181.9	761.	189.7	794.	
	320 A	(-14.5)	181.1					76STA/KLE
	320 H		181.9					
[C ₃ H ₆ O]	C ₂ H ₅ CHO	RN 123-38-6		181.8	761.	189.6	793.	
	320 A	(-14.4)	181.2					77WOL/STA
	320 H		182.0					
	300		183.6**					79AUE/BOW
	340 H	(-5.6)	181.7					80LIA/SHO
	370 T	(+2.5)	181.1					75SOL/HAR
	600 A	(-13.4)	181.7					76YAM/KEB
[C ₇ FH ₇]	3-FC ₆ H ₄ CH ₃	RN 352-70-5		181.8	761.	189.3	792.	
	478 W	(-0.2)	181.8					82STO/SPL
[C ₆ H ₁₀]	c-C ₆ H ₁₀	RN 110-83-8		181.5	759.	189.3	792.	
	340 H	(-5.8)	181.5					80LIA/SHO
[C ₂ H ₄ O ₂]	HCO ₂ CH ₃	RN 107-31-3		181.0	757.	188.9	790.	
	320 A	(-14.7)	180.9					77WOL/STA
	320 H		181.7					
	340 H	(-6.0)	181.2					80LIA/SHO
	370 T	(+2.0)	180.6					76HAR/LIN
	478 W	(-1.3)	180.2					82STO/SPL
	600 A	(-13.5)	181.6					79LAU
[H ₃ P]	PH ₃	RN 7803-51-2		180.2	754.	188.6	789.	
	320 A	(-15.8)	179.8					77WOL/STA-83TAF
	320 H		180.5					
	340 H	(-7.4)	179.9					80LIA/SHO
			182.5**					79AUE/BOW
[C ₂ H ₃ N]	CH ₃ CN	RN 75-05-8		180.6	756.	188.4	788.	
	320 A	(-15.5)	180.1					77WOL/STA
	320 H		180.9					
	340 H	(-6.9)	180.4					80LIA/SHO
	600 A	(-14.5)	180.6					79LAU
			183.1**					79AUE/BOW

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	kJ/mol	Proton affinity kcal/mol	kJ/mol	Reference
[C ₂ H ₆ O]		C ₂ H ₅ OH	RN 64-17-5		180.2	754.	188.3	788.	
	320	A	(-15.8)	179.8					77WOL/STA-83TAF
	320	H		180.6					
	370	T	(+1.3)	179.9					76HAR/LIN
	600	A	(-14.7)	180.4					79LAU
				182.5**					79AUE/BOW
[C ₃ H ₅]		c-C ₃ H ₅ radical	RN xxxxx		179.	749.	188.	787.	
		(br)		179.					80DEF/MCT
[B ₄ H ₈]		B ₄ H ₈	RN 12007-71-5		180	753	188	787	
		(br)		180					72SOL/POR
[C ₂ H ₄ O]		c-C ₂ H ₄ O (Oxirane)	RN 75-21-8		180.6	756.	187.9	786.	
		(Key)		180.6					80AUE/WEB
				182.3**					79AUE/BOW
[C ₃ ClH ₄ N]		Cl(CH ₂) ₂ CN	RN 542-76-7		179.9	752.	187.5	784.5	
	320	A	(-16.4)	179.2					76STA/KLE
	320	H		179.9					
[CH ₄ S]		CH ₃ SH	RN 74-93-1		179.2	750.	187.4	784.	
	320	A	(-16.6)	179.0					77WOL/STA
	320	H		179.7					
	340	H	(-7.9)	179.4					80LIA/SHO
	370	T	(+0.4)	178.7					76HAR/LIN
	370	(Key)		179.1					75SOL/HAR
[H ₂ N]		NH ₂	RN 15194-15-7		179	749	187	782	
		(br)		179.					82DEF/HEH
[C ₄ H ₆]		CH ₃ CCCH ₃	RN 503-17-3		179**	749**	187**	782**	
				179**					79AUE/BOW
[C ₈ H ₅ NO]		4-CNC ₆ H ₄ CHO	RN 105-07-7		179.2	750.	187.0	782.	
	320	A	(-17.1)	178.5					TAFT
	320	H		179.2					
[C ₃ F ₂ H ₄ O]		CFH ₂ COCFH ₂	RN 453-14-5		179	749	187	782	
	298	(Key)		179					82DRU/MCM
[C ₆ H ₃ O ₅ Re]		(CO) ₅ ReCH ₃	RN 14524-92-6				187**	782**	
	320	A	(-17.4)						81STE/BEA
[C ₇ FH ₇]		2-FC ₆ H ₄ CH ₃	RN 95-52-3		178.8	748.	186.6	781.	
	478	W	(-3.2)	178.8					82STO/SPL
[C ₂ H ₄ O]		CH ₃ CHO	RN 75-07-0		178.6	747.	186.6	781.	
		Threshold Value					184.6	772.	68REF/CHU
	370	T	(0.0)	170.6					76HAR/LIN
	370	T	(0.0)	178.6					75SOL/HAR
	320	A	(-17.6)	178.0					77WOL/STA
	320	H		178.7					
	300			181.1**					79AUE/BOW
	340	H	(-0.0)	170.5					80LIA/SHO
	600	A	(-16.1)	179.0					79LAU
	600	N		177.4					

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol kJ/mol	Proton affinity kcal/mol kJ/mol	Reference
[C ₄ F ₃ H ₇ O]		C ₂ H ₅ OCH ₂ CF ₃	RN 461-24-5		178.6 747.	186.4 780.	
	320	A	(-17.7)	177.9			TAFT
	320	H		178.6			
[C ₃ H ₄]		H ₂ C=C=CH ₂	RN 463-49-0		179. 749.	186.3 779.	
		Threshold Value (br)		179.		185.3 775.	77ROS/DRA AUS/LIA
[CH ₂ S]		CH ₂ S	RN 865-36-1		178. 745.	186. 778.	
		(br)		178			82ROY/MCM
[C ₂ H ₆ Hg]		CH ₃ HgCH ₃	RN 593-74-8		-178 -744	-186 -778	
		(br)		-178			80STO/CAM
[C ₇ FH ₇]		4-FC ₆ H ₄ CH ₃	RN 352-32-9		178.0 745.	185.8 777.	
	478	W	(-4.0)	178.0			82STO/SPL
[C ₆ F ₃ H ₉ O ₂]		CF ₃ CO ₂ (n-C ₄ H ₉)	RN 367-64-6		178.0 745.	185.8 777.	
	320	A	(-18.2)	177.3			77WOL/STA
	320	H		178.0			
[C ₅ F ₃ H ₇ O ₂]		CF ₃ CO ₂ (n-C ₃ H ₇)	RN 383-66-4		177.9 744.	185.7 777.	
	320	A	(-18.4)	177.2			77WOL/STA
	320	H		177.9			
[C ₃]		C ₃	RN 12075-35-3 (br)	-177	-742	-185 -775	83RAK/BOH
[C ₆ MoO ₆]		(CO) ₆ Mo	RN 13939-06-5			185** 774**	
	320	A	(-19.2)				81STE/BEA
[C ₂ H ₅ NO ₂]		C ₂ H ₅ NO ₂	RN 79-24-3		177.0 740.	184.8 773.	
	340	H	(-10.3)	177.0			80LIA/SHO
[C ₄ F ₃ H ₅ O ₂]		CF ₃ CO ₂ C ₂ H ₅	RN 383-63-1		176.8 740.	184.6 772.	
	320	A	(-19.5)	176.1			77WOL/STA
	320	H		176.8			
[C ₃ HN]		HCCCN	RN xxxxx (br)	176.	176. 737.	184. 770.	84RAK/BOH
[B ₅ H ₈]		B ₅ H ₈	RN 65930-58-7 (br)	177.	177. 740.	184. 770.	78WAN/DES
[C ₆ O ₆ W]		(CO) ₆ W	RN 14040-11-0			184** 770**	
	320	A	(-20.3)				81STE/BEA
[C ₂ FH ₃ O ₂]		CH ₂ FCOOH	RN 144-49-0		175.7 735.1	183.5 768.	
	600	A	(-19.4)	175.7			76YAM/KEB
[C ₂ Cl ₃ HO ₂]		CCl ₃ COOH	RN 76-03-9		175.7 735.	183.5 768.	
	600	A	(-18.3)	176.8			76YAM/KEB
		N		175.7			
[C ₅ H ₈]		c-C ₅ H ₈	RN 142-29-0			183.4 767.5	
		Threshold Value (br)		178.2		183.6 768.	HOU/BEA
		(Key)				186.0 778.	80LIA/SHO
						183.3 767.	76SOL/FIE

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₆ H ₃ MnO ₅]	(CO) ₅ MnCH ₃	RN 13601-24-6		175	732	183	766	
320	A (br)	(-19.9)	175.			185**		81STE/BEA 79STE/BEA
[C ₆ FH ₅]	Fluorobenzene	RN 462-06-6		174.8	731.4	182.6	764.	
334	J	(+0.2)	174.8					81BOH/STO
400	J	(-0.2)	174.6					78HAR/LIA
600	J	(-0.8)	174.8					76LAU/KEB
600	A	(-19.2)	175.9					79LAU
600	N		174.8					
[BrC ₆ H ₅]	Bromobenzene	RN 108-86-1		174.6	730.5	182.4	763.	
334	J	(0.0)	174.6					81BOH/STO
[C ₂ ClH ₃ O ₂]	CH ₂ ClCOOH	RN 79-11-8		174.6	730.5	182.4	763.	
600	A N	(-19.4)	175.7 174.6					76YAM/KEB
[C ₃ H ₄]	CH ₃ CCH	RN 74-99-7		174*	728*	182*	761*	
			174*					76AUE/DAV
[CH ₄ O]	CH ₃ OH	RN 67-56-1		174.1	728.	181.9	761.	
320	A	(-20.6)	175.0					77WOL/STA-83TAF
320	L		173.7					
340	H	(-12.9)	174.4					80LIA/SHO
600	A	(-19.7)	175.4					79LAU
600	N		174.3					
[C ₆ F ₂ H ₄]	1,2-Difluorobenzene	RN 367-11-3		174.4	729.7	181.8	761.	
400	J	(-0.4)	174.4					78HAR/LIA
[C ₆ ClH ₅]	Chlorobenzene	RN 108-90-7		174.4	730.	181.7	760.	
334	J	(-0.2)	174.4					81BOH/STO
400	J	(-0.5)	174.3					78HAR/LIA
600	J	(-1.0)	174.1					76LAU/KEB
600	A	(-19.4)	175.7					79LAU
600	N		174.6					
[C ₆ F ₂ H ₄]	1,3-Difluorobenzene	RN 372-18-9		174.1	728.	181.5	759.	
400	J	(-0.3)	174.5					78HAR/LIA
600	A N	(-20.3)	175.3 173.7					76YAM/KEB
[C ₆ F ₃ H ₃]	1,2,4-C ₆ H ₃ F ₃	RN 367-23-7		173.6	726.	181.4	759.	
400	J	(-1.2)	173.6					78HAR/LIA
[C ₆ H ₆]	Benzene	RN 71-43-2		174.6	730.5	181.3	758.5	
320	J	(0.0)		174.6	730.5			76DEV/WOL
334	J	(0.0)		174.6	730.5			81BOH/STO
400	J	(0.0)		174.8	731.4			78HAR/LIA
600	J	(0.0)		175.6	734.7			76LAU/KEB
320	A	(-20.3)	175.3					77WOL/STA
320	H		176.0					
340	H	(-12.1)	175.2					80LIA/SHO
550	A	(-18.9)	176.7					80MAU
600	A	(-18.4)	177.2					78LAU/SAL
600	N		175.6					

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₆ F ₂ H ₄]	1,4-C ₆ H ₄ F ₂	RN 540-36-3			173.8	727.	181.2	758.	
	400	J	(-1.0)	173.8					78HAR/LIA
[C ₆ F ₄ H ₂]	1,2,3,4-C ₆ H ₂ F ₄	RN 551-62-2			173.3	725.	181.1	758.	
	400	J	(-1.5)	173.3					78HAR/LIA
[C ₆ F ₃ H ₃]	1,3,5-C ₆ H ₃ F ₃	RN 372-38-3			173.7	727.	181.	757.	
	400	J	(-0.7)	174.1					78HAR/LIA
	600	A	(-20.7)	174.4					79LAU
	600	N		173.3					
[C ₂ H ₃]	C ₂ H ₃ radical	RN 2669-89-8			~172	~720	~181	~757	
	(br)			~172					80DEF/MCI
[C ₆ F ₄ H ₂]	1,2,3,5-C ₆ H ₂ F ₄	RN 2367-82-0			173.2	725.	180.6	756.	
	400	J	(-1.6)	173.2					78HAR/LIA
[C ₄ NiO ₄]	(CO) ₄ Ni	RN 13463-39-3					180**	753**	
	320	A	(-24.5)						81STE/BEA
[C ₆ CrO ₆]	(CO) ₆ Cr	RN 13007-92-6					180**	753**	
	320	A	(-24.8)						81STE/BEA
[C ₆ F ₅ H]	C ₆ HF ₅	RN 363-72-4			172.5	722.	179.9	753.	
	400	J	(-2.3)	172.5					78HAR/LIA
[C ₃ H ₆]	c-C ₃ H ₆	RN 75-19-4			172.0	720.	179.8	752.	
	340	(Key)		173.7					72CHO/FRA
[C ₆ F ₄ H ₂]	1,2,4,5-C ₆ H ₂ F ₄	RN 327-54-8			173.0	724.	179.7	752.	
	400	J	(-1.8)	173.0					78HAR/LIA
[C ₃ H ₆]	CH ₃ CH=CH ₂	RN 115-07-1			171.7	718.	179.5	751.	
	Threshold Value			171.7			179.5		82ROS/BUF
	340	H	(-14.0)	173.3					80LIA/SHO
	340	AA	(0.0)	171.7					
	340	(Key)		173.4					72CHO/FRA
	600	A	(-20.7)	174.9					76YAM/KEB
	600	N		173.3					
[C ₄ H ₅ NO ₂]	NCCOOC ₂ H ₅	RN 623-49-4			171.7	718	179.5	751.	
	320	A	(-22.0)	174.6					77WOL/STA
	320	L		171.7					
[C ₂ ClH ₂ N]	ClCH ₂ CN	RN 107-14-2			171.7	718.	179.5	751.	
	320	A	(-22.0)	174.6					77WOL/STA
	320	L		171.7					
[C ₄ H ₈]	(E)-CH ₃ CH=CHCH ₃	RN 624-64-6			171.6	718.	179.4	751.	
	Threshold Value								81TRA
	340	H	(-14.1)	173.2					80LIA/SHO
	340	AA		171.6					

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₃ F ₃ H ₃ O ₂]		HCOUCH ₂ CF ₃	RN 32042-38-9		171.6	718.	179.4	751.	
	320	A	(-22.1)	174.5					77WOL/STA
	320	L		171.6					
[CH ₃ NO ₂]		CH ₃ NO ₂	RN 75-52-5		171.7	718.	179.2	750.	
	298	(Key)		172.1					78MAC/BOH
	340	H	(-14.1)	173.2					80LIA/SHO
	340	AA		171.4					76MCA/PIT
		(Key)		~173					
[AsH ₃]		AsH ₃	RN 7784-42-1		170.8	715.	179.2	750.	
	320	A	(-22.8)	173.8					77WOL/STA
	320	L		170.8					
	340	H	(-15.1)	172.2					80LIA/SHO
	340	AA		170.8					
[C ₃ F ₃ H ₃ O ₂]		CF ₃ COOCH ₃	RN 431-47-0		171.0	715.	178.8	748.	
	320	A	(-22.6)	174.0					77WOL/STA
	320	L		171.0					
[CH ₂ O ₂]		HCOOH	RN 64-18-6		170.4	713.	178.8	748.	
	320	A	(-23.2)	173.4					78WOL/STA
	320	L		170.4					
	340	H	(-15.4)	171.9					80LIA/SHO
	340	AA		170.3					
	600	A	(-23.6)	172.0					79LAU
	600	N		170.4					
[C ₄ F ₄ H ₄ O ₂]		CF ₃ COOCH ₂ CH ₂ F	RN 1683-88-1		170.8	715.	178.6	747.	
	320	A	(-22.8)	173.8					77WOL/STA
	320	L		170.8					
[BrCN]		BrCN	RN 506-68-3		170.5	713.	178.3	746.	
	320	A	(-23.1)	173.5					76STA/KLE
	320	L		170.5					
[C ₆ F ₆]		C ₆ F ₆	RN 392-56-3		171.3	717.	177.7	743.	
	400	J	(-3.5)	171.3					78HAR/LIA
[C ₂ Cl ₃ H ₃ O]		Cl ₃ CCH ₂ OH	RN 115-20-8		169.6	710.	177.4	742.	
	320	A	(-24.0)	172.6					77WOL/STA
	320	L		169.6					
[C ₂ F ₂ H ₄ O]		CF ₂ HCH ₂ OH	RN 359-13-7		168.4	704.5	176.2	737.	
	320	A	(-25.2)	171.3					77WOL/STA
	320	L		168.4					
[C ₂ F ₂ H ₂]		CH ₂ CF ₂	RN 75-38-7		168	703	176	736	
		(br)		~172					75RID
		(Key)		168					76WIL/LEB
[C ₂ H ₅ I]		C ₂ H ₅ I	RN 75-03-6		~168	~703	~176	~736	
		(br)		~168					72BEA/HOL

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₂ Cl ₃ N]	CCl ₃ CN	RN 545-06-2		168.0	703.	175.8	735.5	
320	A	(-25.6)	171.0					77WOL/STA
320	L		168.0					
[C ₃ H ₃]	c-C ₃ H ₃ radical	RN xxxxx		168.7	706.	175.8	735.	
	(br)		168.7					80DEF/MCI
[C ₃ H ₅]	CH ₂ =CHCH ₂ radical	RN xxxxx		168.7	706.	175.8	735.	
	(br)		168.7					80DEF/MCI
[CClN]	ClCN	RN 506-77-4		167.9	702.	175.7	735.	
320	A	(-25.7)	170.9					76STA/KLE
320	L		167.9					
[C ₃ H ₂ N ₂]	CH ₂ (CN) ₂	RN 109-77-3		167.4	700.	175.6	735.	
298	L	(0.0)	167.4					81DOI/MCM
298	L	(0.0)	167.4					82DRU/MCM
320	A	(-25.8)	170.8					77WOL/STA
320	L		167.4					
600	A	(-26.1)	169.0					79LAU
600	N		167.9					
[C ₂ FH ₃]	C ₂ H ₃ F	RN 75-02-5		167.	699.	175.	732.	
	(br)		~163					75RID
	(Key)		167.					76WIL/LEB
[CS]	CS	RN 2944-05-0		167	699	175	732	
	(br)		167					78MCA
[C ₃ F ₃ H ₃ O]	CH ₃ COCF ₃	RN 421-50-1		166.4	696.	174.2	729.	
298	L	(-0.6)	166.8					81DOI/MCM
298	L	(-1.0)	166.4					82DRU/MCM
[CHNO]	HNCO	RN 75-13-8		165.5	692.	173.3	725.	
320	M	(+1.2)	165.5					80WIG/BEA
[CF ₂]	CF ₂	RN 2154-59-8		164.0	686.	171.9	719.	
	(br)		164.0					77LIA/AUS
[CH ₂ O]	H ₂ CO	RN 50-00-0		164.3	687.	171.7	718.	
320	M	(0.0)	164.3					80WIG/BEA
298	N	(5.2)	164.2					78TAN/MAC
300	Y	(1.2)	160.2					78FRE/HAR
320	A	(-28.3)	168.3					77WOL/STA
320	N		164.3					
[CHN]	HCN	RN 74-90-8		163.8	685.	171.4	717.	
298	N	(4.9)	163.9					78TAN/MAC
320	A	(-28.8)	167.7					77WOL/STA
320	N		163.8					
340	Y	(1.0)	163.8					78FRE/HAR(2)

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[H ₂ Se]	H ₂ Se	RN 7783-07-5			163.8	685.	171.3	717.	
	320	A	(-28.8)	167.7					77WOL/STA
	320	N		163.8					79AUE/BOW
				169.4**					
[CH ₃ I]	CH ₃ I	RN 74-88-4			~163	682	~171	~715	
		(br)		~163					72BEA/HOL
[C ₂ BrH ₅]	C ₂ H ₅ Br	RN 74-96-4			~163	~682	~171	~715	
		(br)		~163					72BEA/HOL
[H ₂ S]	H ₂ S	RN 7783-06-4			162.8	681.	170.2	712.	
	340	Y	(0.0)	162.8					78FRE/HAR(2)
	296	N	(4.6)	163.6					73HOP/BON
	298	N	(3.8)	162.8					78TAN/MAC
	320	A	(-29.6)	167.0					77WOL/STA
	320	N		162.9					
	550	N	(3.9)	162.7					77MAU/FIE
	600	A	(-31.8)	163.8					79LAU
	600	N		162.2					
		Threshold Value					168.4	705.	83PRE/TZE
[C ₃ F ₄ H ₂ O]	CF ₂ HCOCF ₂ H	RN 360-52-1			162.	678.	170.	711.	
		(br)		162.					82DRU/MCM
[CF ₃ NO]	CF ₃ NO	RN 334-99-6			161	674	169	707	
		(br)		161.					79FRE/HAR
[H ₂ O ₄ S]	H ₂ SO ₄	RN 7664-93-9			~161	~674.	~169	~707	
		(br)		~161					78SMI/MUN
[CF ₃ HO ₃ S]	CF ₃ SO ₃ H	RN 1493-13-6			~161	~674	~169	~707	
		(br)		~161					78SMI/MUN
[C ₂ F ₃ H ₃ O]	CF ₃ CH ₂ OH	RN 75-89-8			161.2	674.	169.0	707.	
	298	K	(5.6)	161.2					83COL/MCM
	320	A	(-31.2)	165.4					77WOL/STA
		N		161.3					
	600	A	(-32.7)	162.9					79LAU
		N		161.2					
[C ₂ F ₃ H]	C ₂ F ₃ H	RN 359-11-5			~161	~674	~169	~707	
		(br)		~161					75RID
[C ₆ H ₁₂]	c-C ₆ H ₁₂	RN 110-82-7			~161	~674	~169	~707	
		(br)		~161					82AUS/REB
[C ₂ F ₃ HO ₂]	CF ₃ COOH	RN 76-05-1			161.1	674.	169.0	707.	
	298	(Key)		161.0					82DRU/MCM
	320	A	(-30.5)	165.4					77WOL/STA
		N		162.0					
	600	A	(-32.9)	162.7					79LAU
		N		161.1					

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[B ₅ H ₉]	B ₅ H ₉	RN 19624-22-7			161.	674.	169.	707.	
	(br)			161.					72SOL/POR
[C ₂ ClH ₅]	C ₂ H ₅ Cl	RN 75-00-3			161.	674.	169.	707.	
	(br)			161.					72BEA/HOL
[B ₅ C ₂ H ₇]	2,4-C ₂ B ₅ H ₇	RN 20693-69-0			160.	669.	168.	703.	
	(br)			160.					80DIX
[F ₃ OP]	OPF ₃	RN 13478-20-1			160.0	669.	167.8	702.	
	298 K	(4.4)	160.0						83COL/MCM
[C ₄ F ₇ N]	C ₃ F ₇ CN	RN 375-00-8			159.6	668.	167.4	700.	
	298 K	(4.0)	159.6						83COL/MCM
[CS ₂]	CS ₂	RN 75-15-0			160.7	672.	167.1	699.	
	550 N	(1.9)	160.7						77MAU/FIE
[C ₃ F ₅ N]	C ₂ F ₅ CN	RN 422-04-8			159.3	666.5	167.1	699.	
	298 K	(3.7)	159.3						83COL/MCM
[C ₄ F ₆ H ₄ O]	(CF ₃) ₂ C(CH ₃)OH	RN 1515-14-6			159.2	666.	167.0	699.	
	298 K	(3.6)	159.2						83COL/MCM
[H ₂ O]	H ₂ O	RN 7732-18-5			159.0	665.	166.5	697.	
	298 K	(3.4)	159.0						83COL/MCM
	298 K	(1.8)	157.4						81BOH/MAC
	320 A	(-33.5)	163.1						77WOL/STA
	320 N	(0.0)	159.0						
	600 A	(-34.7)	160.9						79LAU
		(0.0)	159.3						
	296 N	(0.0)	159.0						73HOP/BON
	298 N	(0.0)	159.0						78TAN/MAC
	550 N	(0.0)	159.3						77MAU/FIE
	Threshold Value		160.3				167.8		77NG/TRE
	Threshold Value		157.4				164.9		69HAN/FRA
[C ₂ F ₃ N]	CF ₃ CN	RN 353-85-5			158.4	663.	166.1	695.	
	298 K	(2.8)	158.4						83COL/MCM
	320 A	(-37.1)	158.5						78COR/BEA
[C ₂ F ₂ H ₂]	(E)-CHFCHF	RN 1630-78-0			158	661	166	694	
	(br)			158					75RID
[BrCH ₃]	CH ₃ Br	RN 74-83-9			157.9	661.	165.7	693.	
	(br)			157.					72BEA/HOL
	298 K	(2.3)	157.9						83COL/MCM
[C ₂ F ₃ HO]	CF ₃ CHO	RN 75-90-1			157.3	658.	165.1	691.	
	298 K	(1.6)	157.2						83COL/MCM
	320 A	(-38.2)	157.4						78COR/BEA

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[C ₂ FH ₅]	C ₂ H ₅ F	RN 353-36-6 (br)		157.	157.	157.	657. 165.	690. 72BEA/HOL
[C ₃ F ₆ H ₂ O]	(CF ₃) ₂ CHOH	RN 920-66-1 298 K	(1.6)	157.2	157.2	157.2	658 165.0	690. 83COL/MCM
[C ₄ H ₁₀]	iso-C ₄ H ₁₀	RN 75-28-5 100 (Key) (Key)		156.0 162.7	156.0 162.7	156.0 162.7	651. 163.3 163.8 162.7	683. 76HIR/KEB 78HIR
[F ₃ P]	PF ₃	RN 7783-55-3 298 K (br) (br)	(3.1)	158.7 154.9 155.2	158.7 154.9 155.2	158.7 154.9 155.2	648. 163.3	683. 83COL/MCM 80DOI/MCM 78COR/BEA-72BEA/HOL
[C ₄ F ₉ HO]	(CF ₃) ₃ COH	RN 2378-02-1 298 K	(-0.3)	155.3	155.3	155.3	650. 163.1	682. 83COL/MCM
[CClH ₃]	CH ₃ Cl	RN 74-87-3 320 (Key) (br)		155 152	155 152	155 152	648 163	682 78COR/BEA 72BEA/HOL
[C ₂ H ₄]	C ₂ H ₄	RN 74-85-1 298 K 298 K 298 K Threshold Value	(0.0) (0.0) (0.0)	155.6 155.6 155.6 155.6	155.6 155.6 155.6 155.6	155.6 155.6 155.6 155.6	651. 162.6	680. 81DOI/MCM 83COL/MCM 81BOH/MAC 81TRA/MCL
[C ₂ N ₂]	NCCN	RN xxxxx (br)		155.	155.	155.	648. 162.	679. 84RAK/BOH
[H ₂ O ₂]	H ₂ O ₂	RN 7722-84-1 (br)					162.	678. 75LIN/ALB
[O ₂ S]	SO ₂	RN 7446-09-5 298 K	(-2) (-1.4)	153.6 154.2	153.6 154.2	153.6 154.2	645. 161.6	676. 81DOI/MCM 83COL/MCM
[C ₃ F ₆ O]	(CF ₃) ₂ CO	RN 684-16-2 298 K 298 K 298 (Key) 320 A	(-2.7) (-2.2)	152.9 153.4 153.4 156.3	152.9 153.4 153.4 156.3	152.9 153.4 153.4 156.3	642. 161.5	676. 81DOI/MCM 83COL/MCM 82URU/MCM 78COR/BEA
[CF ₂ O]	F ₂ CO	RN 353-50-4 298 K 298 K	(-3.2) (-2.7)	152.4 152.9	152.4 152.9	152.4 152.9	640. 160.5	671.5 81DOI/MCM 83COL/MCM
[C ₂ F ₄ O]	CF ₃ CFO	RN 354-34-7 298 K 298 K	(-3.6) (-3.2)	152.0 152.4	152.0 152.4	152.0 152.4	638. 160.2	670. 81DOI/MCM 83COL/MCM
[C ₃ H ₇]	i-C ₃ H ₇	RN 19252-53-0 Threshold Value					159.8	669. 79HOU/BEA

Table 1. Gas phase basicities and proton affinities--Continued

T K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
				kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[F ₂ O ₂ S]	F ₂ SO ₂	RN 2699-79-8		151.6	634.	159.0	665.	
	298	K	(-3.8)	151.6				81DOI/MCM
	298	K	(-3.6)	152.0				80DOI/MCM
[S]	S	RN 7704-34-9			152.3	637.	158.3	662.
		Threshold Value		152.3			158.3	662.
								81SMI/ADA
[HO ₂]	HO ₂	RN 3170-83-0				~158.	~661.	
		Threshold Value						75BRO
[Zn]	Zn	RN 7440-66-6					156	653
		(br)						78PO/RAD
[H ₄ Si]	SiH ₄	RN 7803-62-5		~147	~615	~155	~648	
		(br)	~147					73CHE/LAM
[AsF ₃]	AsF ₃	RN 7784-35-2		147.	615.	155.	648.	
		(br)	147					80DOI/MCM
[C ₂ H ₂]	C ₂ H ₂	RN 74-86-2		146.1	611.	153.3	641.	
		Threshold Value		146.1				84LIA/LIE
[CHO]	HCO	RN 17030-74-9				152.	636.	
		Threshold Value				152.		74WAR
[COS]	COS	RN 463-58-1		143.	598.	151.	632.	
		(br)	143					81SMI/ADA
[HI]	HI	RN 10034-85-2		143.	598.	150.	628.	
		(br)	143					78POL/MUN
[C ₃ H ₈]	C ₃ H ₈	RN 74-98-6		142.	594.	150.	628.	
		(Key)	142					76HIR/KEB-75HIR/KEB
[CFH ₃]	CH ₃ F	RN 593-53-3		142.	594.	150.	628.	
		(br)	142					72BEA/HOL
[CF ₂ H ₂]	CH ₂ F ₂	RN 75-10-5		139.	581.5	147.	615.	
		(br)	139.					74BLI/MCM
[CF ₃ H]	CHF ₃	RN 75-46-7		139.	581.5	147.	615.	
		(br)	139.					74BLI/MCM
[B ₂ H ₆]	B ₂ H ₆	RN 19287-45-7				~146	~611	
		(br)						73PIE/POR
[I]	I	RN 14362-44-8		140.4	587.	145.4	608.	
		Threshold Value		140.4				78POL/MUN
[F ₃ N]	NF ₃	RN 7783-54-2		136	569	144	604	
		(br)	136					80DOI/MCM

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	kJ/mol	Proton affinity kcal/mol	kJ/mol	Reference
[B ₄ H ₁₀]	B ₄ H ₁₀	RN 18283-93-7				~144	~602	
		(Key)						73PIE/POR
[C ₂ H ₆]	C ₂ H ₆	RN 74-84-0		135.8	568.	143.6	601.	
	298	O	(22.3)	133.0				81MAC/SCH
	298	Z	(1.4)	135.8				81BOH/MAC
	~200	(Key)		133.5A				76HIR/KEB
	~400	(Key)		141.3B				76HIR/KEB
[CO]	CO	RN 630-08-0		134.4	562.	141.9	593.	
	298	Z	(0.0)	134.4				81BOH/MAC
	298	Z	(0.0)	134.4				73HEM/RUN
		Threshold Value		136.7		144.4		76GUY/CHU
		Threshold Value		134.3		141.9		80DYK/JON
		Threshold Value		134.5		142.1		69MAT/WAR
	298	O	(20.9)	131.6				80BOH/MAC
	298	S	(11.0)	135.4				
[O ₃ S]	SO ₃	RN 7446-11-9		~130	~544	~138	~577	
		(br)	~131					77MUN/SMI
[N ₂ O]	N ₂ O	RN 10024-97-2		131.4	550.	136.5	571.	
	298	O	(18.0)	128.7				80BOH/MAC
	298	S	(8.1)	132.5				
	298	Z	(-3.0)	131.4				73HEM/RUN
[BrH]	HBr	RN 10035-10-6		131	548	136	569	
		(br)	131					78POL/MUN
		Threshold Value				132.9		79TIE/AND
[ClH]	HCl	RN 7647-01-0		127.	531.	134.8	564.	
		Threshold Value	127.					79TIE/AND
		(br)	129					78POL/MUN
		(Key)	>124					74FEH/FER
[Br]	Br	RN 10097-32-2		126.4	529.	132.0	552.	
		Threshold Value	126.4					78POL/MUN
[CH ₄]	CH ₄	RN 74-82-8		126.0	527.	132.0	552.	
	298	O	(11.8)	122.5				80BOH/MAC
	320	S	(2.1)	126.5				75STA/BEA
	340	S	(1.4)	125.8				75KAS/FRA
	340	S	(1.7)	126.1				LIA/AUS
	550	S	(1.3)	126.7				77MAU/FIE
	300	S	(1.5)	125.9				73HEM/RUN
	300	S	(1.9)	126.3				73BOH/HEM
		Threshold Value		120.5		126.5		71CHU/BER
[CO ₂]	CO ₂	RN 124-38-9		124.4	520.	130.9	548.	
		Threshold Value		124.4		130.9		74WAR
	798	S	(0.0)	126.7				76FEH/LIN
	320	S	(0.0)	124.4				75STA/BEA
	340	S	(0.0)	124.4				75KAS/FRA
	550	S	(0.0)	125.4				77MAU/FIE
	298	S	(0.0)	124.4				73HEM/RUN
	298	S	(0.0)	124.4				73BOH/HEM
	298	S	(0.0)	124.4				76MEI/MIT
	298	O	(+9.9)	120.8				80BOH/MAC

Table 1. Gas phase basicities and proton affinities--Continued

T	K	Refer- ence base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity		Proton affinity		Reference
					kcal/mol	kJ/mol	kcal/mol	kJ/mol	
[NO]	NO	RN 10102-43-9			~119	~498	~127	~531	
		(br)		~119					71ROC/SUT
[CF ₄]	CF ₄	RN 75-73-0			~119	~498	~126	~527	
		(br)		~119					71ROC/SUT
[Cl]	Cl	RN 22537-15-1			116.8	489.	123.0	515.	
		Threshold Value		116.8					78POL/MUN
[Xe]	Xe	RN 7440-63-3			113.4	474.	118.6	496.	
		298 O (+2.7)		113.4					80BOH/MAC-76FEH/LIN
		298 S (-7.2)		117.2					
[N ₂]	N ₂	RN 7727-37-9			111.0	464.	118.2	494.5	
		298 O (0.3)		111.0					80BOH/MAC-76FEH/LIN
		S (-9.6)		114.8					
		Threshold Value					118.1		79WIB/FIS
		Threshold Value					114.3		78FON/HUD
		Threshold Value					112.2		76WIL/LOS
[O]	O	RN 17778-80-2			110.7	463.	116.3	487.	
		298 O (0.0)		110.7					80BOH/MAC
		Threshold Value					116.3		76MCC
[Kr]	Kr	RN 7439-90-9			96.1	402.	101.6	425.	
		298 O (-10.3)		100.4					80BOH/MAC
		O' (+1.1)		96.1					
		BB (+1.7)		96.4					75PAY/SCH
		Threshold Value					100.3		79HUB/HER
[H ₂]	H ₂	RN 1333-74-0			94.6	396.	101.3	424.	
		298 O (-11.8)		98.9					80BOH/MAC-73FEN/HEM
		O' (-0.4)		94.6					
		O' (-0.3)		94.7					75FEH/LIN
		296 BB (0.0)		94.7					75PAY/SCH
		(Key)					101.		78PO/RAD
		Threshold Value		94.7			101.3		72COT/ROZ
[O ₂]	O ₂	RN 7782-44-7			95.0	397.	100.9	422.	
		298 O (-11.4)		99.3					80BOH/MAC-73FEN/HEM
		O' (0.0)		95.0					
		O' (0.0)		95.0					75FEH/LIN
		Threshold Value		94.6			100.5		81DYK/JON
		Threshold Value		95.0			100.9		77MCC
[FH]	HF	RN 7664-39-3							
		Threshold Value		87.3			95		79TIE/AND
		320 (Key)		109.4			117		75FOS/BEA
[F]	F	RN 14762-94-8			75.4	315.	81.0	339.	
		Threshold Value		75.4					
[Ar]	Ar	RN 7440-37-1					88.6	371.	
		Threshold Value					>61	>255	79HUB/HER
		BB					88.6	371.	82VIL/FUT

Table 1. Gas phase basicities and proton affinities--Continued

T K	Reference base	Relative gas basicity kcal/mol	Gas basicity kcal/mol	Selected gas basicity kcal/mol	Proton affinity kcal/mol	Reference
[Ne] Ne	RN 7440-01-9				48.1	201.
	Threshold Value				48.1	68CHU/RUS
	Threshold Value				48.0	79HUB/HER
[He] He	RN 7440-59-7				42.5	178.
	Threshold Value				42.5	79HUB/HER

Annotated References to Table 1

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 $\text{H}_2\text{O} < \text{C}_2\text{H}_5\text{Cl} < \text{H}_2\text{S}$
 $\text{C}_2\text{H}_4 < \text{CH}_3\text{Br} \sim \text{C}_2\text{H}_5\text{F} < \text{H}_2\text{O}$
 $\text{C}_2\text{H}_2 < \text{CH}_3\text{Cl} < \text{C}_2\text{H}_4$
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 $\text{CH}_3\text{COOCH}_3 < \text{C}_6\text{H}_5\text{CH}_2 < c\text{-C}_4\text{H}_9\text{O}$
 $\text{CH}_3\text{COC}_2\text{H}_5 < c\text{-C}_7\text{H}_7 < i\text{-C}_3\text{H}_7\text{COCH}_3$
 $\text{C}_2\text{H}_5 < \text{C}_2\text{H}_5\text{OC}_2\text{H}_5$
 $\text{NCCH}_2\text{CN} < c\text{-C}_3\text{H}_7 < \text{CF}_3\text{COOCH}_3$
 $\text{NCCH}_2\text{CN} < \text{C}_3\text{H}_7 < \text{CF}_3\text{COOCH}_3$
 $\text{CH}_3\text{CHO} < c\text{-C}_3\text{H}_7 < \text{C}_2\text{H}_5\text{OH}$
 $\text{CF}_3\text{COOCH}_3 < \text{C}_2\text{H}_5 < \text{CH}_3\text{OH}$
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 $\text{HCOOC}_2\text{H}_5 < \text{H}_2\text{B}_3\text{N}_3\text{H}_3 < \text{H}_3\text{B}_3\text{N}_3\text{H}_3$
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 $\text{NH}_3 < \text{C}_2\text{B}_4\text{H}_6 < \text{C}_6\text{H}_5\text{NH}_2$
 $\text{H}_2\text{O} < \text{C}_2\text{B}_3\text{H}_7 < \text{CF}_3\text{CH}_2\text{OH}$
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 $\text{CO} < \text{NF}_3 < \text{CH}_3\text{F}$
 $\text{SO}_2 < \text{PF}_3 < \text{C}_2\text{H}_4$
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 $\text{CH}_2\text{FCOCH}_3$ vs. HCOOC_2H_5 , $(\text{CH}_3)_2\text{O}$
 $\text{CH}_2\text{FCOCH}_2\text{F}$ vs. CH_3CHO , CH_3SH
 CF_3COCH_3 vs. HCOOH , $\text{CH}_2(\text{CN})_2$, H_2S
 $\text{CHF}_2\text{COCHF}_2$ vs. H_2S
 CF_3COOH vs. $\text{CHF}_2\text{COCHF}_2$
 CF_3COCF_3 vs. SO_2 , CF_2O
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 $(\text{CH}_2 = \text{CHCH}_2\text{CH}_2)_3\text{N} > \text{CH}_3\text{CH} = \text{CHN}(\text{CH}_3)_2 \sim (\text{CH}_3)_2\text{C} = \text{CHN}(\text{CH}_3)_2$
 $\sim c\text{-C}_3\text{H}_7\text{N}(\text{CH}_3) > 1,3\text{-}c\text{-C}_3\text{H}_7\text{N}_2(\text{CH}_3)$
 $(\text{C}_2\text{H}_5)_2\text{N} < \text{CH}_3\text{CH} = \text{C}(\text{C}_2\text{H}_5)\text{N}(\text{CH}_3)_2 = (\text{CH}_3)_2\text{NCH}_2\text{CH}_2\text{N}(\text{CH}_3)_2$
 $\text{CH}_3\text{CH} = \text{C}(\text{CH}_3)\text{N}(\text{CH}_3)_2 > (\text{CH}_3)_2\text{NCH}_2\text{CH}_2\text{N}(\text{CH}_3)_2$
 $< \text{CH}_3\text{CH} = \text{C}(\text{C}_2\text{H}_5)\text{N}(\text{CH}_3)_2$
 $(n\text{-C}_3\text{H}_7)_2\text{NH} < c\text{-C}_3\text{H}_7\text{N}_3(\text{CH}_3) < (\text{CH}_3)_2\text{C} = \text{CHN}(\text{CH}_3)_2 < (i\text{-C}_2\text{H}_5)_2\text{NH}$
 $(\text{CH}_3)_2\text{N}(\text{CH}_3)_3\text{NH}_2 < \text{CH}_3\text{CH} = \text{C}(\text{CH}_3)\text{N}(\text{CH}_3)_2$
 $< (\text{CH}_3)_2\text{NCH}_2\text{CH}_2\text{N}(\text{CH}_3)_2$
 $c\text{-C}_3\text{H}_7\text{NH} < \text{CH}_3 = \text{CHN}(\text{CH}_3)_2 < c\text{-C}_3\text{H}_7\text{N}_3(\text{CH}_3)$
 $1,3\text{-C}_6\text{H}_4(\text{CH}_3)(\text{NH}_2) < \text{CH}_3\text{CH} = \text{NH} < 2\text{-ClC}_5\text{H}_4\text{N}$
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 $3\text{-CH}_3\text{C}_6\text{H}_4\text{NH}_2 < \text{C}_2\text{H}_5\text{N} < 2\text{-ClC}_5\text{H}_4\text{N}$
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 $(\text{CH}_3)_2\text{CO} < \text{C}_2\text{H}_5\text{ONO} < \text{CH}_3\text{COOCH}_3$
 $(\text{C}_2\text{H}_5)_2\text{O} < i\text{-C}_3\text{H}_7\text{ONO} < \text{NH}_3$
 $\text{NH}_3 < t\text{-C}_4\text{H}_9\text{ONO} < \text{Pyrrole}$
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 $NH_3 < CH_2N_2 < CH_2N=NCH_3$
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 $H_2O < CF_3NO < HCN$
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The heat of formation of $C_3H_9^+$ was determined from the equilibrium constant for the process: $C_2H_5^+ + CH_4 \rightleftharpoons C_3H_9^+$ and that of $C_4H_{11}^+$, from the equilibrium constant for the process: $sec-C_3H_7^+ + CH_4 \rightleftharpoons C_4H_{11}^+$. Values reported in the paper have been modified slightly to take into account more recent values for heats of formation of relevant ions.
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 $C_4H_7O_3P$ vs. $CH_3COCH_2COCH_3$, $(CH_3)_6C_6$
 $C_5H_9O_3P$ vs. $HCON(CH_3)_2$
 $C_6H_9O_3P$ vs. 2-Cl-pyridine, $(t-C_4H_9)_2S$
 $C_3H_7O_3P$ vs. 2-Cl-pyridine, $(t-C_4H_9)_2S$
 $C_4H_9O_3P$ vs. pyridine
- trans-2-methoxy-cis,cis-4,6-dimethyl-1,3,2-dioxaphosphorinane vs. 3-CH₃-pyridine, $(C_2H_5)_2NH$, 2-CH₃-pyridine cis,cis-2-methoxy-4,6-dimethyl-1,3,2-dioxaphosphorinane vs. 4-CH₃pyridine, $c-C_5H_{10}NH$
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 $OP(OCH_3)_2$: Equilibrium with $C_6H_5NH_2$, $(C_6H_5)_2C=CH_2$
 $OP(OC_2H_5)_2$: Close to 3- $(CH_3CO)C_2H_4N$ and pyridazine ($C_4H_4N_2$)
 $SP(OCH_3)_2$: Equilibrium with 4- $CH_3OC_6H_4CHO$
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	ΔH kcal/mol	ΔS cal/K mol
THF:Oxepane	-2.7	(0)
$\text{C}_2\text{H}_5\text{SCH}_3$:Oxepane	+0.5	(1.4)
THF:1,3-Dioxane	+0.2	(0)
$(\text{C}_2\text{H}_5)_2\text{O}$:1,3-Dioxane	+1.4	(0)
$n\text{-}(\text{C}_3\text{H}_7)_2\text{O}$: $\text{CH}_3\text{OCH}_2\text{CH}_2\text{OCH}_3$	-2.2	-5.0
$(\text{C}_2\text{H}_5)\text{SCH}_3$: $\text{CH}_3\text{OCH}_2\text{CH}_2\text{OCH}_3$	-1.8	-3.6
$n\text{-Bu}_2\text{S}$: $\text{CH}_3\text{O}(\text{CH}_2)_3\text{OCH}_3$	-5.2	-5.4
2-FPyr: $\text{CH}_3\text{O}(\text{CH}_2)_3\text{OCH}_3$	-3.6	-5.5
3-FPyr: $\text{CH}_3\text{O}(\text{CH}_2)_3\text{OCH}_3$	+0.8	-7.0
2-FPyr: $\text{CH}_3(\text{OCH}_2\text{CH}_2)_2\text{OCH}_3$	-8.5	-13.2
3-FPyr: $\text{CH}_3(\text{OCH}_2\text{CH}_2)_2\text{OCH}_3$	-5.7	-15.7
1,2-Diazine:	-1.4	-11.7
1,2-Diaz: $\text{CH}_3(\text{OCH}_2\text{CH}_2)_3\text{OCH}_3$	-9.1	-17.7
1,2-Diaz:12-Crown-4	-3.3	-3.1
Pyr:12-Crown-4	+1.2	-2.6
1,2-Diaz:15-Crown-5	-3.6	(-3)
Pyr:15-Crown-5	+0.1	(-3)
1,2-Diaz:18-Crown-6	-2.7	(-3)
Pyr:18-Crown-6	+0.9	(-3)
$\text{HCOO}n\text{-C}_4\text{H}_9$: $\text{CH}_3\text{COCOCH}_3$	-1.7	(1.4)
$(\text{CH}_3)_2\text{CO}$: $\text{CH}_3\text{COCOCH}_3$	+1.9	2.6
$\text{CH}_3\text{COC}_2\text{H}_5$:	+3.7	(1.4)
$(n\text{-C}_3\text{H}_7)_2\text{S}$: $\text{CH}_3\text{COCH}_2\text{COCH}_3$	-1.2	-3.0
Pyrrrole: $\text{CH}_3\text{COCH}_2\text{COCH}_3$	+0.1	-4.1
Pyrrrole: $\text{CH}_3\text{COCH}_2\text{CH}_2\text{COCH}_3$	-3.9	-8.5
2-FPyr: $\text{CH}_3\text{COCH}_2\text{CH}_2\text{COCH}_3$	-2.6	-5.6
$(\text{C}_2\text{H}_5)_2\text{CO}$:Cyclohexanone	-2.5	(0)
$\text{C}_2\text{H}_5\text{SCH}_3$:Cyclohexanone	-1.0	(0)
$\text{C}_2\text{H}_5\text{SCH}_3$:1,2-Cyclohexanedione	-1.7	(1.4)
$(n\text{-C}_3\text{H}_7)_2\text{S}$:1,2-Cyclohexanedione	+3.8	(0)
Pyrrrole:1,3-Cyclohexanedione	-2.7	(1.4)
2-FPyr:1,3-Cyclohexanedione	-1.0	(1.4)
2-FPyr: $\text{CH}_3\text{CONHCH}_2\text{COOCH}_3$	-7.0	-13.4
2-FPyr: $\text{CH}_3\text{CONHCNCH}_2\text{COOCH}_3$	-12.3	-14.7
3-FPyr: $\text{CH}_3\text{CONHCNCH}_2\text{COOCH}_3$	-9.6	-15.7
H_2NNH_2 : $\Delta G = -4.7$ kcal/mol to pyrrole; -0.1 kcal/mol to cyclopropylmethylketone, $+2.1$ kcal/mol to n-propylether at 600 K. Isooxazole: $\Delta G = +2.7$ kcal/mol to $(\text{C}_2\text{H}_5)_2\text{O}$ at 600 K. Oxazole: $\Delta G = 0.3$ kcal/mol to pyrrole at 600 K. Furan: $\Delta G = +3.2$ kcal/mol to toluene; -0.6 kcal/mol to ethylformate at 600 K. 2,5-Dimethylfuran: $\Delta G = 2.4$ kcal/mol to pyrrole at 600 K. 2-Methylfuran: $\Delta G = -1.8$ kcal/mol to pyrrole at 600 K. Thiophene: $\Delta G = 1.5$ kcal/mol to 1,4-dioxane; -2.5 kcal/mol to $\text{CH}_3\text{COC}_2\text{H}_5$ at 600 K. 2-Methylthiophene: $\Delta G = 0.7$ kcal/mol to $c\text{-C}_4\text{H}_9\text{COCH}_3$ at 600 K. N-Methylimidazole: $\Delta G = 4.7$ kcal/mol to 3-methylpyridine; -5.5 kcal/mol to $(\text{C}_2\text{H}_5)_3\text{N}$ at 600 K. 4-Methylimidazole: $\Delta G = 0.4$ kcal/mol to 3-ethylpyridine at 600 K. Imidazole: $\Delta G = 2.6$ kcal/mol to $\text{iso-C}_3\text{H}_7\text{NH}_2$ at 600 K. 2,5-Dimethylpyrrole: $\Delta G = 1.0$ kcal/mol to $\text{iso-C}_3\text{H}_7\text{NH}_2$ at 600 K. Thiazole: $\Delta G = -0.8$ kcal/mol to 3-fluoropyridine at 600 K. Pyrazole: $\Delta G = 0.8$ kcal/mol to 2-fluoropyridine at 600 K. $\text{CH}_2=\text{CHOCH}_3$: $\Delta G = -0.7$ kcal/mol to pyrrole at 600 K. $\text{C}_2\text{H}_5\text{OCH}=\text{CH}_2$: $\Delta G = 1.0$ kcal/mol to pyrrole; 2.6 kcal/mol to 2-fluoropyridine at 600 K.		
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Also given, but not included here for lack of brackets:
 $(t\text{-C}_4\text{H}_9)_2\text{O} > t\text{-C}_4\text{H}_9\text{O}(i\text{-C}_3\text{H}_7)$ and
 $((\text{CH}_3)_2\text{HSi})_2\text{O} > ((\text{CH}_3)_3\text{SiCH}_2)_2\text{O}$ and
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High pressure mass spectrometer. Bracketing.

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ICR. Bracketing:
 $(t-C_6H_9)_2S < C_6H_4 < (CH_3)_2NCOOC_2H_5$
- 78POL/MUN C. W. Polley and B. Munson, "The Proton Affinities of the Halogen Acids", *Int. J. Mass Spectrom. Ion Phys.* **26**, 49 (1978).
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Bracketing: $Br < HCl < N_2O$
 $HCl < HBr < CO$
 $I < HI < HBr$
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ICR. Bracketing:
3-Clpyridine $< o-C_8H_8 < CF_3CH_2N(CH_3)_2$
2-Clpyridine $< p-C_8H_8 < C_2H_5NH_2$
- 77POL/WOL S. K. Pollack, J. F. Wolf, B. A. Levi, R. W. Taft, and W. J. Hehre, "Kinetic Detection of Common Intermediates in Gas Phase Ion-Molecule Reactions", *J. Am. Chem. Soc.* **99**, 1350 (1977).
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Heat of formation of H_3S^+ from appearance energy from $(H_2S)_2$; approximately corrected to 298 K by present authors.
- 83RAK/BOH A. B. Rakshit and D. K. Bohme, "The Proton Affinity of C_3 and Heat of Formation of C_3H^+ ", *Int. J. Mass Spectrom. Ion Phys.* **49**, 275 (1983).
Flowing afterglow. Bracketing:
 $CH_3OH < C_3 < CH_3CN$
- 84RAK/BOH A. B. Rakshit and D. K. Bohme, "Selected-Ion Flow Tube Methods Applied to the Bracketing of Proton Affinities. $PA(C_2N_2)$ and $PA(HC_3N)$ " *Int. J. Mass Spectrom. Ion Proc.* **57**, 211 (1984).
Flowing afterglow. Bracketing:
 $SO_2 < C_2N_2 < C_2H_4$
 $CH_3NO_2 < HC_3N < CH_3CN$
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Determination of appearance potentials of fragment ions from alcohols.
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ICR. Bracketing:
 $H_2O < C_2H_3F \sim H_2S$
 $C_2H_3I < CH_2CF_2 < CH_3OH$
 $C_2H_3F < CHFCHF < H_2O$
 $H_2O < C_2HF_3 < H_2S$
- 71ROC/SUT A. E. Roche, M. M. Sutton, D. K. Bohme, and H. I. Schiff, "Determination of Proton Affinity from the Kinetics of Proton Transfer Reactions. I. Relative Proton Affinities", *J. Chem. Phys.* **55**, 5480 (1971).
Flowing afterglow. Bracketing:
 $N_2 < CF_4$, $NO < CH_4$
 $Ar < H_2O_2$ (Not included in Table)
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Appearance potentials of $C_2H_5^+$ and $C_3H_7^+$ from alkyl halides.
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ICR. Bracketing:
 $CF_3CO_2C_2H_5 < CH_2S < CH_3CHO$
- 79SAL/KEB P. P. S. Saluja and P. Kebarle, "Heat of Formation of the 2-Norbornyl Cation in the Gas Phase", *J. Am. Chem. Soc.* **101**, 1084 (1979).
High pressure mass spectrometer. Proton transfer equilibria observed in norbornene: anisole ($\Delta G_{560} = 1.5$ kcal/mol) and anisole: cyclohexanone ($\Delta G_{560} = -1.9$ kcal/mol) systems.
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Determination of equilibrium constant of reaction: $C_6H_5CH_2^+ + (CH_3)_3CCl \rightleftharpoons (CH_3)_3C^+ + C_6H_5CH_2Cl$
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SIFT. Bracketing: $CO < COS < S$
- 78SMI/MUN D. E. Smith and B. Munson, "Proton Affinities of Some Sulfur-Oxygen Compounds", *J. Am. Chem. Soc.* **100**, 497 (1978).
Bracketing. $CH_3F < SO_2F_2 < SO_2 < HSO_3F < C_2H_4$
 $H_2O < H_2SO_4 \sim CF_3SO_3H < H_2S$
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Hydride transfer equilibrium constant determined for $t-C_4H_9^+ +$ Norbornane \rightarrow Norbornyl $^+$ + iso- C_4H_{10} . Heat of formation of $t-C_4H_9^+$ taken as 165.8 kcal/mol.
- 75SOL/HAR B. H. Solka and A. G. Harrison, "Bimolecular Reactions of Trapped Ions. Part XI. Rates and Equilibria in Proton Transfer Reactions of $CH_3SH_2^+$ ", *Int. J. Mass Spectrom. Ion Phys.* **17**, 379 (1975).
Equilibrium:
 $CH_3CHOH^+ + CH_3SH$, $\Delta G = -0.5$ kcal/mol

- $\text{CH}_3\text{SH}_2^+ + \text{C}_2\text{H}_5\text{CHO}, \Delta G = -2 \text{ kcal/mol}$
 $\text{CH}_3\text{SH}_2^+ + (\text{CH}_3)_2\text{O}, \Delta G = -4 \text{ kcal/mol}$
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 Bracketing: $\text{H}_2\text{O} < \text{B}_2\text{H}_6 < \text{H}_2\text{S}$
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 ICR. Data relative to TAFT scale; temperature assumed to be 320 K.
- 79STE/BEA A. E. Stevens and J. L. Beauchamp, "Gas Phase Organometallic Chemistry. Mechanism and Energetics of Methane Formation Resulting from Protonation of $(\text{CO})_2\text{MnCH}_3$ ", *J. Am. Chem. Soc.* **101**, 245 (1979).
 ICR. Bracketing:
 $\text{CH}_3\text{CH}=\text{CH}_2 < (\text{CO})_2\text{MnCH}_3 \sim \text{CH}_3\text{OH} < \text{CH}_3\text{CHO}$
- 81STE/BEA A. E. Stevens and J. L. Beauchamp, "Metal-Hydrogen Bond Energies in Protonated Transition Complexes", *J. Am. Chem. Soc.* **103**, 190 (1981).
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 ICR. Bracketing:
 $\text{C}_6\text{H}_6 < (\text{CH}_3)_2\text{Hg} < \text{C}_6\text{H}_5\text{CH}_3$
- 82STO/SPL J. A. Stone, D. E. Splinter, and S. Y. Kong, "A Comparison of the Relative Binding Energies of H^+ and NO^+ to Aromatic and Haloaromatic Bases by High Pressure Mass Spectrometry", *Can. J. Chem.* **60**, 910 (1982).
 High pressure mass spectrometer. Proton transfer equilibrium constants in mixtures of methyl formate with toluene, 3-fluorotoluene, 2-fluorotoluene, and 4-fluorotoluene.
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- TAFT
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 ICR. Data as reported corrected to 320 K. See comments under TAFT.
- 83TAF R. W. Taft, "Protonic Acidities and Basicities in the Gas Phase and in Solution: Substituent and Solvent Effects", *Prog. Org. Chem.* **14**, 248 (1983).
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 Flowing afterglow.
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 Flowing afterglow.
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- Appearance potential of sec-C₄H₉⁺ from halogenated butanes, and correction to 298 K.
- 81TRA/MCL J. C. Traeger and R. G. McLoughlin, "Absolute Heats of Formation for Gas Phase Cations", *J. Am. Chem. Soc.* **103**, 3647 (1981).
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- 82TRA/MCL J. C. Traeger, R. G. McLoughlin and A. J. C. Nicholson, "Heat of Formation of Acetyl Cation in the Gas Phase", *J. Am. Chem. Soc.* **104**, 5318 (1982).
- Appearance potentials of CH₃CO⁺ ions; correction to 298 K.
- 78TSA W. Tsang, "Thermal Stability of Primary Amines," *Int. J. Chem. Kinet.* **10**, 41 (1978).
- Heats of formation of benzyl, tert-butyl radicals.
- 79VAJ/HAR J. H. Vajda and A. G. Harrison, "Proton Affinities of Some Olefinic Carbonyl Compounds and Heats of Formation of C_nH_{2n-1}O⁺ Ions", *Int. J. Mass Spectrom. Ion Phys.* **30**, 293 (1979).
- High pressure mass spectrometer. Crotonaldehyde measured vs. acetone, methyl acetate, and ethyl acetate; methacrolein measured vs. acetone, p-dioxane, and n-butyl formate; acrolein measured vs. ethyl formate.
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- 77WAN/DES J.-S. Wang, A. J. DeStefano, and R. F. Porter, "Acidity of B₃H₈⁺ and Stability of the B₃H₈ Radical", *Inorg. Chem.* **17**, 1374 (1978).
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- 79WIB/FIS N. Wiberg, G. Fischer and H. Bachhuber, "Diazen und andere Distickstoffhydride: Bildungswarmen, Dissoziationsenergien, Aufttrittspotentiale, Protonenaffinitaten", *Z. Naturforsch.* **34b**, 1385 (1979).
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- 80WIG/BEA C. A. Wight and J. L. Beauchamp, "Acidity, Basicity, and Ion-Molecule Reactions of Isocyanic Acid in the Gas Phase by Ion Cyclotron Resonance Spectroscopy", *J. Phys. Chem.* **84**, 2503 (1980).
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- 76WIL/LOS C. Willis, F. P. Lossing and R. A. Back, "The Heat of Formation of N₂H₂ and the Proton Affinity of N₂", *Can. J. Chem.* **54**, 1 (1976).
- Heat of formation of N₂H⁺ as a fragment ion in N₂H₂.
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- High pressure mass spectrometer.
- 76YAM/KEB R. Yamdagni and P. Kebarle, "Gas Phase Basicities and Proton Affinities of Compounds between Water and Ammonia and Substituted Benzenes from a Continuous Ladder of Proton Transfer Equilibria Measurements" *J. Am. Chem. Soc.* **98**, 1320 (1976).
- High pressure mass spectrometer. Data assumed to have been superseded by data in LAU/KEB, when species studied have been duplicated. Other data corrected to LAU/KEB scale; free energy change values multiplied by 1.05.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species
(Prepared in collaboration with Mahnaz Motevalli-Aliabadi)

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[Ar]	Ar RN 7440-37-1	88.6	371.	0.	0.	DEF	277.	1159.
[AsC ₃ H ₉]	(CH ₃) ₃ As RN 593-88-4	213.4	893.	3.	13.(10)	{77PED/RYL}	155.	650.
[AsF ₃]	AsF ₃ RN 7784-35-2	155.	648.	-188.	-786.	[82/TN270]	23.	95.5
[AsH ₃]	AsH ₃ RN 7784-42-1	179.2	750.	16.	66.	[82/TN270]	202.	846.
[B ₂ H ₆]	B ₂ H ₆ RN 19287-45-7	~146	~611	9.	36.	[82/TN270]	228.	955.
[B ₃ H ₅ N ₃]	B-Boraziny radical RN xxxxx	193.6	810.					
[B ₃ H ₆ N ₃]	Borazine RN 6569-51-3	194.1	812.	-122.	-512.	[82/TN270]	49.	206.
[B ₄ C ₂ H ₆]	1,6-C ₂ B ₄ H ₆ RN 20693-67-8	207.	866.	-90.	-378.	Est	68.	286.
[B ₄ H ₈]	B ₄ H ₈ RN 12007-71-5	188	787					
[B ₄ H ₁₀]	B ₄ H ₁₀ RN 18283-93-7	~144	~602	16.	66.	[82/TN270]	237.	993.5
[B ₅ C ₂ H ₇]	2,4-C ₂ B ₅ H ₇ RN 20693-69-0	168.	703.	-85.	-356.	Est	113.	471.
[B ₅ H ₈]	B ₅ H ₈ RN 65930-58-7	184.	770.					
[B ₅ H ₉]	B ₅ H ₉ RN 19624-22-7	169.	707.	17.	73.	[82/TN270]	214.	896.
[Br]	Br RN 10097-32-2	132.0	552.	26.7	111.7	[82/TN270]	259.3	1085.
[BrCH ₃]	CH ₃ Br RN 74-83-9	165.7	693.	-9.	-37.(1)	{77PED/RYL}	191.	800.
[BrCN]	BrCN RN 506-68-3	178.3	746.	43.	181.(4)	{77PED/RYL}	231.	965.
[BrC ₅ H ₄ N]	4-Bromopyridine RN 1120-87-2	217.9	912.	38.	161.	Est	186.	779.
[BrC ₅ H ₄ N]	3-Bromopyridine RN 626-55-1	215.1	900.	30.	161.	Est	189.	791.
[BrC ₆ H ₄ N]	2-Bromopyridine RN 109-04-6	214.7	898.	38.	161.	Est	189.	793.
[BrC ₆ H ₅]	Bromobenzene RN 108-86-1	182.4	763.	25.	104.(3)	{77PED/RYL}	208.	871.
[BrC ₆ H ₆ N]	3-BrC ₆ H ₄ NH ₂ RN 591-19-5	208.1	871.	26.	108.	Est	183.	767.
[BrC ₇ H ₁₂ N]	3-Bromo-1-azabicyclo[2.2.2]- octane RN xxxxx	227.1**	950.**	1.	3.	Est	139.	583.
[BrH]	HBr RN 10035-10-6	136	569	-9.	-36.	[82/TN270]	221.	925.
[CCl ₃]	CH ₃ Cl RN 74-87-3	~168	~703	-19.5	-82.(1)	[79KUD/KUD]	178.	745.
[CClN]	ClCN RN 506-77-4	175.7	735.	32.	133.(1)	{77PED/RYL}	222.	928.
[CCl ₂]	CCl ₂ RN 1605-72-7	~200.0	~837.	45.	189.	[78AUS/LIA]	211.	882.
[CFH ₃]	CH ₃ F RN 593-53-3	150.	628.	-56.	-234.(8)	[78KUD/KUD]	160.	668.
[CF ₂]	CF ₂ RN 2154-59-8	171.9	719.	-52.	-217.	[77LIA/AUS]	142.	594.
[CF ₂ H ₂]	CH ₂ F ₂ RN 75-10-5	147.	615.	-108.	-453.(8)	[78KUD/KUD]	110.	462.
[CF ₂ O]	F ₂ CO RN 353-50-4	160.5	671.5	-153.	-640.(1)	{77PED/RYL}	52.	218.5
[CF ₃ H]	CHF ₃ RN 75-46-7	147.	615.	-166.	-695.(8)	[78KUD/KUD]	52.5	220.
[CF ₃ HO ₃ S]	CF ₃ SO ₃ H RN 1493-13-6	~169	~707	-282.	-1179.	Est	-85.	-356.
[CF ₃ NO]	CF ₃ NO RN 334-99-6	169	707	-126.	-529.	Est	70.	294.
[CF ₄]	CF ₄ RN 75-73-0	~126	~527	-223.	-933.(8)	[81BOM/BER]	17.	70.
[CHN]	HNC RN xxxxx	190.2	796.	51.	212.(8)	[82PAU/HEH]	226.	947.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[CHN]	HCN RN 74-90-8	171.4	717.	32.	135.	[82/TN270]	226.	947.
[CHNO]	HNCO RN 75-13-8	173.3	725.	-25.	-105. (13)	[75COM]	167.	700.
[CHO]	HCO RN 17030-74-9	152.	636.	9.	37. (5)	[82MCM/GOL]	223.	931.
[CH ₂ N ₂]	CH ₂ N ₂ RN 334-88-3	205.	858.	55.	230. (17)	[78VOG/WIL]	216.	902.
[CH ₂ O]	HCOH RN xxxxx	229.	958.	31.	131.	[82PAU/HEH2]	168.	703.
[CH ₂ O]	H ₂ CO RN 50-00-0	171.7	718.	-26.	-109. (1)	[77PED/RYL]	168.	703.
[CH ₂ O ₂]	HCOOH RN 64-118-6	178.8	748.	-90.5	-379. (1)	[78CHA/ZWO]	96.	403.
[CH ₂ S]	CH ₂ S RN 865-36-1	186.	778.	21.5	90. (8)	[82ROY/MCM]	201.	842.
[CH ₃ I]	CH ₃ I RN 74-88-4	~171	~715	3.5	15. (1)	[77PED/RYL]	198.	830.
[CH ₃ NO]	HCONH ₂ RN 75-12-7	198.4	830.	-44.	-186.	[69BEN/CRU]	123.	514.
[CH ₃ NO ₂]	CH ₃ ONO RN 624-91-9	192.5	805.	-16.	-67. (2)	[74BAT/CHR]	157.	658.
[CH ₃ NO ₂]	CH ₃ NO ₂ RN 75-52-5	179.2	750.	-18.	-75. (1)	[77PED/RYL]	168.5	705.
[CH ₄]	CH ₄ RN 74-82-8	132.0	552.	-18.	-75.	[74SCO]	216.	903.
[CH ₄ N]	CH ₂ NH ₂ RN 54088-53-8	199	833	36.	149. (8)	[82MCM/GOL]	202.	846.
[CH ₄ O]	CH ₃ OH RN 67-56-1	181.9	761.	-48.	-202. (1)	[77PED/RYL]	135.5	567.
[CH ₄ S]	CH ₃ SH RN 74-93-1	187.4	784.	-5.	-23. (1)	[77PED/RYL]	173.	723.
[CH ₅ N]	CH ₃ NH ₂ RN 74-89-5	214.1	896.	-5.	-23. (1)	[77PED/RYL]	146.	611.
[CH ₅ P]	CH ₃ PH ₂ RN 593-54-4	204.1	854.	-7.	-30.	Est	154.	646.
[CH ₆ N ₂]	CH ₃ NHNH ₂ RN 60-34-4	214.1**	896.**	23.	95. (1)	[77PED/RYL]	174.	729.
[CO]	CO RN 630-08-0	141.9	594.	-26.5	-111.	[82/TN270]	197.	824.
[COS]	COS RN 463-58-1	151.	632.	-34.	-142. (1)	[77PED/RYL]	181.	756.
[CO ₂]	CO ₂ RN 124-38-9	130.9	548.	-94.	-393.	[82/TN270]	141.	589.
[CS]	CS RN 2944-05-0	175	732	56.	234.	[82/TN270]	247.	1032.
[CS ₂]	CS ₂ RN 75-15-0	167.1	699.	28.	117. (1)	[77PED/RYL]	226.5	948.
[C ₂ BrH ₅]	C ₂ H ₅ Br RN 74-96-4	~171	~715	-15.	-62. (2)	[77PED/RYL]	180.	753.
[C ₂ ClH ₂ N]	ClCH ₂ CN RN 107-14-2	179.5	751.	20.5	86.	Est	207.	865.
[C ₂ ClH ₃ O ₂]	CH ₂ ClCOOH RN 79-11-8	182.4	763.	-104.	-435. (9)	[77PED/RYL]	79.	332.
[C ₂ ClH ₅]	C ₂ H ₅ Cl RN 75-00-3	169.	707.	-27.	-112. (1)	[77PED/RYL]	170.	711.
[C ₂ Cl ₃ HO ₂]	CCl ₃ COOH RN 76-03-9	183.5	768.	-106.	-444. (10)	Est	76.	318.
[C ₂ Cl ₃ H ₃ O]	Cl ₃ CCH ₂ OH RN 115-20-8	177.4	742.	-70.	-293.	Est	118.	495.
[C ₂ Cl ₃ N]	CCL ₃ CN RN 545-06-2	175.8	735.5	19.5	82.	Est	209.	876.5
[C ₂ D ₆ O]	(CD ₃) ₂ O RN 17222-37-6	190.6	797.	-44.	-184. (1)	[77PED/RYL]	131.	549.
[C ₂ FH ₃]	C ₂ H ₃ F RN 75-02-5	175.	732.	-33.	-139. (2)	[76WIL/LEB]	157.	659.
[C ₂ FH ₃ O ₂]	CH ₂ FCOOH RN 144-49-0	183.5	768.	-140.	-586.	Est	42.	176.
[C ₂ FH ₅]	C ₂ H ₅ F RN 353-36-6	165.	690.	-63.	-263. (2)	[75SCHE/ROD]	138.	577.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₂ FH ₆ N]	CH ₂ FCH ₂ NH ₂ RN 406-34-8	212.3	888.	-55.	-229.	Est	99.	413.
[C ₂ F ₂ H ₂]	CH ₂ CF ₂ RN 75-38-7	176	736	-82.	-345. (10)	[76WIL/LEB]	107.	449.
[C ₂ F ₂ H ₂]	(E)-CHFCHF RN 1630-78-0	166	694	-70.	-293.	[80STA/VOG]	130.	543.
[C ₂ F ₂ H ₄ O]	CF ₂ HCH ₂ OH RN 359-13-7	176.2	737.	-155.	-649.	Est	34.	144.
[C ₂ F ₂ H ₅ N]	CF ₂ HCH ₂ NH ₂ RN 430-67-1	207.5	868.	110.	462. (6)	Est	269.	1124.
[C ₂ F ₃ H]	C ₂ F ₃ H RN 359-11-5	~169	~707	-117.	-491. (8)	[77PED/RYL]	79.	332.
[C ₂ F ₃ HO]	CF ₃ CHO RN 75-90-1	165.1	691.	-189.	-790. (50)	[75HAR/THY]	12.	49.
[C ₂ F ₃ HO ₂]	CF ₃ COOH RN 76-05-1	169.0	707.	-246.	-1031. (1)	[77PED/RYL]	-50.	-208.
[C ₂ F ₃ H ₃ O]	CF ₃ CH ₂ OH RN 75-89-8	169.0	707.	-212.	-888. (5)	[77PED/RYL]	-15.5	-65.
[C ₂ F ₃ H ₄ N]	CF ₃ CH ₂ NH ₂ RN 753-90-2	202.5	847.	-167.5	-701.	Est	-4.	-18.
[C ₂ F ₃ N]	CF ₃ CN RN 353-85-5	166.1	695.	-118.	-495. (3)	[71JANAF]	81.	340.
[C ₂ F ₄ O]	CF ₃ CFO RN 354-34-7	160.2	670.	-249.	-1042.	Est	-43.5	-182.
[C ₂ H ₂]	C ₂ H ₂ RN 74-86-2	153.3	641.	54.	228. (1)	[82/TN270]	266.9	1117.
[C ₂ H ₂ O]	CH ₂ C=O RN 463-51-4	198.0	828.	-11.	-48. (8)	[71NUT/LAU]	157.	657.
[C ₂ H ₃]	C ₂ H ₃ radical RN 2669-89-8	~181	~757	70.5	295. (8)	[82MCM/GOL]	255.	1068.
[C ₂ H ₃ N]	CH ₃ CN RN 75-05-8	188.4	788.	18.	74. (1)	[83AN/MAN]	195.	816.
[C ₂ H ₃ NS]	CH ₃ SCN RN 556-64-9	195.9	820.	38.	160.	[82/TN270]	208.	870.
[C ₂ H ₃ NS]	CH ₃ NCS RN 556-61-6	195.9	820.	31.	131.	[82/TN270]	201.	841.
[C ₂ H ₄]	C ₂ H ₄ RN 74-85-1	162.6	680.	12.	52. (1)	[77PED/RYL]	215.6	902.
[C ₂ H ₄ N ₂]	NCCH ₂ NH ₂ RN xxxxx	197.4	826.	26.	108.	Est	194.	812.
[C ₂ H ₄ O]	c-C ₂ H ₄ O (Oxirane) RN 75-21-8	187.9	786.	-13.	-53. (1)	[77PED/RYL]	165.	691.
[C ₂ H ₄ O]	CH ₃ CHO RN 75-07-0	186.6	781.	-40.	-166. (1)	[77PED/RYL]	139.	583.
[C ₂ H ₄ O ₂]	CH ₃ COOH RN 64-19-7	190.2	796.	-103.	-432. (1)	[77PED/RYL]	72.	302.
[C ₂ H ₄ O ₂]	HCO ₂ CH ₃ RN 107-31-3	188.9	790.	-85.	-356. (1)	[77PED/RYL]	92.	384.
[C ₂ H ₄ S]	c-C ₂ H ₄ S (Thiirane) RN 420-12-2	194.6	814.	19.5	82. (1)	[77PED/RYL]	191.	798.
[C ₂ H ₅ I]	C ₂ H ₅ I RN 75-03-6	~176	~736	-2.	-9. (1)	[77PED/RYL]	187.5	785.
[C ₂ H ₅ N]	Aziridine (Azirane) RN 151-56-4	215.7	902.	30.	127. (1)	[77PED/RYL]	180.	755.
[C ₂ H ₅ N]	CH ₂ =CHNH ₂ RN 593-67-9	219.1	917.	7.	29.	[81ELL/DIX]	154.	643.
[C ₂ H ₅ N]	CH ₃ CH=NH RN 20729-41-3	213.9	895.	2.	8. (17)	[79ELL/EAD]	154.	643.
[C ₂ H ₅ NO]	CH ₃ CONH ₂ RN 60-35-5	206.2	863.	-57.	-238. (1)	[77PED/RYL]	103.	429.
[C ₂ H ₅ NO]	HCONHCH ₃ RN 123-39-7	205.8	861.	-45.	-187. (3)	Est	115.	481.
[C ₂ H ₅ NO ₂]	NH ₂ CH ₂ COOH (Glycine) RN 56-40-6	211.6	885.	-93.	-391. (5)	[77NGA/SAB]	61.	254.
[C ₂ H ₅ NO ₂]	C ₂ H ₅ ONO RN 109-95-5	197.3	825.5	-25.	-103.	[74BAT/CHR]	144.	601.5

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₂ H ₅ NO ₂]	C ₂ H ₅ NO ₂ RN 79-24-3	184.8	773.	-24.	-102. (1)	[77PED/RYL]	156.5	655.
[C ₂ H ₅ P]	c-C ₂ H ₄ PH (Phosphirane) RN 6569-82-0	191.4	801.	-16.	-69. (2)	Est	158.	660.
[C ₂ H ₆]	C ₂ H ₆ RN 74-84-0	143.6	601.	-20.	-84.	[74SCO]	202.	845.
[C ₂ H ₆ Hg]	CH ₃ HgCH ₃ RN 593-74-8	~186	~778	22.	94. (1)	[77PED/RYL]	202.	846.
[C ₂ H ₆ N ₂]	(E)-CH ₃ N=NCH ₃ RN 4143-41-3	206.9	866.	36.	149.	[82PAM/ROG]	194.	813.
[C ₂ H ₆ O]	(CH ₃) ₂ O RN 115-10-6	192.1	804.	-44.	-184. (1)	[77PED/RYL]	130.	542.
[C ₂ H ₆ O]	C ₂ H ₅ OH RN 64-17-5	188.3	788.	-56.	-235. (1)	[77PED/RYL]	121.	507.
[C ₂ H ₆ OS]	(CH ₃) ₂ SO RN 67-68-5	211.3	834.	-36.	-151. (1)	[77PED/RYL]	118.	495.
[C ₂ H ₆ S]	(CH ₃) ₂ S RN 75-18-3	200.6	839.	-9.	-38. (1)	[77PED/RYL]	156.	653.
[C ₂ H ₆ S]	C ₂ H ₅ SH RN 75-08-1	190.8	798.	-11.	-46. (1)	[77PED/RYL]	164.	686.
[C ₂ H ₆ S ₂]	CH ₃ SSCH ₃ RN 624-92-0	~196	~820	-6.	-24. (1)	[77PED/RYL]	164.	686.
[C ₂ H ₇ N]	(CH ₃) ₂ NH RN 124-40-3	220.6	923.	-4.5	-19. (1)	[77PED/RYL]	140.5	588.
[C ₂ H ₇ N]	C ₂ H ₅ NH ₂ RN 75-04-7	217.0	908.	-11.	-48. (1)	[77PED/RYL]	137.	574.
[C ₂ H ₇ NO]	NH ₂ (CH ₂) ₂ OH RN 141-43-5	221.3	926.	-48.	-202.	[77REI/PRA]	96.	402.
[C ₂ H ₇ O ₃ P]	(CH ₃ O) ₂ PHO RN 868-85-9	207.2	867.					
[C ₂ H ₇ P]	(CH ₃) ₂ PH RN 676-59-5	216.3	905.	-16.	-66.	Est	134.	559.
[C ₂ H ₈ N ₂]	1,2-Diaminoethane RN 107-15-3	225.9	945.	-4.	-18. (2)	[77PED/RYL]	135.	567.
[C ₂ H ₈ N ₂]	(CH ₃) ₂ NNH ₂ RN 57-14-7	219.9	920.	20.	84. (2)	[77PED/RYL]	166.	694.
[C ₂ N ₂]	NCCN RN xxxxx	162.	678.					
[C ₃]	C ₃ RN 12075-35-3	~185	~774	200 (4)	837 (17)	[83RAK/BOH]	~381.	~1593.
[C ₃ ClH ₄ N]	Cl(CH ₂) ₂ CN RN 542-76-7	187.5	784.5	10.	41.	Est	188.	786.5
[C ₃ FH ₅ O]	CH ₃ COCH ₂ F RN 430-51-3	192.0	803.	-91.5	-383.	Est	82.	344.
[C ₃ FH ₈ N]	FCH ₂ CH ₂ CH ₂ NH ₂ RN 462-41-9	217.8	911.	-61.	-254.	Est	87.	365.
[C ₃ F ₂ H ₄ O]	CFH ₂ COCFH ₂ RN 453-14-5	187	782	-126.	-529.	Est	52.	219.
[C ₃ F ₃ H ₃ O]	CH ₃ COCF ₃ RN 421-50-1	174.2	729.	-194.	-812.	Est	-3.	-11.
[C ₃ F ₃ H ₃ O ₂]	HCOOCH ₂ CF ₃ RN 32042-38-9	179.4	751.	-256.	-1073.	Est	-70.	-293.5
[C ₃ F ₃ H ₃ O ₂]	CF ₃ COCH ₃ RN 431-47-0	178.8	748.	-242.	-1013.	Est	-55.	-231.
[C ₃ F ₃ H ₆ N]	CF ₃ CH ₂ CH ₂ NH ₂ RN 460-39-9	210.6	881.	-175.	-731.	Est	-20.	-82.
[C ₃ F ₃ H ₆ N]	CF ₃ CH ₂ NHCH ₃ RN 2730-67-8	209.8	878.	-167.	-699.	Est	-11.	-47.
[C ₃ F ₃ H ₆ N]	CF ₃ N(CH ₃) ₂ RN 677-41-8	193.8	811.	-187.	-784. (15)	Est	-15.	-65.
[C ₃ F ₄ H ₂ O]	CF ₂ HCOCF ₂ H RN 360-52-1	170.	711.	-228.	-953.	Est	-32.	-134.
[C ₃ F ₅ N]	C ₂ F ₅ CN RN 422-04-8	167.1	699.	-219.	-917. (29)	[73THY/HAR]	-20.5	-86.
[C ₃ F ₆ H ₂ O]	(CF ₃) ₂ CHOH RN 920-66-1	165.0	690.	-381	-1595.	Est	-180.	-754.
[C ₃ F ₆ O]	(CF ₃) ₂ CO RN 684-16-2	161.5	676.	-334.	-1397.	[72GOR/T600]	-130.	-543.
[C ₃ GeH ₈]	(CH ₃) ₂ Ge=CH ₂ RN 82064-99-1	204.9	857.	19.	81.	[82PIE/HEH]	180.	754.
[C ₃ HN]	HCOCN RN xxxxx	184.	770.					

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H^\circ (M)$		Reference	$\Delta_f H^\circ (MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₃ H ₂ N ₂]	CH ₂ (CN) ₂ RN 109-77-3	175.6	735.	63.5	266.	(2) [77PED/RYL]	254.	1061.
[C ₃ H ₃]	c-C ₃ H ₃ radical RN xxxxx	175.8	735.	99.	414.	(17) [82MCM/GOL]	289.	1210.
[C ₃ H ₃ N]	CH ₂ =CHCN RN 107-13-1	189.7	794.	44.	184.	[82CHU/NGU]	220.	920.
[C ₃ H ₃ NO]	Oxazole RN 288-42-6	208.4	872.	-4.	-16.	(1) [78MCC/HAM]	153.	642.
[C ₃ H ₃ NO]	Isooxazole RN 288-14-2	202.3	846.	19.	79.	(1) [78MCC/HAM]	182.	763.
[C ₃ H ₃ NS]	Thiazole RN 288-47-1	213.2	892.	36.5	153.	(10) Est	189.	791.
[C ₃ H ₃ N ₃]	1,3,5-Triazine RN 290-87-9	201.1	841.	54.	226.	(1) [82BYS]	219.	915.
[C ₃ H ₄]	Cyclopropene RN 2781-85-3	198	828	66.	277.	(3) [77PED/RYL]	234.	979.
[C ₃ H ₄]	H ₂ C=C=CH ₂ RN 463-49-0	186.3	779.	46.	191.	(1) [77PED/RYL]	225.	942.
[C ₃ H ₄]	CH ₃ CCH RN 74-99-7	182	761	45.	187.	(2) [77PED/RYL]	228.	956.
[C ₃ H ₄ N ₂]	Imidazole RN 288-32-4	219.8	920.	35.	145.	(2) [80SAB]	180.5	755.
[C ₃ H ₄ N ₂]	Pyrazole RN 288-13-1	209.8	878.	44.	185.	(2) [80SAB]	200.	837.
[C ₃ H ₄ O]	CH ₃ CH=CO RN 6004-44-0	199.4	834.	-25.	-105.	[80DEM/WUL]	141.	591.
[C ₃ H ₄ O]	CH ₂ =CHCHO RN 107-02-8	193.9	811.	-18.	-77.	[79VAJ/HAR]	153.	642.
[C ₃ H ₅]	c-C ₃ H ₅ radical RN xxxxx	188.	787.	62.	261.	[80DEF/MCI]	240.	1004.
[C ₃ H ₅]	CH ₂ =CH-CH ₂ radical RN xxxxx	175.8	735.	43.	179.	[81TSA]	229.	958.5
[C ₃ H ₅ N]	1-Azabicyclo[1.1.0]butane RN 19540-05-7	212**	887**	75.	314.	Est	229.	957.
[C ₃ H ₅ N]	HCCCH ₂ NH ₂ RN 2450-71-7	210.8	882.	53.	222.	Est	208.	870.
[C ₃ H ₅ N]	C ₂ H ₅ CN RN 107-12-0	192.6	806.	12.	51.	[82CHU/NGU]	185.	775.
[C ₃ H ₅ O ₃ P]	2,6,7-Trioxa-1-phospha- bicyclo[2.2.1.]heptane RN 279-53-8	194.0	812.	-146.5	-613.	Est	25.	105.
[C ₃ H ₆]	c-C ₃ H ₆ RN 75-19-4	179.8	752.	13.	53.	(1) [77PED/RYL]	198.5	831.
[C ₃ H ₆]	CH ₃ CH=CH ₂ RN 115-07-1	179.5	751.	5.	20.	(1) [77PED/RYL]	191.	799.
[C ₃ H ₆ N ₂]	H ₂ N(CH ₂) ₂ CN RN 151-18-8	207.0	866.	22.	91	Est	180.	755.
[C ₃ H ₆ N ₂]	CH ₃ NHCH ₂ CN RN 5616-32-0	206.0	862.	25.5	107.	Est	185.	775.
[C ₃ H ₆ O]	CH ₂ =CHOCH ₃ RN 107-25-5	207.4	868.	-24.	-100.	(7) Est	134.	562.
[C ₃ H ₆ O]	c-C ₃ H ₆ O (Oxetane) RN 503-30-0	196.9	824.	-19.	-81.	(1) [77PED/RYL]	149.	625.
[C ₃ H ₆ O]	(CH ₃) ₂ CO RN 67-64-1	196.7	823.	-52.	-217.	(1) [76CHA/ZWO]	117.	490.
[C ₃ H ₆ O]	2-Methyloxirane RN 75-56-9	194.7	815.	-23.	-95.	(1) [77PED/RYL]	148.	620.
[C ₃ H ₆ O]	C ₂ H ₅ CHO RN 123-38-6	189.6	793.	-45.	-187.	(2) [77PED/RYL]	131.	550.
[C ₃ H ₆ O ₂]	CH ₃ COOCH ₃ RN 79-20-9	197.8	828.	-99.	-414.	(1) [*80SVO/UCH]	69.	288.
[C ₃ H ₆ O ₂]	HCO ₂ C ₂ H ₅ RN 109-94-4	193.1	808.	-92.	-387.	Est	80.	335.
[C ₃ H ₆ O ₂]	C ₂ H ₅ COOH RN 79-09-4	191.8	802.	-107.	-448.	(2) [77PED/RYL]	67.	279.5
[C ₃ H ₆ O ₃]	(CH ₃ O) ₂ CO RN 616-38-6	200.2	838.	-131.5	-550.	Est	34.	142.
[C ₃ H ₆ S]	Thietane RN 287-27-4	201.3**	842.**	14.5	61.	(1) [77PED/RYL]	179.	749.
[C ₃ H ₆ S]	2-Methylthiirane RN 1072-43-1	200.6**	839.**	11.	46.	(2) [77PED/RYL]	176.	737.
[C ₃ H ₇]	i-C ₃ H ₇ RN 19252-53-0	159.8	669.	21.5	91.	[81TSA]	227.4	951.5

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₃ H ₇ N]	CH ₂ =C(CH ₃)NH ₂ RN 4427-28-5	226.3	947.	8.	32.	Est	147.	615.
[C ₃ H ₇ N]	Azetidine RN 503-29-7	223.5	935.	24.	99. (4)	Est	166.	694.
[C ₃ H ₇ N]	N-Methylaziridine RN 1072-44-2	221.6	927.	30.	127.	Est	174.	730.
[C ₃ H ₇ N]	(CH ₃) ₂ C=NH RN 38697-07-3	221	925	2.	10.	[70BEN/O`N]	147.	615.
[C ₃ H ₇ N]	2-Methylaziridine RN 75-55-8	219.2**	917.**	22.	91. (6)	Est	168.	704.
[C ₃ H ₇ N]	H ₂ C=CHCH ₂ NH ₂ RN 107-11-9	215.8	903.	14.	57.	Est	163.5	684.
[C ₃ H ₇ N]	c-C ₃ H ₅ NH ₂ RN 765-30-0	215.0**	899.**	18.	77. (1)	[77PED/RYL]	169.	708.
[C ₃ H ₇ NO]	(CH ₃) ₂ NCHO RN 68-12-2	211.4	884.	-46.	-192. (2)	[77PED/RYL]	108.	453.5
[C ₃ H ₇ NO ₂]	Sarcosine RN xxxxx	218.7	915.	-88.	-367. (1)	[78SAB/LAF]	59.	248.
[C ₃ H ₇ NO ₂]	L-Alanine RN 56-41-7	214.8	899.	-99.	-415. (4)	[77NGA/SAB]	52.	216.
[C ₃ H ₇ NO ₂]	i-C ₃ H ₇ ONO RN 541-42-4	201.9	845.	-32.	-133. (4)	[74BAT/CHR]	132.	552.
[C ₃ H ₇ NO ₂ S]	L-Cysteine RN 3374-22-9	214.3	897.					
[C ₃ H ₇ NO ₃]	L-Serine RN 302-84-1	216.8	907.	-134.	-561.	Est	15.	62.
[C ₃ H ₇ O ₃ P]	2-Methoxy-1,3,2-dioxaphospholane RN 3741-36-4	212.7	890.	-164.	-688.	Est	-11.	-48.
[C ₃ H ₈]	C ₃ H ₈ RN 74-98-6	150.	628.	-25.	-105.	[74SCO]	191.	797.
[C ₃ H ₈ O]	CH ₃ OC ₂ H ₅ RN 540-67-0	196.4	822.	-52.	-216. (1)	[77PED/RYL]	118.	492.
[C ₃ H ₈ O]	i-C ₃ H ₇ OH RN 67-63-0	191.2	800.	-65.	-273. (1)	[77PED/RYL]	109.	457.
[C ₃ H ₈ O]	n-C ₃ H ₇ OH RN 71-23-8	190.8	798.	-61.	-255. (1)	[77PED/RYL]	114.	476.
[C ₃ H ₈ Pb]	(CH ₃) ₂ Pb=CH ₂ RN 82065-01-8	223.9	937.	59.	247.	[82PIE/HEH]	200.	840.
[C ₃ H ₈ S]	CH ₃ SC ₂ H ₅ RN 624-89-5	203.5	851.	-14.	-60. (1)	[77PED/RYL]	148.	619.
[C ₃ H ₈ S]	i-C ₃ H ₇ SH RN 75-33-2	194.1	812.	-18.	-76. (1)	[77PED/RYL]	153.	642.
[C ₃ H ₈ S]	n-C ₃ H ₇ SH RN 107-03-9	191.6	802.	-16.	-68. (1)	[77PED/RYL]	158.	660.
[C ₃ H ₈ S1]	(CH ₃) ₂ S1=CH ₂ RN 4112-23-6	226.4	947.	-1.	-5.	[82PIE/HEH]	138.	579.
[C ₃ H ₈ Sn]	(CH ₃) ₂ Sn=CH ₂ RN 82065-00-7	215.8	903.	31.5	132.	[82PIE/HEH]	181.	759.
[C ₃ H ₉ N]	(CH ₃) ₃ N RN 75-50-3	225.1	942.	-6.	-24. (1)	[77PED/RYL]	135.	564.
[C ₃ H ₉ N]	(CH ₃)(C ₂ H ₅)NH RN 624-78-2	222.8	932.	-11.	-46. (2)	Est	132.	552.
[C ₃ H ₉ N]	i-C ₃ H ₇ NH ₂ RN 75-31-0	218.6	915.	-20.	-84. (1)	[*79MAJ/SVO]	127.	531.
[C ₃ H ₉ N]	n-C ₃ H ₇ NH ₂ RN 107-10-8	217.9	912.	-17.	-70. (1)	[*79MAJ/SVO]	131.	548.
[C ₃ H ₉ NO]	NH ₂ (CH ₂) ₃ OH RN 156-87-6	228.6	956.5	-52.	-218.	Est	85.	356.
[C ₃ H ₉ NO]	CH ₃ OCH ₂ CH ₂ NH ₂ RN 109-85-3	223.3	934.	-44.	-184. (3)	Est	98.	412.
[C ₃ H ₉ O ₃ P]	P(OCH ₃) ₃ RN 121-45-9	220.6	923.	-167.	-697. (20)	Est	-21.	-92.
[C ₃ H ₉ O ₃ PS]	SP(OCH ₃) ₃ RN 29952-66-7	214.5	897.	-183.	-767.	Est	-32.	-134.
[C ₃ H ₉ O ₄ P]	OP(OCH ₃) ₃ RN 512-56-1	212.0	887.	-264.5	-1107.	Est	-111.	-464.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₃ H ₉ P]	(CH ₃) ₃ P RN 594-09-2	227.1	950.	-24.	-101.(5)	[77PED/RYL]	114.	479.
[C ₃ H ₁₀ N ₂]	1,3-Diaminopropane RN 109-76-2	234.1	979.	-8.	-32.	Est	124.	518.
[C ₃ H ₁₀ OSi]	(CH ₃) ₃ SiOH See References to Table 1: 75PIT/BUR							
[C ₄ F ₂ H ₇ MO]	CF ₂ HCON(CH ₃) ₂ RN 667-50-5	207.2	867.					
[C ₄ F ₃ H ₅ O ₂]	CF ₃ CO ₂ C ₂ H ₅ RN 383-63-1	184.6	772.	-249.	-1042.	Est	-68.	-284.
[C ₄ F ₃ H ₇ O]	C ₂ H ₅ OCH ₂ CF ₃ RN 461-24-5	186.4	780.	-216.	-904.	Est	-37.	-154.
[C ₄ F ₃ H ₈ N]	CF ₃ CH ₂ N(CH ₃) ₂ RN 819-06-7	215.0	900.	-167.	-700.	Est	-17.	-69.
[C ₄ F ₃ H ₈ N]	CF ₃ (CH ₂) ₃ NH ₂ RN 819-46-5	214.3	897.	-180.	-755.	Est	-29.	-121.5
[C ₄ F ₄ H ₄ O ₂]	CF ₃ COOCH ₂ CH ₂ F RN 1683-88-1	178.6	747.	-292.5	-1224.	Est	-105.	-441.
[C ₄ F ₆ H ₄ O]	(CF ₃) ₂ C(CH ₃)OH RN 1515-14-6	167.0	699.	-391.	-1636.	Est	-192.	-805.
[C ₄ F ₇ N]	C ₃ F ₇ CN RN 375-00-8	167.4	700.	-308.	-1290.(40)	[73THY/HAR]	-110.	-460.
[C ₄ F ₉ HO]	(CF ₃) ₃ COH RN 2378-02-1	163.1	682.	-549.	-2297.	Est	-346.	-1449.
[C ₄ F ₉ H ₂ N]	(CF ₃) ₃ CNH ₂ RN 2809-92-9	191.5**	801.**	-503.	-2104.	Est	-329.	-1375.
[C ₄ H ₄ N ₂]	Pyridazine (1,2-Diazine) RN 289-80-5	215.6	902.	66.	278.(1)	[77PED/RYL]	216.5	906.
[C ₄ H ₄ N ₂]	Pyrimidine (1,3-Diazine) RN 289-95-2	210.5	881.	46.	193.(2)	[77NAB/SAB]	201.	842.
[C ₄ H ₄ N ₂]	Pyrazine (1,4-Diazine) RN 290-37-9	209.0	874.	47.	196.(2)	[81STE/BAR]	203.5	852.
[C ₄ H ₄ N ₂ O]	2(1H)-Pyrimidinone RN 557-01-7	~208	~870	-11.	-47.	Est	146.	613.
[C ₄ H ₄ N ₂ O ₂]	Uracil RN 66-22-8	~208	~870	-72.	-303.(2)	[77NAB/SAB]	85.	357.
[C ₄ H ₄ N ₂ S ₂]	Dithiouracil RN 2001-93-6	~217	~907	51.	214.	Est	200.	836.
[C ₄ H ₄ O]	Furan RN 110-00-9	192.2	804.	-8.	-35.(1)	[77PED/RYL]	165.	691.
[C ₄ H ₄ S]	c-C ₄ H ₄ S (Thiophene) RN 110-02-1	196.5	822.	27.	115.(1)	[81KUD/KUD3]	197.	823.
[C ₄ H ₅ N]	Pyrrrole RN 109-97-7	207.6	868.	26.	108.	[80WIL/BAE]	184.	769.
[C ₄ H ₅ N]	c-C ₃ H ₅ CN RN 5500-21-0	195.4	817.5	44.	183.(1)	[82FUC/HAL]	214.	895.5
[C ₄ H ₅ NO ₂]	NCCOOC ₂ H ₅ RN 623-49-4	179.5	751.	-52.	-217.	Est	134.	562.
[C ₄ H ₅ N ₃ O]	Cytosine RN 71-30-7	223.8	936.	-14.	-59.(10)	[80SAB2]	128.	535.
[C ₄ H ₆]	1-Methylcyclopropene RN 3100-04-7	206	862	58.	244.(1)	[77PED/RYL]	218.	912.
[C ₄ H ₆]	(E)-CH ₂ =CHCH=CH ₂ RN 106-99-0	193**	807.5*	26.	110.(1)	[77PED/RYL]	199.	832.5
[C ₄ H ₆]	Cyclobutene RN 822-35-5	191**	799**	37.5	157.(2)	[77PED/RYL]	212.	888.
	Note: Heat of formation of cyclobutyl ion =						225 kcal/mol,	941 kJ/mol
[C ₄ H ₆]	CH ₃ COCH ₃ RN 503-17-3	187**	782**	35.	145.(1)	[77PED/RYL]	213.	893.
[C ₄ H ₆ N ₂]	1-Methylimidazole RN 616-47-7	228.9	958.	43.5	182.	Est	180.	754.
[C ₄ H ₆ N ₂]	4-Methylimidazole RN 822-36-6	224.4	939.	26.	110.	Est	167.	701.
[C ₄ H ₆ O]	CH ₂ =CHCOCH ₃ RN 78-94-4	200.2	838.	-29.	-125.	[79VAJ/HAR]	136.	567.
[C ₄ H ₆ O]	CH ₃ CH=CHCHO RN 4170-30-3	199.7	835.5	-22.	-91.(E)	[79VAJ/HAR]	144.	603.5
[C ₄ H ₆ O]	CH ₂ =C(CH ₃)CHO RN 78-85-3	195.2	817.	-25.	-106.	[79VAJ/HAR]	145.	607.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₄ H ₆ O ₂]	CH ₃ COCOCH ₃ RN 431-03-8	194.8	815.	-78.	-327.	[77PED/RYL]	93.	388.
[C ₄ H ₇ N]	i-C ₃ H ₇ CN RN 78-82-0	194.3	813.	5.	23.	[82CHU/NGU]	177.	740.
[C ₄ H ₇ N]	n-C ₃ H ₇ CN RN 109-74-0	193.7	810.	7.	31.	[82CHU/NGU]	179.	751.
[C ₄ H ₇ NO ₄]	L-Aspartic Acid RN 617-45-8	216.7	907.	-193.	-808.	Est	-44.	-104.5
[C ₄ H ₇ O ₂]	1,4-Dioxyl radical RN 4598-47-4	193.8	811.					
[C ₄ H ₇ O ₃ P]	2,6,7-Trioxa-1-phosphabicyclo-[2.2.2]octane RN 280-45-5	207.1	866.5	-128.5	-538.	Est	30.	125.5
[C ₄ H ₇ O ₃ P]	1-Methyl-2,6,7-trioxa-1-phosphabicyclo[2.2.1]heptane RN 61580-09-4	198.1	829.	-154.	-646.	Est	13.	55.
[C ₄ H ₈]	(CH ₃) ₂ C=CH ₂ RN 115-11-7	195.9	820.	-4.	-17. (1)	[77PED/RYL]	166.	693.
[C ₄ H ₈]	E-CH ₃ CH=CHCH ₃ RN 624-64-6	179.4	751.	-3.	-12. (1)	[77PED/RYL]	183.	767.
[C ₄ H ₈ N ₂]	NCCH ₂ N(CH ₃) ₂ RN 926-64-7	211.1	883.	34.	141.	Est	188.	788.
[C ₄ H ₈ N ₂ O ₃]	L-Asparagine RN 3130-87-8	219.8	920.	-141.	-591.	Est	5.	19.
[C ₄ H ₈ O]	C ₂ H ₅ OCH=CH ₂ RN 109-92-2	208.2	871.	-34.	-141. (1)	[77PED/RYL]	124.	518.
[C ₄ H ₈ O]	CH ₃ COC ₂ H ₅ RN 78-93-3	199.8	836.	-57.	-239. (1)	[76CHA/ZWO]	109.	455.
[C ₄ H ₈ O]	c-C ₄ H ₈ O (Tetrahydrofuran) RN 109-99-9	198.8	831.	-44.	-184. (1)	[77PED/RYL]	123.	514.
[C ₄ H ₈ O]	i-C ₃ H ₇ CHO RN 78-84-2	192.6	806.	-52.	-216. (1)	[77PED/RYL]	121.	508.
[C ₄ H ₈ O]	n-C ₃ H ₇ CHO RN 123-72-8	191.5	801.	-50.	-208. (2)	[77PED/RYL]	124.	521.
[C ₄ H ₈ O ₂]	C ₂ H ₅ COOCH ₃ RN 554-12-1	200.2	838.	-103.	-432.	[80HOL/LOS]	62.	260.
[C ₄ H ₈ O ₂]	1,3-Dioxane RN 505-22-6	198.8	832.	-81.	-338. (1)	[82BYS/MAN]	86.	360.
[C ₄ H ₈ O ₂]	1,4-Dioxane RN 123-91-1	193.8	811.	-75.5	-316. (1)	[82BYS/MAN]	96.	403.
[C ₄ H ₈ O ₂]	CH ₃ COOC ₂ H ₅ RN 141-78-6	200.7	840.	-106.	-443. (1)	[*80SVO/UCH]	59.	247.
[C ₄ H ₈ O ₂]	HCOOCH(CH ₃) ₂ RN 625-55-8	196.0	820.	-97.	-405.	[70BEN/O'N]	73.	305.
[C ₄ H ₈ O ₂]	HCO ₂ (n-C ₃ H ₇) RN 110-74-7	194.2	812.5	-110.	-462.	[*80SVO/UCH]	61.	255.5
[C ₄ H ₈ O ₂ S]	C ₂ H ₅ S(OCH ₃)CO RN 38103-96-7	201.0	841.	-100.	-420.	Est	64.	269.
[C ₄ H ₈ O ₃]	C ₂ H ₅ COOCH ₃ RN 623-53-0	202.7	848.	-141.	-592.	Est	21.5	90.
[C ₄ H ₉ N]	(CH ₃) ₂ NCH=CH ₂ RN 5763-87-1	227.8	953.	15.	62.	Est	153.	639.
[C ₄ H ₉ N]	Pyrrolidine RN 123-75-1	225.2	942.	-1.	-3. (1)	[77PED/RYL]	140.	585.
[C ₄ H ₉ N]	CH ₃ CH=NC ₂ H ₅ RN 1190-79-0	222.7	932.	4.	18. (1)	Est	147.	616.
[C ₄ H ₉ N]	CH ₂ =C(CH ₃)CH ₂ NH ₂ RN 2878-14-0	218.2**	913.**	5.	21.	Est	152.5	638.
[C ₄ H ₉ NO]	Morpholine RN 110-91-8	219.4	918.	12.	51.	Est	158.	663.
[C ₄ H ₉ NO]	Dimethylacetamide RN 127-19-5	216.8	907.	-55.6	-233.	[*78BEA/LEE]	93.	390.
[C ₄ H ₉ NO]	n-C ₃ H ₇ NHCHO RN 6281-94-3	210.0**	879.	-61.	-256.	Est	94.5	395.
[C ₄ H ₉ NO ₂]	t-C ₄ H ₉ ONO RN 540-80-7	205.7	861.	-41.	-172. (4)	[74BAT/CHR]	119.	497.
[C ₄ H ₉ NO ₃]	L-Threonine RN xxxxx	218.6	915.	-141.	-592.	Est	6.	23.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₄ H ₉ O ₃ P]	2-Methoxy-1,3,2-dioxaphosphorinane RN 31121-06-9	219.4	918.	-175.	-733.	Est	-29.	-121.
[C ₄ H ₁₀]	iso-C ₄ H ₁₀ RN 75-28-5	163.3	683.	-32.	-135.	[74SCO]	170.	712.
[C ₄ H ₁₀ N ₂]	Piperazine RN 110-85-0	224.2	938.	6.	25.(1)	Est	147.	617.
[C ₄ H ₁₀ N ₂]	c-C(CH ₃)(C ₂ H ₅)NHNH RN 4901-75-1	214.9**	899.**	32.	133.(17)	Est	182.5	764.
[C ₄ H ₁₀ O]	t-C ₄ H ₉ OH RN 75-65-0	193.7	810.	-75.	-312.(3)	[77PED/RYL]	97.	408.
[C ₄ H ₁₀ O]	n-C ₄ H ₉ OH RN 71-36-3	191.1	799.5	-66.	-275.(1)	[77PED/RYL]	109.	455.5
[C ₄ H ₁₀ O]	(C ₂ H ₅) ₂ O RN 60-29-7	200.2	838.	-60.	-252.(1)	[*80MAJ/WAC]	105.5	440.
[C ₄ H ₁₀ O ₂]	HO(CH ₂) ₄ OH RN 110-63-4	212**	887**	-102.	-427.(3)	Est	52.	216.
[C ₄ H ₁₀ O ₂]	CH ₃ OCH ₂ CH ₂ OCH ₃ RN 110-71-4	204.9	857.	-81.	-340.	[67LOU/LAI]	79.5	333.
[C ₄ H ₁₀ S]	(C ₂ H ₅) ₂ S RN 352-93-2	205.0	858.	-22.	-94.(1)	[77PED/RYL]	138.	578.
[C ₄ H ₁₀ S]	t-C ₄ H ₉ SH RN 75-66-1	196.9	824.	-26.	-110.(1)	[77PED/RYL]	142.5	596.
[C ₄ H ₁₁ N]	(CH ₃) ₂ (C ₂ H ₅)N RN 598-56-1	227.5	952.	-11.	-47.(2)	Est	127.	531.
[C ₄ H ₁₁ N]	(C ₂ H ₅) ₂ NH RN 109-89-7	225.9	945.	-17.	-73.(1)	[*79MAJ/SVO]	125.	512.
[C ₄ H ₁₁ N]	t-C ₄ H ₉ NH ₂ RN 75-64-9	220.8	924.	-29.	-121.(1)	[77PED/RYL]	116.	485.
[C ₄ H ₁₁ N]	sec-C ₄ H ₉ NH ₂ RN 13952-84-6	220.5	922.	-25.	-105.(1)	[*79MAJ/SVO]	120.	502.5
[C ₄ H ₁₁ N]	i-C ₄ H ₉ NH ₂ RN 78-81-9	218.8	915.	-24.	-100.(1)	[*79MAJ/SVO]	123.	515.
[C ₄ H ₁₁ N]	n-C ₄ H ₉ NH ₂ RN 109-73-9	218.4	914.	-22.	-92.(1)	[*79MAJ/SVO]	125.	524.
[C ₄ H ₁₁ NO]	NH ₂ (CH ₂) ₄ OH RN 13325-10-5	233.8	978.	-57.	-240.	Est	74.5	312.
[C ₄ H ₁₂ N ₂]	1,4-Diaminobutane RN 110-60-1	237.6	994.	-13.	-53.	Est	115.	483.
[C ₄ H ₁₂ OSi]	(CH ₃) ₃ SiOCH ₃ RN 1825-61-2	~203	~849	-112.	-468.(8)	Est	51.	213.
[C ₄ H ₁₄ OSi ₂]	((CH ₃) ₂ SiH) ₂ O RN 3277-26-7	~203	~849	-156.	-655.	Est	6.	26.
[C ₄ NiO ₄]	(CO) ₄ Ni RN 13463-39-3	180**	753**	-143.	-598.(4)	[77PED/RYL]	43.	179.
[C ₅ ClH ₃ N ₄]	6-Chloropurine RN 87-42-3	~208	~870	43.	179.	Est	200.	839.
[C ₅ ClH ₄ N]	4-Chloropyridine RN 626-61-9	217.8	911.	26.	108.	Est	174.	727.
[C ₅ ClH ₄ N]	3-Chloropyridine RN 626-60-8	214.8	899.	26.	108.	Est	177.	739.
[C ₅ ClH ₄ N]	2-Chloropyridine RN 109-09-1	214.4	897.	25.	103.	Est	176.	736.
[C ₅ FH ₄ N]	4-Fluoropyridine RN 694-52-0	216.6	906.	-14.	-57.	Est	135.	567.
[C ₅ FH ₄ N]	3-Fluoropyridine RN 372-47-4	214.3	897.	-13.	-56.	Est	138.	577.
[C ₅ FH ₄ N]	2-Fluoropyridine RN 372-48-5	210.6	881.	-16.	-68.	Est	139.	581.
[C ₅ F ₃ H ₇ O ₂]	CF ₃ CO ₂ (n-C ₃ H ₇) RN 383-66-4	185.7	777.	-254.	-1064.	Est	-74.	-311.
[C ₅ FeO ₅]	(CO) ₅ Fe RN 13463-40-6	~202	~845	-173.	-725.(1)	[77PED/RYL]	-9.5	-40.
[C ₅ HMnO ₅]	(CO) ₅ MnH RN 16972-33-1	201**	841**	-177.	-740.(10)	[82CON/ZAF]	-12.	-51.
[C ₅ H ₄ N ₂ O ₂]	4-Nitropyridine RN 1122-61-8	208.5	872.	33.	137.	Est	190.	795.
[C ₅ H ₄ N ₄]	Purine RN 120-73-0	219.3	917.5	55.	230.	Est	201.	842.5

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₅ H ₄ N ₄ O]	Hypoxanthine RN 68-94-0	~217	~907	12.	50.	[*/85CHI]	-161.	-672.
[C ₅ H ₅]	c-C ₅ H ₅ radical RN xxxxx	~199	~833	58.	242. (6)	[82McM/GOL]	229.	957.
[C ₅ H ₅ N]	Pyridine RN 110-86-1	220.8	924.	33.	140. (1)	[79KUD/KUD3]	178.	746.
[C ₅ H ₅ NNiO]	(C ₅ H ₅)NiNO RN 12071-73-7	200.1**	837.**					
[C ₅ H ₅ NO]	Pyridine-N-oxide RN 694-59-7	220.3	922.	14.5	61.	Est	160.	669.
[C ₅ H ₅ N ₅]	Adenine RN 73-24-5	223.5	935.	49.	207. (8)	[83KIR/DOM]	191.	802.
[C ₅ H ₅ N ₅ O]	Guanine RN 73-40-5	~223	~933	0.5	2.	[*/85CHI]	-143	-599.
[C ₅ H ₆]	c-C ₅ H ₆ RN 542-92-7	199.6	835.	31.	131. (4)	[77PED/RYL]	197.	826.
[C ₅ H ₆ N ₂]	2-Pyridinamine RN 504-29-0	223.8	936.	28.	118. (1)	[84BIC/PIL]	170.	711.
[C ₅ H ₆ N ₂]	3-Pyridinamine RN 462-08-8	221.0	925.	34.	144. (2)	[84BIC/PIL]	179.	749.
[C ₅ H ₆ N ₂]	4-Pyridinamine RN 504-24-5	230*	962*	31.	130. (1)	[84BIC/PIL]	167.	697.
[C ₅ H ₆ N ₂ O ₂]	Thymine RN 65-71-4	208.8	874.	-79.	-329. (4)	[77NAB/SAB]	78.	327.
[C ₅ H ₆ O]	2-Methylfuran RN 534-22-5	205.6	860.	-19.	-80.	Est	141.	590.
[C ₅ H ₆ S]	2-Methylthiophene RN 554-14-3	205.7	861.	20.	84. (1)	[77PEJ/RYL]	180.	753.
[C ₅ H ₈]	3,3-Dimethylcyclopropene RN 3907-06-0	203	849	50.	209.	[79AUE/BOW]	213.	890.
[C ₅ H ₈]	(E)-1,3-Pentadiene RN 2004-70-8	201.8**	844.**	18.	77. (1)	[77PED/RYL]	182.	763.
[C ₅ H ₈]	1-Methylcyclobutene RN xxxxx	201	841	28.	118.	[79AUE/BOW]	193.	807.
[C ₅ H ₈]	CH ₂ =CHC(CH ₃)=CH ₂ RN 78-79-5	200.4**	838.**	18.	75. (1)	[77PED/RYL]	183.	767.
[C ₅ H ₈]	(CH ₃) ₂ CHCOH RN 598-23-2	198**	828**	32.5	136.	[69BEN/CRU]	200.	838.
[C ₅ H ₈]	c-C ₃ H ₅ CH=CH ₂ RN 693-86-7	197.6	827.	36.	149. (1)	[*81CHI/HYM]	204.	852.
[C ₅ H ₈]	C ₂ H ₅ COCH ₃ RN 627-21-4	196**	820**	30.5	128. (4)	[77PED/RYL]	200.	838.
[C ₅ H ₈]	c-C ₅ H ₈ RN 142-29-0	183.4	767.5	9.	36. (2)	[82ALL/DOD]	191.	799.
[C ₅ H ₈ O]	c-C ₃ H ₅ COCH ₃ RN 765-43-5	205.1	858.	-28.	-119. (1)	[83FUC/SMI]	133.	555.
[C ₅ H ₈ O]	Cyclopentanone RN 120-92-3	198.8	832.	-46.	-194. (2)	[*76MEY/HOT]	121.	506.
[C ₅ H ₈ O ₂]	CH ₃ COCH=C(OH)CH ₃ RN 123-54-6	207.8	869.	-92.	-384. (1)	[79HAC/PIL]	66.	277.
[C ₅ H ₈ O ₂]	c-C ₃ H ₅ COOCH ₃ RN 2868-37-3	202.9	849.	-74.	-308.	[83FUC/SMI]	89.	373.
[C ₅ H ₉ N]	n-C ₄ H ₉ CN RN 110-59-8	194.0	812.	2.	10.	[82CHU/NGU]	174.	728.
[C ₅ H ₉ NO]	c-C ₄ H ₉ N(2-OCH ₃) RN 5264-35-7	225.9	945.	-36.	-152.	Est	103.	433.
[C ₅ H ₉ NO]	1-Methyl-2-pyrrolidinone RN 872-50-4	216.8	907.	-50.	-211.	[*72GAF]	98.	412.
[C ₅ H ₉ NO ₂]	c-C ₄ H ₉ NH(2-COOH) (L-Proline) RN 609-36-9	220.2	921.	-87.	-366. (4)	[78SAB/LAF]	58.	243.
[C ₅ H ₉ NO ₃]	CH ₃ CONHCH ₂ COOCH ₃ RN xxxxx	217.7	911.	-140.	-585.	Est	8.	34.
[C ₅ H ₉ NO ₄]	L-Glutamic Acid RN 617-65-2	216.5	906.	-120.	-503.	Est	29.	121.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₅ H ₉ O ₃ P]	4-Methyl-3,6,7,-trioxa-1-phospha- [2.2.2.]-octane RN 1449-91-8	210.0	879.	-136.	-571.	Est	19.	80.
[C ₅ H ₁₀]	(CH ₃) ₂ C=CHCH ₃ RN 513-35-9	196.4	822.	-10.	-42. (1)	[77PED/RYL]	159.	666.
[C ₅ H ₁₀ N ₂ O ₃]	L-Glutamine RN 585-21-7	218.4	914.	-74.5	-312.	Est	73.	304.
[C ₅ H ₁₀ O]	c-C ₄ H ₇ O(2-CH ₃) RN 96-47-9	203.6	852	-52.	-218.	Est	110.	461.
[C ₅ H ₁₀ O]	(C ₂ H ₅) ₂ CO RN 96-22-0	201.4	843.	-62.	-258. (1)	[*79SAL/PEA]	102.	429.
[C ₅ H ₁₀ O]	(i-C ₃ H ₇)COCH ₃ RN 563-80-4	201.1	841.	-63.	-262. (1)	[77PED/RYL]	102.	427.
[C ₅ H ₁₀ O]	c-C ₅ H ₁₀ O RN 142-68-7	199.7	835.5	-53.	-223. (1)	[77PED/RYL]	113.	471.5
[C ₅ H ₁₀ O]	n-C ₄ H ₉ CHO RN 110-62-3	192.6	806.	-55.	-231. (3)	[77PED/RYL]	118.	493.
[C ₅ H ₁₀ O ₂]	i-C ₃ H ₇ COOCH ₃ RN 547-63-7	201.6	843.	-109.	-456. (1)	[83FUC/SMI]	55.	231.
[C ₅ H ₁₀ O ₂]	CH ₃ COOC ₃ H ₇ RN 109-60-4	200.6	839.	-108.5	-454.	[70BEN/O'N]	57.	237.
[C ₅ H ₁₀ O ₂]	C ₃ H ₇ COOCH ₃ RN 623-42-7	200.1	837.	-108.	-452.	[70HOL/LOS]	57.	241.
[C ₅ H ₁₀ O ₂]	HCO ₂ (n-C ₄ H ₉) RN 592-84-7	194.8	815.	-103.	-430.	Est	68.	285.
[C ₅ H ₁₁ N]	CH ₃ CH=CHN(CH ₃) ₂ RN 6163-56-0	229.4	960.	6.	26.	Est	142.	596.
[C ₅ H ₁₁ N]	(CH ₃) ₂ C=NC ₂ H ₅ RN 15673-04-8	229.5**	960.**	-9.	-36. (9)	Est	127.5	534.
[C ₅ H ₁₁ N]	N-Methylpyrrolidine RN 120-94-5	228.7	957.	-0.5	-2. (2)	Est	136.5	571.
[C ₅ H ₁₁ N]	Piperidine RN 110-89-4	226.4	947.	-12.	-49. (2)	[77PED/RYL]	128.	535.5
[C ₅ H ₁₁ NO ₂]	(CH ₃) ₂ CHCH(NH ₂)COOH (L-Valine) RN 72-18-4	217.0	908.	-109.	-455. (1)	[77PED/RYL]	40.	167.
[C ₅ H ₁₁ NO ₂]	(CH ₃) ₂ NCOOC ₂ H ₅ RN 687-48-9	213.7	894.	-109.	-456.	Est	43.	180.
[C ₅ H ₁₁ NO ₂ S]	L-Methionine RN 59-51-8	221.4	926.	-99.	-414. (4)	[81SAB/MIN]	45.	190.
[C ₅ H ₁₂ O]	C ₂ H ₅ O(i-C ₃ H ₇) RN 625-54-7	203.5	851.	-68.	-286.	Est	94.	393.
[C ₅ H ₁₂ O]	t-C ₄ H ₉ OCH ₃ RN 1634-04-4	202.2	846.	-69.	-288. (4)	Est	95.	396.
[C ₅ H ₁₂ O ₂]	CH ₃ O(CH ₂) ₃ OCH ₃ RN 17081-21-9	213.8	894.					
[C ₅ H ₁₃ N]	(CH ₃)(C ₂ H ₅) ₂ N RN 616-39-7	230.0	962.	-17.	-70.	Est	119.	498.
[C ₅ H ₁₃ N]	(CH ₃) ₂ (i-C ₃ H ₇)N RN 996-35-0	229.8	961.	-24.	-99.	Est	112.	470.
[C ₅ H ₁₃ N]	(C ₂ H ₅)(i-C ₃ H ₇)NH RN 19961-27-4	227.4	951.	-25	-105.	Est	113.	474.
[C ₅ H ₁₃ N]	t-C ₅ H ₁₁ NH ₂ RN 594-39-8	222.3	930.	-31.5	-132.	Est	112.	468.
[C ₅ H ₁₃ N]	neo-C ₅ H ₁₁ NH ₂ RN 5813-64-9	219.3	917.5	-30.	-127.	Est	115.	480.5
[C ₅ H ₁₃ N]	n-C ₅ H ₁₁ NH ₂ RN 110-58-7	218.9	916.	-26.	-110.	Est	120.5	504.
[C ₅ H ₁₄ N ₂]	1,5-Diaminopentane RN 462-94-2	238.1	996.	-17.	-73.	Est	110.	461.
[C ₅ H ₁₄ N ₂]	(CH ₃) ₂ N(CH ₂) ₃ NH ₂ RN 109-55-7	236.4	989.	-7.	-30.	Est	122.	511.
[C ₅ H ₁₅ NSi]	(CH ₃) ₃ SiN(CH ₃) ₂ RN 18135-05-2	226	946	-59 (1)	-248 (3)	[77PED/RYL]	81.	336.
[C ₆ ClH ₅]	Chlorobenzene RN 108-90-7	181.7	760.	12.	51. (1)	[77PED/RYL]	196.	821.
[C ₆ ClH ₆ N]	2-Chloro-6-methylpyridine RN 18368-63-3	219**	916**	14.5	61.	Est	161.	675.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H (M)$		Reference	$\Delta_f H (MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₆ ClH ₆ N]	2-Chloro-4-methylpyridine RN 3678-62-4	218.6**	915.**	16.	66.	Est	163.	681.
[C ₆ ClH ₆ N]	4-Chlorobenzeneamine RN 106-47-8	208.6	873.	13.	55.	Est	170.	712.
[C ₆ ClH ₆ N]	3-Chlorobenzeneamine RN 108-42-9	207.2	867.	13.	55.	Est	172.	718.
[C ₆ ClH ₆ NO]	6-Chloro-1-methyl-2(1H)- pyridinone RN 17228-63-6	217.8	911.	-21.	-88.(16)	Est	127.	531.
[C ₆ ClH ₆ NO]	2-Chloro-6-methoxypyridine RN 17228-64-7	215.9	903.	-21.	-89.	Est	128.5	538.
[C ₆ CrO ₆]	(CO) ₆ Cr RN 15007-92-6	180**	753**	-217.	-908.(1)	[77PED/RYL]	-31.	-131.
[C ₆ FH ₅]	Fluorobenzene RN 462-06-6	182.6	764.	-28.	-116.(1)	[77PED/RYL]	155.	650.
[C ₆ FH ₆ N]	4-Fluorobenzeneamine RN 371-40-4	208.1	871.	-26.	-109.	Est	131.5	550.
[C ₆ FH ₆ N]	3-Fluorobenzeneamine RN 372-19-0	207.0	866.	-27.	-112.	Est	132.	552.
[C ₆ F ₂ H ₄]	1,2-Difluorobenzene RN 367-11-3	181.8	761.	-70.	-294.(1)	[77PED/RYL]	114.	475.
[C ₆ F ₂ H ₄]	1,3-Difluorobenzene RN 372-18-9	181.5	759.	-74.	-309.(1)	[77PED/RYL]	110.	462.
[C ₆ F ₂ H ₄]	1,4-Difluorobenzene RN 540-36-3	181.2	758.	-73.	-307.(1)	[77PED/RYL]	111.	465.
[C ₆ F ₃ H ₃]	1,2,4-C ₆ H ₃ F ₃ RN 367-23-7	181.4	759.	-115.	-482.(1)	Est	69.	289.
[C ₆ F ₃ H ₃]	1,3,5-C ₆ H ₃ F ₃ RN 372-38-3	181.	757.	-122.	-512.(3)	Est	62.	261.
[C ₆ F ₃ H ₄ N]	4-Trifluoromethylpyridine RN 3796-24-5	212.8	890.	-128.	-536.	Est	25.	104.
[C ₆ F ₃ H ₄ N]	3-Trifluoromethylpyridine RN 3796-23-4	212.6	889.	-128.	-537.	Est	25.	103.5
[C ₆ F ₃ H ₄ N]	2-Trifluoromethylpyridine RN 368-48-9	211.5	885.	-127.	-532.	Est	27.	113.
[C ₆ F ₃ H ₉ O ₂]	CF ₃ CO ₂ (n-C ₄ H ₉) RN 367-64-6	185.8	777.	-259.	-1085.	Est	-79.	-332.
[C ₆ F ₃ H ₁₀ NO]	CF ₃ CONH(n-C ₄ H ₉) RN 400-59-9	203.6	852.	-216.	-904.	Est	-54.	-226.
[C ₆ F ₄ H ₂]	1,2,3,4-C ₆ H ₂ F ₄ RN 551-62-2	181.1	758.	-152.	-638.(1)	Est	32.	134.
[C ₆ F ₄ H ₂]	1,2,3,5-C ₆ H ₂ F ₄ RN 2367-82-0	180.6	756.	-157.	-657.(1)	Est	28.	117.
[C ₆ F ₄ H ₂]	1,2,4,5-C ₆ H ₂ F ₄ RN 327-54-8	179.7	752.	-155.	-647.(3)	[78HAR/HEA]	31.	131.
[C ₆ F ₅ H]	C ₆ HF ₅ RN 363-72-4	179.9	753.	-193.	-806.(7)	[77PED/RYL]	-7.	-29.
[C ₆ F ₆]	C ₆ F ₆ RN 392-56-3	177.7	743.	-226.	-946.(8)	[79PRI/SAP]	-38.	-159.
[C ₆ H ₃ MnO ₅]	(CO) ₅ MnCH ₃ RN 13601-24-6	183	766	-180.	-753.(4)	[82CON/ZAF]	3.	11.
[C ₆ H ₃ O ₅ Re]	(CO) ₅ ReCH ₃ RN 14524-92-6	187**	782**	-183.	-765.(10)	[77TEL/RAB]	-4.	-17.
[C ₆ H ₄]	o-Benzyne RN xxxxx	213.0	891.	119.	497.	[80POL/HEH]	271.	1136.
[C ₆ H ₄ N ₂]	2-Pyridinecarbonitrile RN 100-70-9	208.1	871.	67.	281.(2)	[84BIC/PIL]	225.	940.
[C ₆ H ₄ N ₂]	3-Pyridinecarbonitrile RN 100-54-9	209.3	876.	66.	278.(2)	[84BIC/PIL]	222.	932.
[C ₆ H ₄ N ₂]	4-Pyridinecarbonitrile RN 100-48-1	210.3	880.	68.	284.(1)	[84BIC/PIL]	223.	934.
[C ₆ N ₅ NO]	Nitrosobenzene RN 586-96-9	204.8	857.					

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₆ H ₅ NO]	4-Pyridinecarboxaldehyde RN 872-85-5	215.2**	900.**	6.	24.	Est	156.	654.
[C ₆ H ₅ NO ₂]	C ₆ H ₅ NO ₂ RN 98-95-3	193.4	809.	16.	68. (1)	[77PED/RYL]	188.5	789.
[C ₆ H ₅ O]	C ₆ H ₅ O radical RN xxxxx	~204	~853	11.	48.	[82MCM/GOL]	173.	724.5
[C ₆ H ₆]	Benzene RN 71-43-2	181.3	759.	20.	83. (1)	[77PED/RYL]	204.	854.5
[C ₆ H ₆ IN]	3-IC ₆ H ₄ NH ₂ RN 626-01-7	208.9	874.	40.	166.	Est	196.	822.
[C ₆ H ₆ N]	C ₆ H ₅ NH radical RN xxxxx	219	916	57.	237. (8)	[82MCM/GOL]	198.4	830.
[C ₆ H ₆ N ₄]	6-Methylpurine RN 2004-03-7	~223	~933	42.5	178.	Est	185.	775.
[C ₆ H ₆ O]	C ₆ H ₅ OH RN 108-95-2	196.3	821.	-23.	-96. (1)	[78KUD/KUD]	146.	613.
[C ₆ H ₆ O]	(HCCCH ₂) ₂ O RN 6921-27-3	190.8	798.	71.	299.	Est	246.	1031.
[C ₆ H ₇ N]	4-Methylpyridine RN 108-89-4	225.2	942.	25.	104. (1)	[77PED/RYL]	165.	692.
[C ₆ H ₇ N]	2-Methylpyridine RN 109-06-8	225.0	942.	24.	99. (1)	[77PED/RYL]	164.	688.
[C ₆ H ₇ N]	3-Methylpyridine RN 108-99-6	224.1	938.	25.	106. (1)	[77PED/RYL]	167.	698.
[C ₆ H ₇ N]	(HCCCH ₂) ₂ NH RN 6921-28-4	216.1	904.	113.	472. (4)	Est	262.	1098.
[C ₆ H ₇ N]	C ₆ H ₅ NH ₂ RN 62-53-3	209.5	876.	21.	87. (1)	[78COL/BEN]	177.	740.5
[C ₆ H ₇ NO]	4-Methoxypyridine RN 620-08-6	227.6	952.	-3.	-13.	Est	135.	565.
[C ₆ H ₇ NO]	3-Methoxypyridine RN 7295-76-3	223.6	935.	-4.	-16.	Est	138.	578.5
[C ₆ H ₇ NO]	2-Methoxypyridine RN 1628-89-3	221.9	928.	-12.	-52.	Est	131.	550.
[C ₆ H ₇ NO]	1-Methyl-2-pyridinone RN 694-85-9	220.2	921.	-20.	-85. (10)	Est	125.	524.
[C ₆ H ₇ NO]	2-(OH)C ₆ H ₄ NH ₂ RN xxxxx	214.2	896.	-20.	-85.	Est	131.	549.
[C ₆ H ₇ NO]	3-(OH)C ₆ H ₄ NH ₂ RN 591-27-5	214.2	896.	-23.	-95.	Est	129.	539.
[C ₆ H ₇ NS]	4-(Methylthio)-pyridine RN 22581-72-2	225.5**	943.**	37.	155.	Est	177.	741.5
[C ₆ H ₇ NS]	2-(Methylthio)-pyridine RN 18438-38-5	222.0	929.	33.	138.	Est	177.	739.
[C ₆ H ₈]	1-Methyl-3-methylenecyclobutene RN 15082-13-0	212**	887**	48.	202.	[79AUE/BOW]	202.	845.
[C ₆ H ₈ N ₂]	1,2-C ₆ H ₄ (NH ₂) ₂ RN 95-54-5	212.8	890.	22.	92. (5)	Est	175.	732.
[C ₆ H ₈ N ₂]	1,3-C ₆ H ₄ (NH ₂) ₂ RN 108-45-2	222.4	930.5	21.	88.	Est	164.	687.5
[C ₆ H ₈ N ₂]	1,4-C ₆ H ₄ (NH ₂) ₂ RN 106-50-3	215.9	903.	23.	97.	Est	173.	723.
[C ₆ H ₈ O]	2,5-Dimethylfuran RN 625-86-5	209.1	875.	-30.	-125.	Est	127.	530.
[C ₆ H ₈ O ₂]	1,3-Cyclohexanedione RN 504-02-9	211.9	886.	-79.	-330.	Est	75.	313.
[C ₆ H ₈ O ₂]	1,2-Cyclohexanedione RN 765-87-7	204.8	857.	-70.	-293. (2)	Est	91.	380.
[C ₆ H ₉ N]	2,5-Dimethylpyrrole RN 625-84-3	218.4	914.	9.5	40. (1)	[77PED/RYL]	157.	656.
[C ₆ H ₉ N ₃ O ₂]	L-Histidine RN xxxxx	231.9	970.	-31.	-129.	Est	103.	431.
[C ₆ H ₉ O ₃ P]	2,8,9,-Trioxa-1-phosphadamantane RN 281-33-4	213.8	894.	-166.	-694.	Est	-14.	-58.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₆ H ₁₀]	1,3,3-Trimethylcyclopropene RN 3664-56-0	214.**	895**	41.	173.	[80WOL/HOL]	193.	808.
[C ₆ H ₁₀]	c-C ₃ H ₅ C(CH ₃)=CH ₂ RN 4663-22-3	209.0	874.	22.	94.	[82KOZ/MAS]	179.	750.
[C ₆ H ₁₀]	Cl ₃ CCl=C(CH ₃)-CH ₂ RN 1118-58-7	207.9**	870.**	11.	46.	Est (E)	169.	706.
[C ₆ H ₁₀]	CH ₂ =CH(CH ₃)C(CH ₂) ₂ RN 16906-27-7	206	862	21.	88.	[79AUE/BOW]	181.	756.
[C ₆ H ₁₀]	CH ₃ CH=C(CH ₃)CH=CH ₂ RN 4549-74-0	205.7**	860.6**	10.	43.	[80WOL/HOL]	170.	712.
[C ₆ H ₁₀]	CH ₂ =C(CH ₃)C(CH ₃)=CH ₂ RN 513-81-5	202.1**	846.**	10.5	44. (1)	[77PED/RYL]	174.	728.5
[C ₆ H ₁₀]	1,2-Dimethylcyclobutene RN 1501-58-2	201	841	17.	73.	[76JEN]	182.	762.
[C ₆ H ₁₀]	c-C ₅ H ₈ =CH ₂ RN 1528-30-9	200.8	840.	3.	12. (2)	[82ALL/DOD]	168.	702.
[C ₆ H ₁₀]	1-Methylcyclopentene RN 693-89-0	196.9	824.	-1.	-4. (1)	[82ALL/DOD]	168.	702.
[C ₆ H ₁₀]	c-C ₆ H ₁₀ RN 110-83-8	189.3	792.	-1.	-5. (1)	[77PED/RYL]	175.	733.
[C ₆ H ₁₀ O]	Cyclohexanone RN 108-94-1	201.4	843.	-54.	-226. (2)	[*76MEY/HOT]	110.	461.
[C ₆ H ₁₀ O]	{CH ₂ =CHCH ₂ } ₂ O RN 557-40-4	200.4	838.	-7.	-31.	Est	158.	661.
[C ₆ H ₁₀ O ₂]	CH ₃ COCH ₂ CH ₂ COCH ₃ RN 110-13-4	213.2	892.	-89.	-372.	Est	63.5	266.
[C ₆ H ₁₁ N]	(CH ₂ =CHCH ₂) ₂ NH RN 124-02-7	224.7	940.	34.	146. (6)	Est	175.	735.
[C ₆ H ₁₁ NO]	c-C ₅ H ₈ N(2-OCH ₃) RN 53687-79-9	228.1	954.	-42.	-176. (8)	Est	95.5	400.
[C ₆ H ₁₁ NO]	c-C ₅ H ₈ N(2-O)1-CH ₃ RN 931-20-4	219.3	917.5	-57.	-237. (3)	[*74BEA/MUE]	90.	375.5
[C ₆ H ₁₁ NO ₃]	CH ₃ CONHCH(CH ₃)COOCH ₃ (N-Acetyl alanine methyl ester) RN xxxxx	224.5	939.	-145.5	-609.	Est	-4.	-18.
[C ₆ H ₁₂]	(CH ₃) ₂ C=C(CH ₃) ₂ RN 563-79-1	199.0	833.	-16.	-69. (1)	[77PED/RYL]	150.	628.
[C ₆ H ₁₂]	CH ₃ CH=C(CH ₃)C ₂ H ₅ RN 922-61-2	198.2	829.	-15.	-64. (1) (E)	[77PED/RYL]	152.	638.
[C ₆ H ₁₂]	(CH ₃) ₂ C=CHCH ₂ CH ₃ RN 625-27-4	197.9	828.	-16.	-67. (1)	[77PED/RYL]	152.	635.
[C ₆ H ₁₂]	c-C ₆ H ₁₂ RN 110-82-7	169	707	-29.	-123. (1)	[77PED/RYL]	167.	700.
[C ₆ H ₁₂ N ₂]	1,4-Diazabicyclo[2.2.2]octane RN 280-57-9	229.0	958.	21.	89. (7)	[71RAP/WES]	158.	661.
[C ₆ H ₁₂ O]	t-C ₄ H ₉ COCH ₃ RN 75-97-8	202.3	846.	-69.	-290. (1)	[77PED/RYL]	94.	394.
[C ₆ H ₁₂ O]	2,2-Dimethyltetrahydrofuran	205.4	859.					
[C ₆ H ₁₂ O]	c-C ₆ H ₁₂ O (Oxepane) RN 592-90-5	202	845	-3.	-12.	Est	161.	673.
[C ₆ H ₁₂ O ₂]	t-C ₄ H ₉ COOCH ₃ RN 598-98-1	202.8	848.5	-117.	-491. (1)	[*82FUC]	45.5	190.5
[C ₆ H ₁₃ N]	(CH ₃) ₂ NC(CH ₃)=CHCH ₃ RN 52113-79-8	237	992	0.2	1.	Est	129.	539.
[C ₆ H ₁₃ N]	(CH ₃) ₂ C=CHN(CH ₃) ₂ RN xxxxx	229.5	960.					
[C ₆ H ₁₃ N]	1-Methylpiperidine RN 626-67-5	229.7	961.	-12.	-50. (4)	Est	124.	519.
[C ₆ H ₁₃ N]	n-C ₃ H ₇ CH=NC ₂ H ₅ RN 1611-12-7	225.3**	943.**	-5.	-21.	Est	135.	566.
[C ₆ H ₁₃ N]	c-C ₆ H ₁₁ NH ₂ RN 108-91-8	221.2	925.5	-25.	-105. (1)	[79STE]	120.	500.
[C ₆ H ₁₃ NO ₂]	L-C ₂ H ₅ CH(CH ₃)CH(NH ₂)COOH RN 73-32-5	218.9	916.	-116.	-487. (10)	Est	30.	127.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₆ H ₁₃ NO ₂]	(CH ₃) ₂ CHCH ₂ CH(NH ₂)COOH (L-Leucine) RN 61-90-5	218.1	912.5	-117.	-488. (3)	[77PED/RYL]	31.	129.5
[C ₆ H ₁₃ O ₃ P]	cis,cis-2-Methoxy-4,6-dimethyl- 1,3,2-dioxaphosphorinane RN 7735-82-2	226.2	946.	-182.	-760.	Est	-42.	-176.
[C ₆ H ₁₃ O ₃ P]	trans-2-Methoxy-cis,cis-4,6- dimethyl-1,3,2-dioxaphosphorinane RN 41821-91-4	225	941	-182.	-760.	Est	-41.	-171.
[C ₆ H ₁₄ N ₂ O ₂]	L-Lysine RN 56-87-1	230.3	963.5	-125.	-522.	Est	11.	44.5
[C ₆ H ₁₄ O]	(i-C ₃ H ₇) ₂ O RN 108-20-3	206.0	862.	-76.	-319. (2)	[*80MAJ/WAG]	84.	350.
[C ₆ H ₁₄ O]	C ₂ H ₅ O(t-C ₄ H ₉) RN 637-92-3	205.3	859.	-77.	-324.	Est	83.	347.
[C ₆ H ₁₄ O]	(n-C ₃ H ₇) ₂ O RN 111-43-3	202.3	846.	-70.	-293. (2)	[*80MAJ/WAG]	93.	391.
[C ₆ H ₁₄ O ₂]	CH ₃ O(CH ₂) ₄ OCH ₃ RN 13179-96-9	221.8	928.	-98.	-408.	Est	46.	194.
[C ₆ H ₁₄ OSi]	CH ₂ =C(CH ₃)OSi(CH ₃) ₃ RN 1833-53-0	221.	925.	-104.	-437.	Est	40.	168.
[C ₆ H ₁₄ O ₃]	CH ₃ (OCH ₂ CH ₂) ₂ OCH ₃ RN 111-96-6	219.4	918.	-119.	-498.	Est	27.	114.
[C ₆ H ₁₄ S]	(i-C ₃ H ₇) ₂ S RN 625-80-9	209.6	877.	-34.	-142. (1)	[77PED/RYL]	122.	511.
[C ₆ H ₁₄ S]	(n-C ₃ H ₇) ₂ S RN 111-47-7	206.5	864.	-30.	-125. (1)	[77PED/RYL]	129.	541.
[C ₆ H ₁₅ N]	(C ₂ H ₅) ₃ N RN 121-44-8	232.3	972.	-22.	-93. (1)	[*79MAJ/SVO]	111.	465.
[C ₆ H ₁₅ N]	(CH ₃) ₂ (t-C ₄ H ₉)N RN 918-02-5	232.0	971.	-24.	-102.	Est	109.	457.
[C ₆ H ₁₅ N]	(i-C ₃ H ₇) ₂ NH RN 108-18-9	230.2	963.	-34.	-144. (1)	[*79PET/MAJ]	101.	423.
[C ₆ H ₁₅ N]	(n-C ₃ H ₇) ₂ NH RN 142-84-7	227.5	952.	-28.	-116. (1)	[77PED/RYL]	110.	462.
[C ₆ H ₁₅ N]	n-C ₆ H ₁₃ NH ₂ RN 111-26-2	218.9	916.	-31.	-130.	Est	116.	484.
[C ₆ H ₁₅ NO]	NH ₂ (CH ₂) ₆ OH RN 4048-33-3	231.0**	966.5**	-67.	-279.	Est	68.	284.5
[C ₆ H ₁₅ O ₄ P]	OP(OC ₂ H ₅) ₃ RN 78-40-0	~217	~910	-284.	-1187. (6)	[77PED/RYL]	-135.	-565.
[C ₆ H ₁₅ P]	(C ₂ H ₅) ₃ P RN 554-70-1	231.7**	969.**	-12.	-49. (13)	[77PED/RYL]	122.	512.
[C ₆ H ₁₆ N ₂]	1,6-diaminohexane RN 124-09-4	237.7	994.4	-22.	-94.	Est	105.5	441.5
[C ₆ H ₁₆ N ₂]	(CH ₃) ₂ N(CH ₂) ₂ N(CH ₃) ₂ RN 110-18-9	235.7	986.	-4.	-16. (2)	Est	126.	528.
[C ₆ H ₁₇ NSi]	(CH ₃) ₃ SiCH ₂ N(CH ₃) ₂ RN 18182-40-6	231.5	968.	-49.	-207.	Est	85.	354.
[C ₆ H ₁₈ OSi ₂]	((CH ₃) ₃ Si) ₂ O RN 107-46-0	~203	~849	-185.	-777. (6)	[77PED/RYL]	-23.	-96.
[C ₆ MoO ₆]	(CO) ₆ Mo RN 13939-06-5	185**	774**	-219.	-916. (2)	[77PED/RYL]	-38.	-160.
[C ₆ O ₆ V]	(CO) ₆ V RN 20644-87-5	194.5**	814.**	-204.	-854. (29)	[67BID/MCI]	-33.	-138.
[C ₆ O ₆ W]	(CO) ₆ W RN 14040-11-0	184**	770**	-211.	-883. (3)	[77PED/RYL]	-29.	-123.
[C ₇ ClH ₅ O]	4-ClC ₆ H ₄ CHO RN 104-88-1	200.2	838.	-16.	-69.	Est	149.	623.
[C ₇ ClH ₁₀ N]	3-Chloro-1-azabicyclo[2.2.2]- oct-2-ene RN xxxxx	224.0**	937.**	25.	104. (10)	Est	166.5	697.
[C ₇ ClH ₁₂ N]	3-Chloro-1-azabicyclo[2.2.2]- octane RN 42332-45-6	225.8**	945.**	-10.5	-44.	Est	129.	541.
[C ₇ ClH ₁₄ N]	c-C ₅ H ₉ N,2-CH ₂ Cl,1-CH ₃ RN 49665-74-9	227.6**	952.**	-23.	-97.	Est	115.	481.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H (M)$		Reference	$\Delta_f H (MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₇ CoH ₅ O ₂]	(C ₅ H ₅)Co(CO) ₂ RN 12078-25-0	~204**	853**					
[C ₇ CrH ₅ NO ₃]	(C ₅ H ₅)Cr(CO) ₂ NO RN 36312-04-6	196.9**	824.**					
[C ₇ D ₃ H ₅]	C ₆ H ₅ CD ₃ RN 1124-18-1	189.8	794.	12.	50. (1)	[77PED/RYL]	188.	786.
[C ₇ FH ₅ O]	4-FC ₆ H ₄ CHO RN 459-57-4	199.2	833.	-56.	-235.	Est	110.	462.
[C ₇ FH ₅ O]	3-FC ₆ H ₄ CHO RN 456-48-4	196.4	822.	-56.	-236.	Est	113.	472.
[C ₇ FH ₆]	3-FC ₆ H ₄ CH ₂ radical RN xxxxx	~200	~837					
[C ₇ FH ₇]	3-FC ₆ H ₄ CH ₃ RN 352-70-5	189.3	792.	-36.	-150.	Est	140.	587.
[C ₇ FH ₇]	2-FC ₆ H ₄ CH ₃ RN 95-52-3	186.6	781.	-36.	-149.	Est	143.	600.
[C ₇ FH ₇]	4-FC ₆ H ₄ CH ₃ RN 352-32-9	185.8	777.	-35.	-148. (1)	[77PED/RYL]	144.5	605.
[C ₇ FH ₁₂ N]	3-Fluoro-1-azabicyclo[3.2.1]- octane RN xxxxx	228.1**	954.**	22.	94.	Est	160.	670.
[C ₇ F ₂ H ₁₁ N]	3,3-Difluoro-1-azabicyclo- [2.2.2]octane RN xxxxx	221.8**	928.**	-101.	-423.	Est	43.	179.
[C ₇ F ₃ H ₆ N]	3-CF ₃ C ₆ H ₄ NH ₂ RN 98-16-8	204.2	854.	-142.	-595.	Est	19.	81.
[C ₇ H ₅ N]	C ₆ H ₅ CN RN 100-47-0	195.9	820.	52.	219.	[82CHU/NGU]	222.	929.
[C ₇ H ₅ O ₂ Rh]	(C ₅ H ₅)Rh(CO) ₂ RN 12192-97-1	212**	887.**					
[C ₇ H ₆ N ₂]	m-NCC ₆ H ₄ NH ₂ RN 2237-30-1	200.7	840.	53.	222.	Est	218.	912.
[C ₇ H ₆ O]	4-Methylene-2,5-cyclohexadiene- 1-one RN 502-87-4	~222	~929	10.	40. (4)	Est	153.	641.
[C ₇ H ₆ O]	2,4,6-Cycloheptatriene-1-one RN 539-80-0	219	918	10.5	44. (3)	[77PED/RYL]	157.	656.
[C ₇ H ₆ O]	C ₆ H ₅ CHO RN 100-52-7	200.2	838.	-9.	-37. (2)	[77PED/RYL]	157.	655.
[C ₇ H ₆ O ₂]	C ₆ H ₅ COOH RN 65-85-0	198.2	829.	-70.	-294. (2)	[77PED/RYL]	97.	407.
[C ₇ H ₇]	c-C ₇ H ₇ radical RN 3551-27-7	199.4	834.	65.	271. (8)	[82MCM/GOL]	227.	951.
[C ₇ H ₇]	C ₆ H ₅ CH ₂ RN 2154-56-5	199.1	833.	49.	204.	[81TSA]	215.	901.
[C ₇ H ₇ N]	3,4-Cyclobutenopyridine RN xxxxx	225.9**	945.**	60.	252.	Est	200.	837.
[C ₇ H ₇ N]	2,3-Cyclobutenopyridine RN xxxxx	223.3**	934.**	60.	250.	Est	202.	846.
[C ₇ H ₇ N]	4-Vinylpyridine RN 100-43-6	223.2**	934.**	48.	202.	Est	191.	798.
[C ₇ H ₇ NO]	1-(4-Pyridinyl)-ethanone RN 1122-54-9	217.4	910.	-6.	-26.	Est	142.	594.
[C ₇ H ₇ NO]	1-(3-Pyridinyl)-ethanone RN 350-03-8	217.2	909.	-6.	-26.	Est	142.	595.
[C ₇ H ₈]	C ₆ H ₅ CH ₃ RN 108-88-3	189.8	794.	12.	50. (1)	[77PED/RYL]	188.	786.
[C ₇ H ₈ O]	C ₆ H ₅ OCH ₃ RN 100-66-3	200.3	838.	-16.	-68. (1)	[77PED/RYL]	149.	624.
[C ₇ H ₉ N]	2,3-Dimethylpyridine RN 583-61-9	226.2	946.	16.	68. (1)	[77PED/RYL]	156.	652.
[C ₇ H ₉ N]	2,4-Dimethylpyridine RN 108-47-4	227.3	951.	15.	64. (2)	[77PED/RYL]	153.	643.
[C ₇ H ₉ N]	2,5-Dimethylpyridine RN 589-93-5	226.2	946.	16.	67. (1)	[77PED/RYL]	155.5	651.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₇ H ₉ N]	2,6-Dimethylpyridine RN 108-48-5	228.2	955.	14.	59. (2)	[77PED/RYL]	152.	634.
[C ₇ H ₉ N]	3,4-Dimethylpyridine RN 583-58-4	226.0	946.	17.	70. (1)	[77PED/RYL]	157.	654.
[C ₇ H ₉ N]	3,5-Dimethylpyridine RN 591-22-0	225.5	943.	17.	73. (1)	[77PED/RYL]	158.	661.
[C ₇ H ₉ N]	2-Ethylpyridine RN 100-71-0	226.2	946.	19.	81.	Est	159.	665.
[C ₇ H ₉ N]	3-Ethylpyridine RN 536-78-7	223.9	937.	20.5	86.	Est	162.	679.
[C ₇ H ₉ N]	4-Ethylpyridine RN 536-75-4	224.6**	940.**	20.	83.	Est	161.	672.
[C ₇ H ₉ N]	C ₆ H ₅ NiCH ₃ RN 100-61-8	218.1	912.5	20.	85.	[78COL/BEN]	168.	702.5
[C ₇ H ₉ N]	C ₆ H ₅ CH ₂ NH ₂ RN 100-46-9	216.8	907.	20.	84. (3)	[77CAR/LAY]	169.	707.
[C ₇ H ₉ N]	4-CH ₃ C ₆ H ₄ NH ₂ RN 106-49-0	213.7	894.	14.	59.	Est	166.	695.
[C ₇ H ₉ N]	3-CH ₃ C ₆ H ₄ NH ₂ RN 108-44-1	213.4	893.	13.	54. (2)	Est	165.	691.
[C ₇ H ₉ NO]	Pyridine-2-methoxymethyl RN 23579-92-2	226.0**	945.5**	-5.	-22.	Est	134.	562.5
[C ₇ H ₉ NO]	3-CH ₃ OC ₆ H ₄ NH ₂ RN 536-90-3	217.6	910.	-16.	-67.	Est	132.	553.
[C ₇ H ₉ NO]	2-CH ₃ OC ₆ H ₄ NH ₂ (o-Anisidine) RN 90-04-0	214.7	898.	-14.	-57.	Est	137.	575.
[C ₇ H ₉ NO]	4-CH ₃ OC ₆ H ₄ NH ₂ RN 104-94-9	214.3	897.	-14.	-58.	Est	137.5	575.
[C ₇ H ₉ NS]	3-CH ₃ SC ₆ H ₄ NH ₂ RN 1783-81-9	214.5	897.	24.	102.	Est	175.5	735.
[C ₇ H ₁₀]	Bicyclo[2.2.1]hept-2-ene RN 498-66-8	200.4	838.	21.5	90. (4)	[80ROG/CHO]	187.	782.
[C ₇ H ₁₀ N ₂]	N,N-Dimethyl-2-pyridinamine RN 5683-33-0	229.2	959.	31.5	132.	[84BIC/PIL]	168.	703.
[C ₇ H ₁₀ N ₂]	N,N-Dimethyl-3-pyridinamine RN 18437-57-5	229.9**	962.**	38.	158.	[84BIC/PIL]	174.	726.
[C ₇ H ₁₀ N ₂]	N,N-Dimethyl-4-pyridinamine RN 1122-58-3	236.2	980.	34.	144.	[84DIC/PIL]	163.5	686.
[C ₇ H ₁₀ O]	(c-C ₃ H ₅) ₂ CO RN 1121-37-5	210.7	881.5	39.	163.	Est	194.	811.5
[C ₇ H ₁₁ N]	1-Azabicyclo[2.2.2]oct-2-ene RN 13929-94-7	228.5**	956.**	37.	156.	Est	174.5	730.
[C ₇ H ₁₁ NO]	1-Azabicyclo[2.2.2]octan-3-one RN 3731-38-2	221.9**	928.**	-28.	-116.	Est	116.	486.
[C ₇ H ₁₂]	(CH ₃) ₂ C=CHC(CH ₃)=CH ₂ RN xxxxx	213.1**	892.	4.	17.	[79AUE/BOW]	157.	655.
[C ₇ H ₁₂]	1-Methylcyclohexene RN 591-49-1	198.8	832.	-10.	-43. (1)	[77PED/RYL]	157.	655.
[C ₇ H ₁₂]	c-C ₅ H ₆ -1,2-(CH ₃) ₂ RN 765-47-9	198.1	829.	-10.	-41.	[82ALL/DOD]	158.	660.
[C ₇ H ₁₃ N]	1-Azabicyclo[2.2.2]octane (Quinuclidine) RN 100-76-5	232.1	971.	-1.	-4. (1)	[77PED/RYL]	132.6	555.
[C ₇ H ₁₃ N]	Bicyclo[2.2.1]heptan-2-amine, exo (2-Aminonorbornane) RN 7242-92-4	221.7**	927.**	-8.	-32. (1)	Est	136.	570.
[C ₇ H ₁₃ N]	Bicyclo[2.2.1]heptan-2-amine, endo (2-Aminonorbornane) RN 31002-73-0	221.7**	927.**	-7.	-28. (1)	Est	137.	574.
[C ₇ H ₁₄]	(CH ₃) ₂ C=CHC(CH ₃) ₂ RN xxxxx	196.1	820.	-20.	-84. (1)	[77PED/RYL]	149.5	626.
[C ₇ H ₁₄ N ₂]	3-Amino-1-azabicyclo[2.2.2] octane RN 6238-14-8	231.8**	970.**	4.	17.	Est	138.	577.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₇ H ₁₄ O]	(i-C ₃ H ₇) ₂ CO RN 565-80-0	204.9	857.	-74.	-311.(1)	[77PED/RYL]	87.	363.
[C ₇ H ₁₅ N]	(CH ₃) ₂ NC(C ₂ H ₅)=CHCH ₃ RN 78733-73-0	236.4	989.	-2.	-10.	Est	127.	531.
[C ₇ H ₁₆ O]	(i-C ₃ H ₇) ₂ O(t-C ₄ H ₉) RN 17348-59-3	208.8**	874.**	-81.	-339.	Est	76.	317.
[C ₇ H ₁₆ O ₂]	CH ₃ O(CH ₂) ₅ OCH ₃ RN 111-89-7	221.8	928.	-104.	-436.	Est	40.	167.
[C ₇ H ₁₇ N]	(C ₂ H ₅) ₂ (n-C ₃ H ₇)N RN 4458-31-5	232.0**	971.**	-27.	-114.	Est	106.	445.
[C ₇ H ₁₇ N]	(CH ₃) ₂ (neo-C ₅ H ₁₁)N RN 10076-31-0	229.9	962.	-28.	-118.	Est	107.5	450.
[C ₇ H ₁₇ N]	n-C ₇ H ₁₅ NH ₂ RN 111-68-2	219.0	916.	-36.	-151.	Est	111.	463.
[C ₇ H ₁₈ N ₂]	(CH ₃) ₂ N(CH ₂) ₃ N(CH ₃) ₂ RN 110-95-2	238.8	999.	-7.	-29.	Est	120.	502.
[C ₇ H ₁₈ N ₂]	1,7-Diaminoheptane RN 646-19-5	238.	996.	-27.	-115.	Est	100.	419.
[C ₇ H ₁₉ NSi]	(CH ₃) ₃ Si(CH ₂) ₂ N(CH ₃) ₂ RN 23138-94-5	231.8	970.	-54.	-228.	Est	79.	332.
[C ₈ F ₃ H ₅ O]	p-CF ₃ C ₆ H ₄ CHO RN 455-19-6	191.0	799.	-172.	-721.	Est	2.	10.
[C ₈ FeH ₈ O ₂]	(C ₅ H ₅)Fe(CO) ₂ CH ₃ RN 12080-06-7	190.6**	797.**					
[C ₈ H ₅ NO]	4-(CN)C ₆ H ₄ CHO RN 105-07-7	187.0	782.	25.5	107.	Est	204.	855.
[C ₈ H ₆ N ₂]	Cinnoline RN 253-66-7	223.2	934.	81.	338.(10)	Est	223.	934.
[C ₈ H ₆ N ₂]	Quinoxaline RN 91-19-0	214.4	897.	63.	262.(4)	[81STE/BAR]	214.	895.
[C ₈ H ₈]	C ₆ H ₅ CH=CH ₂ RN 100-42-5	202.0	845.	35.	148.(1)	[77PED/RYL]	199.	833.
[C ₈ H ₈]	1,2-C ₆ H ₄ (=CH ₂) ₂ RN xxxxx	214.8	899.	55.	230.(17)	[81POL/RAI]	206.	861.
[C ₈ H ₈]	1,4-C ₆ H ₄ (=CH ₂) ₂ RN xxxxx	215.7	902.	56.	234.(17)	[81POL/RAI]	207.	865.
[C ₈ H ₈ O]	C ₆ H ₅ COCH ₃ RN 98-86-2	205.4	859.	-21.	-87.(2)	[77PED/RYL]	140.	584.
[C ₈ H ₈ O]	4-(CH ₃)C ₆ H ₄ CHO RN 104-87-0	203.7	852.	-18.	-75.	Est	144.	603.
[C ₈ H ₈ O ₂]	4-CH ₃ OC ₆ H ₄ CHO RN 123-11-5	213.5	893.	-48.5	-203.(5)	[77PED/RYL]	104.	434.
[C ₈ H ₈ O ₂]	C ₆ H ₅ CO ₂ CH ₃ RN 95-58-3	203.7	852.	-69.	-288.(7)	[77PED/RYL]	93.	390.
[C ₈ H ₉]	C ₆ H ₅ CHCH ₃ radical RN xxxxx	201	841	44.	184.	[82MAU]	209.	875.
[C ₈ H ₉ N]	3,4-Cyclopentenopyridine RN xxxxx	226.8**	949.**	27.	113.	Est	166.	695.
[C ₈ H ₉ N]	2,3-Cyclopentenopyridine RN xxxxx	225.8**	945.**	27.	111.	Est	166.	696.
[C ₈ H ₁₀]	m-Xylene RN 108-38-3	195.9	820.	4.	17.(1)	[77PED/RYL]	174.	727.
[C ₈ H ₁₀]	o-Xylene RN 95-47-6	193.3	809.	4.	18.(1)	[77PED/RYL]	177.	739.
[C ₈ H ₁₀]	p-Xylene RN 106-42-3	192.0	803.	4.	18.(1)	[77PED/RYL]	178.	745.
[C ₈ H ₁₀]	C ₂ H ₅ C ₆ H ₅ RN 100-41-4	191.6	802.	7.	29.(1)	[77PED/RYL]	181.	757.
[C ₈ H ₁₁ N]	2-Isopropylpyridine RN 75981-47-4	227.2	951.	13.	56.	Est	152.	635.
[C ₈ H ₁₁ N]	C ₆ H ₅ N(CH ₃) ₂ RN 121-69-7	223.4	935.	24.	101.(3)	[82FUR/SAK]	166.	696.
[C ₈ H ₁₁ N]	C ₆ H ₅ NHC ₂ H ₅ RN 103-69-5	221.8	928.	13.	56.(6)	[69BEN/CRU]	157.	658.
[C ₈ H ₁₁ N]	3-C ₂ H ₅ C ₆ H ₄ NH ₂ RN 587-02-0	214.0	895.	6.	27.	Est	158.	662.
[C ₈ H ₁₁ P]	C ₆ H ₅ P(CH ₃) ₂ RN 672-66-2	229.6	961.	19.5	82.	Est	156.	651.
[C ₈ H ₁₂]	(o-C ₃ H ₅) ₂ C=CH ₂ RN 822-93-5	216.5	906.	51.	213.	[70BEN/O`N]	200.	837.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₈ H ₁₂]	2-Methylenebicyclo[2.2.1]heptane RN 497-35-8	207**	866**	12.	50.	[79AUE/BOW]	171.	714.
[C ₈ H ₁₂]	2-Methylbicyclo[2.2.1]hept-2-ene RN xxxxx	206	862	11.	46.	Est	171.	714.
[C ₈ H ₁₃ N]	1-Azabicyclo[2.2.2]- oct-2-ene,3-methyl RN xxxxx	231.0**	966.5**	30.	124.	Est	164.	687.5
[C ₈ H ₁₃ N]	1-Azabicyclo[2.2.2]- octane,3-methylene RN 22207-84-7	230.1**	963.**	20.	84.	Est	156.	651.
[C ₈ H ₁₄]	(CH ₃) ₂ C=C(CH ₃)C(CH ₃)=CH ₂ RN xxxxx	210.6**	881.**	-3.	-13.	[79AUE/BOW]	152.	636.
[C ₈ H ₁₄ O]	c-C ₆ H ₁₁ COCH ₃ RN 823-76-7	202.4	847.	-65.	-273.	Est	98.	410.
[C ₈ H ₁₄ O ₂]	c-C ₆ H ₁₁ COOCH ₃ RN 4630-82-4	203.7	852.	-111.	-466.	Est	51.	212.
[C ₈ H ₁₅ N]	3-Methyl-1-azabicyclo[2.2.2]- octane RN-695-88-5	231.7**	969.**	-8.	-33.	Est	126.	528.
[C ₈ H ₁₅ N]	1,4,4-Trimethyl-1,2,3,4-tetra- hydropyridine RN 35079-50-6	234.2**	980.**					
[C ₈ H ₁₅ NO]	c1s-3-Aminobicyclo[2.2.2]octan- 2-ol RN 17997-65-8	223.9	937.	-57.	-240.	Est	84.	353.
[C ₈ H ₁₅ NO]	trans-3-Aminobicyclo[2.2.2]- octan-2-ol RN 40335-14-6	220.6	923.	-59.0	-248.	Est	86.	359.
[C ₈ H ₁₆ O ₄]	1,4,7,10-Tetraoxa- cyclododecane (12-Crown-4) RN 294-93-9	221.6	927.	-151.	-631.	[82BYS/MAN]	-7.	-29.
[C ₈ H ₁₇ N]	1,4,4-Trimethylpiperidine RN 1003-84-5	230.8**	966.**					
[C ₈ H ₁₈ O]	(n-C ₄ H ₉) ₂ O RN 142-96-1	203.7	852.	-79.5	-333. (1)	[*80MAJ/WAG]	82.	345.
[C ₈ H ₁₈ O]	(sec-C ₄ H ₉) ₂ O RN 6863-58-7	209.0	874.	-88.	-370. (2)	[77PED/RYL]	68.	286.
[C ₈ H ₁₈ O]	(t-C ₄ H ₉) ₂ O See References to Table 1: 75PIT/BUR							
[C ₈ H ₁₈ O ₄]	CH ₃ (OCH ₂ CH ₂) ₃ OCH ₃ RN 112-49-2	224.1	938.	-157.	-656.	Est	-15.	-64.
[C ₈ H ₁₈ S]	(t-C ₄ H ₉) ₂ S RN 107-47-1	212.8	890.	-45.	-189. (1)	[77PED/RYL]	108.	451.
[C ₈ H ₁₈ S]	(n-C ₄ H ₉) ₂ S RN 544-40-1	208.7	873.	-40.	-167. (1)	[77PED/RYL]	117.	490.
[C ₈ H ₁₉ N]	(i-C ₃ H ₇) ₂ (C ₂ H ₅)N RN 7087-68-5	235.3	984.	-33.	-140.	Est	97.	405.5
[C ₈ H ₁₉ N]	(t-C ₄ H ₉) ₂ NH RN 21981-37-3	233.2	976.	-41.	-172. (3)	[81SUR/HAC]	91.	382.
[C ₈ H ₁₉ N]	(sec-C ₄ H ₉) ₂ NH RN 626-23-3	231.8	970.	-37.5	-157.	Est	96.	403.
[C ₈ H ₁₉ N]	(CH ₃) ₃ C(CH ₂) ₂ N(CH ₃) ₂ RN 15673-04-8	230.4	964.	-36.	-149. (3)	Est	100.	417.
[C ₈ H ₁₉ N]	(i-C ₄ H ₉) ₂ NH RN 110-96-3	228.6	956.	-43.	-179. (8)	[73PEP/GAF]	94.	395.
[C ₈ H ₁₉ N]	(n-C ₄ H ₉) ₂ NH RN 111-92-2	228.4	956.	-37.5	-157. (1)	[77PED/RYL]	100.	417.
[C ₈ H ₁₉ N]	n-(C ₈ H ₁₇)NH ₂ RN 111-86-4	220.4**	922.**	-41.	-172.	Est	104.	436.
[C ₈ H ₂₀ N ₂]	(CH ₃) ₂ N(CH ₂) ₄ N(CH ₃) ₂ RN 111-51-3	240.4	1006.	-12.	-51. (1)	Est	113.	473.
[C ₈ H ₂₁ NSi]	(CH ₃) ₃ Si(CH ₂) ₃ N(CH ₃) ₂ RN 28247-29-2	231.8	970.	-59.	-248.	Est	75.	312.
[C ₈ H ₂₁ NSi]	(CH ₃) ₂ (t-C ₄ H ₉)SiN(CH ₃) ₂ RN 66365-05-7	229.7	961.	-68.	-286. (23)	Est	68.	283.
[C ₈ H ₂₂ OSi ₂]	((CH ₃) ₃ SiCH ₂) ₂ O See References to Table 1: 75PIT/BUR							
[C ₉ ClH ₉]	4-ClC ₆ H ₄ C(CH ₃)=CH ₂ RN 1712-70-5	205.0	858.	19.	81.	Est	180.	753.
[C ₉ CrH ₈ O ₃]	(C ₅ H ₅)Cr(CO) ₃ CH ₃ RN 41311-89-1	206**	862**					
[C ₉ FH ₉]	4-FC ₆ H ₄ C(CH ₃)=CH ₂ RN 350-40-3	206.7	865.	-20.5	-86.	Est	138.	579.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₉ H ₇ MnO ₃]	(CH ₃ C ₅ H ₄)Mn(CO) ₃ RN 12108-13-3	200.6**	839.**	-121.	-508.	Est	44.	183.
[C ₉ H ₇ N]	Quinoline RN 91-22-5	226.5	948.	50.	211.(1)	[79VIS]	190.	793.
[C ₉ H ₇ N]	Isoquinoline RN 119-65-3	225.9	945.	50.	208.(1)	[79VIS/WIL]	190.	793.
[C ₉ H ₇ NO]	Quinoline-1-oxide RN 1613-37-2	224.6	940.	31.	131.	Est	172.	721.
[C ₉ H ₉ N]	(HCCCH ₂) ₃ N RN 6921-29-5	220.2	921.	174.	727.	Est	319.	1336.
[C ₉ H ₁₀]	C ₆ H ₅ C(CH ₃)=CH ₂ RN 98-83-9	207.0	866.	27.	113.	[69BEN/CRU]	186.	777.
[C ₉ H ₁₀ O]	(4-CH ₃)C ₆ H ₄ COCH ₃ RN xxxxx	208.7	873.	-29.	-120.	Est	128.	535.
[C ₉ H ₁₁]	C ₆ H ₅ C(CH ₃) ₂ radical RN xxxxx	202.4	847.	33.	139.	[82MCM/GOL]	202.	846.
[C ₉ H ₁₁]	C ₆ H ₅ (CHC ₂ H ₅) radical RN xxxxx	~202	~845	39	164	[82MAU]	203.	849.
[C ₉ H ₁₁ N]	2,3-Cyclohexenopyridine RN 10500-57-9	227.7**	953.**	18.	74.	Est	156.	651.
[C ₉ H ₁₁ N]	3,4-Cyclohexenopyridine RN 36566-06-6	227.7**	953.**	18.	76.	Est	156.	653.
[C ₉ H ₁₁ NO ₂]	C ₆ H ₅ CH ₂ CH(NH ₂)COOH (L-Phenylalanine) RN 150-30-1	216.5	906.	-75.	-313.(1)	[77PED/RYL]	74.	311.
[C ₉ H ₁₁ NO ₃]	L-Tyrosine RN xxxxx	222.3	930.	-116.	-486.	Est	27.	114.
[C ₉ H ₁₂]	Mesitylene RN 108-67-8	200.7	840.	-4.	-16.(1)	[77PED/RYL]	161.	674.
[C ₉ H ₁₂]	n-C ₃ H ₇ C ₆ H ₅ RN 103-65-1	192.4	805.	2.	8.(1)	[77PED/RYL]	175.	733.
[C ₉ H ₁₂]	i-C ₃ H ₇ C ₆ H ₅ RN 98-82-8	192.1	804.	1.	4.(1)	[77PED/RYL]	174.5	730.
[C ₉ H ₁₂ N ₂ O ₆]	Uridine RN 58-96-8	~208	~870	-223.	-935.	Est	-66.	-275.
[C ₉ H ₁₂ O ₃]	1,3,5-C ₆ H ₃ (OCH ₃) ₃ RN 621-23-8	220.6	923.	-90.5	-379.	Est	54.5	228.
[C ₉ H ₁₃ N]	2,6-Diethylpyridine RN 935-28-4	231.1	967.	4.5	19.(2)	Est	139.	582.
[C ₉ H ₁₃ N]	C ₆ H ₅ CH ₂ N(CH ₃) ₂ RN 103-83-3	228.1**	954.**	20.	84.	Est	158.	660.
[C ₉ H ₁₃ N]	C ₆ H ₅ N(CH ₃)(C ₂ H ₅) RN 613-97-8	227.1	950.	17.	71.	Est	156.	651.
[C ₉ H ₁₃ N]	2-t-Butylpyridine RN 5944-41-2	227.4**	951.**	7.	28.	Est	145.	607.
[C ₉ H ₁₃ N]	4-t-Butylpyridine RN 3978-81-2	225.9	945.	8.	32.	Est	147.	617.
[C ₉ H ₁₃ N]	3-CH ₃ C ₆ H ₄ N(CH ₃) ₂ RN 121-72-2	224.5	939.	16.	67.	Est	157.	658.
[C ₉ H ₁₃ N]	4-CH ₃ C ₆ H ₄ N(CH ₃) ₂ RN 99-97-8	225.6	944.	17.	70.	Est	157.	656.
[C ₉ H ₁₄ N ₂ O ₆]	5,6-Dihydrouridine RN 5627-05-4	~208	~870	-233.5	-977.	Est	-76.	-317.
[C ₉ H ₁₅ N]	(CH ₂ =CHCH ₂) ₃ N RN 102-70-5	230.0	962.	53.5	224.	Est	189.	792.
[C ₉ H ₁₇ N]	1-Cyclopentylpyrrolidine RN 18707-33-0	233.1**	975.**					
[C ₉ H ₁₇ N]	c-C ₅ H ₁₀ NCH=C(CH ₃) ₂ RN 673-33-6	230.7**	965.**	-7.	-31.	Est	127.5	534.
[C ₉ H ₁₇ NO ₂]	3,3-Dimethoxy- -1-azabicyclo[2.2.2]octane RN xxxxx	232**	971**	-78.	-326.	Est	56.	233.
[C ₉ H ₁₈ O]	(tert-C ₄ H ₉) ₂ CO RN 815-24-7	206.5	864.	-83.	-345.8	[77PED/RYL]	77.	320.
[C ₉ H ₁₈ N ₂]	1,5-Diazabicyclo[3.3.3]- undecane RN 283-58-9	232.4	972.	33.	138.	[81ALD/ARR]	166.	696.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₉ H ₁₉ N]	2,2,6,6,-Tetramethyl-piperidine RN 768-66-1	231.7**	969.**	-38.	-160. (3)	[81SUR/HAC]	96.	401.
[C ₉ H ₁₉ N]	N-Isobutylpiperidine RN 10315-89-6	232.9**	974.**					
[C ₉ H ₂₁ N]	(t-C ₄ H ₉)C(CH ₃) ₂ N(CH ₃) ₂ RN 3733-36-6	235.1	984.	-34.	-142.	Est	97.	404.
[C ₉ H ₂₁ N]	(n-C ₃ H ₇) ₃ N RN 102-69-2	234.0	979.	-38.	-161.	Est	93.	390.
[C ₉ H ₂₁ N]	(t-C ₅ H ₁₁)(t-C ₄ H ₉)NH RN 58471-09-3	232.5	973.	-46.	-191. (4)	Est	87.5	366.
[C ₁₀ ClH ₁₄ N]	4-ClC ₆ H ₄ N(C ₂ H ₅) ₂ RN 2873-89-4	225.6	944.	2.	8.	Est	142.	594.
[C ₁₀ CrH ₇ O ₃]	(C ₆ H ₅ CH ₂)Cr(CO) ₃ RN 32984-97-7	205**	858**					
[C ₁₀ F ₃ H ₉]	4-CF ₃ C ₆ H ₄ C(CH ₃)CH ₂ RN 55186-75-9	199.6	835.	-131.	-549.	Est	35.	146.
[C ₁₀ FeH ₁₀]	(C ₅ H ₅) ₂ Fe RN 102-54-5	~210	~879.	58.	242. (3)	[77PED/RYL]	213.5	893.
[C ₁₀ H ₈]	Azulene RN 275-51-4	220.	921.	69.	289. (3)	[77PED/RYL]	215.	898.
[C ₁₀ H ₈]	Naphthalene RN 91-20-3	194.7	815.	36.	150. (1)	[*82COL/JIM]	207.	865.
[C ₁₀ H ₉ N]	1-Naphthalenamine RN 134-32-7	216.9	907.5	38.	158. (7)	[77PED/RYL]	186.5	780.5
[C ₁₀ H ₁₀ N ₂]	1,8-Diaminonaphthalene RN 479-27-6	223.8	936.	46.	193.	Est	188.	787.
[C ₁₀ H ₁₀ Ni]	(C ₅ H ₅) ₂ Ni RN 1271-28-9	223.	933.	85.	357. (5)	[77PED/RYL]	228.	954.
[C ₁₀ H ₁₀ Ru]	(C ₅ H ₅) ₂ Ru RN 1287-13-4	218**	912**	-62.	-260.	Est	85.5	358.
[C ₁₀ H ₁₂]	1,2,3,4-tetrahydronaphthalene RN 119-64-2	194.7	815.	6.	24. (2)	[77PED/RYL]	177.	739.5
[C ₁₀ H ₁₂]	4-CH ₃ C ₆ H ₄ C(CH ₃)CH ₂ RN 1195-32-0	211.0	883.	19.	80.	Est	174.	727.
[C ₁₀ H ₁₂ O]	4-CH ₃ OC ₆ H ₄ C(CH ₃)CH ₂ RN 1712-69-2	217.4	910.	-23.	-95.	Est	125.5	525.
[C ₁₀ H ₁₃ N]	N-Phenylpyrrolidine RN 4096-21-3	224.7	940.	30.	117.	Est	171.	716.
[C ₁₀ H ₁₄]	t-C ₄ H ₉ C ₆ H ₅ RN 98-06-6	193.0	807.	-5.	-23. (1)	[77PED/RYL]	167.	699.5
[C ₁₀ H ₁₄]	n-C ₄ H ₉ C ₆ H ₅ RN 104-51-8	192.1	804.	-3.	-13. (1)	[77PED/RYL]	170.	713.
[C ₁₀ H ₁₄ N ₂ O ₅]	Thymidine RN 50-89-5	~208	~870	-230.	-961.	Est	-72.	-301.
[C ₁₀ H ₁₅ N]	C ₆ H ₅ N(C ₂ H ₅) ₂ RN 91-66-7	227.6	952.	9.5	40.	[69BEN/CRU]	148.	617.
[C ₁₀ H ₁₅ N]	3,5-(CH ₃) ₂ C ₆ H ₃ N(CH ₃) ₂ RN 4913-13-7	227.0	950.	8.	35.	Est	147.	615.
[C ₁₀ H ₁₆]	1,5,5-Trimethyl-3-methylenecyclohexene RN 16609-28-2	216.1**	904.**	-2.	-8.	[79AUE/BOW]	148.	618.
[C ₁₀ H ₁₆ N ₂]	1,2-(N(CH ₃) ₂) ₂ C ₆ H ₄ RN 704-01-8	235.2	984.	36.	151.	Est	167.	697.
[C ₁₀ H ₁₇ NO]	cis-3-Amino-2-twistanol RN xxxxx	224.0	937.	-47.	-197.	Est	95.	396.
[C ₁₀ H ₁₇ NO]	trans-3-Amino-2-twistanol RN xxxxx (isomer 1)	221.5	927.	-49.	-205.	Est	95.	398.
[C ₁₀ H ₁₇ NO]	trans-3-Amino-2-twistanol RN xxxxx (isomer 2)	220.0	920.	-49.	-205.	Est	97.	405.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₁₀ H ₁₉ N]	1-Azabicyclo[3.3.3]undecane (Manxine) RN 31023-92-4	230.1	963.	5.	20.(20)	Est	140.	587.
[C ₁₀ H ₁₉ NO]	4-Aminodecahydro-3-naphthalenol RN xxxxx	222.1	929.	-77.	-321.	Est	67.	280.
[C ₁₀ H ₂₀ O ₅]	1,4,7,10,13-Pentaoxacyclopenta- decane (15-Crown-5) RN 33100-27-5	223.6	936.	-186.	-780.(2)	[82BYS/MAN]	-44.	-184.
[C ₁₀ H ₂₂ O]	(n-C ₅ H ₁₁) ₂ O RN 693-65-2	205.2**	859.**	-90.	-375.	Est	71.	296.5
[C ₁₀ H ₂₂ O ₅]	CH ₃ (OCH ₂ CH ₂) ₄ OCH ₃ RN 143-24-8	227.2	951.					
[C ₁₀ H ₂₃ N]	n-(C ₁₀ H ₂₁)NH ₂ RN 2016-57-1	220.7**	923.**	-51.	-214.	Est	94.	393.
[C ₁₀ H ₂₄ N ₂]	(CH ₃) ₂ N(CH ₂) ₆ N(CH ₃) ₂ RN 111-18-2	237.9	995.	-22.	-91.	Est	106.	444.
[C ₁₁ H ₁₀]	1-Methylnaphthalene RN 90-12-0	200.7	840.	27.	113.(2)	[74SAB/CHA]	192.	803.
[C ₁₁ H ₁₀]	2-Methylnaphthalene RN 91-57-6	200.0	837.	26.5	111.(2)	[74SAB/CHA]	192.	804.
[C ₁₁ H ₁₂ N ₂ O ₂]	L-Tryptophan RN 54-12-6	225.4	943.	-58.	-243.	Est	82.	344.
[C ₁₁ H ₁₃ N]	1,4-Dihydro-1,4-ethanoquinoline RN 4363-25-1	232.0	971.	41.	173.	Est	175.	732.
[C ₁₁ H ₁₅ N]	1-Phenylpiperidine RN 4096-2-2	225.8	945.	14.	57.	Est	154.	642.
[C ₁₁ H ₁₇ N]	2,6-Diisopropylpyridine RN 6832-21-9	232.9	974.	-8.	-33.	Est	125.	523.
[C ₁₁ H ₁₇ N]	2-C ₆ H ₁₃ (c-C ₅ H ₄ N) RN 1129-69-7	228.9	958.	0.	0.	Est	137.	572.
[C ₁₁ H ₁₇ N]	3-CH ₃ C ₆ H ₄ N(C ₂ H ₅) ₂ RN 91-67-8	228.9	956.	1.	4.	Est	138.	578.
[C ₁₁ H ₁₇ N]	4-CH ₃ C ₆ H ₄ N(C ₂ H ₅) ₂ RN 613-48-9	228.6	956.	3.	12.	Est	140.	587.
[C ₁₂ H ₈]	Biphenylene RN 259-79-0	203.4	851.	104.	437.(13)	[77PED/RYL]	267.	1116.
[C ₁₂ H ₈ N ₂]	Phenazine RN 92-82-0	223.7	936.	82.	344.(3)	[80ARS]	224.	938.
[C ₁₂ H ₁₀]	Acenaphthene RN 83-32-9	203.5	851.	37.	155.(1)	[81KUD/KUD]	199.	834.
[C ₁₂ H ₁₀]	Biphenyl RN 92-52-4	196.1	820.	43.	182.(1)	[77PED/RYL]	213.	892.
[C ₁₂ H ₁₄ N ₂]	N,N'-Dimethyl-1,8-naphthalene- diamine RN 20734-56-9	230.0	962.	45.	189.	Est	181.	757.
[C ₁₂ H ₁₆ N ₂ O ₆]	2',3'-O-Isopropylideneuridine RN 362-43-6	208	870	-218.	-911.	Est	-60.	-251.
[C ₁₂ H ₁₈]	(CH ₃) ₆ C ₆ RN 87-85-4	207.3	867.	-21.	-87.(3)	[77PED/RYL]	138.	576.
[C ₁₂ H ₁₉ N]	C ₆ H ₄ N(CH ₃) ₂ ,2-t-C ₄ H ₉ RN 22025-87-2	229.3	959.	15.	63.	Est	151.5	634.
[C ₁₂ H ₁₉ N]	C ₆ H ₅ N(C ₃ H ₇) ₂ RN 2217-07-4	228.6	956.	1.	3.	Est	138.	578.
[C ₁₂ H ₂₁ N]	(CH ₂ =C(CH ₃)CH ₂) ₃ N RN xxxxx	230.7**	965.**	28.	116.	Est	163.	684.
[C ₁₂ H ₂₁ NO]	3-Amino-tricyclo[7.3.0.0 ^{4,8}] dodecan-2-ol RN xxxxx	220.0	920.	-63.	-263.	Est	83.	347.
[C ₁₂ H ₂₄ N ₂]	1,6-Diazabicyclo[4.4.4]- tetradecane RN 71058-67-8	226.0	946.	-23.	-95.	[81ALD/ARR]	116.	489.5
[C ₁₂ H ₂₄ O ₆]	1,4,7,10,13,16-Hexaoxa- cyclooctadecane (18-Crown-6) RN 17455-13-9	230.	962.	-227.	-950.	Est	-91.	-382.
[C ₁₂ H ₂₇ N]	(n-C ₄ H ₉) ₃ N RN 102-82-9	234.8	982.	-53.	-222.(1)	Est	78.	326.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₁₃ H ₉ N]	Acridine RN 260-94-6	231.9	970.	70.	291.	(1) [81KUD/KUD2]	203.	851.
[C ₁₃ H ₁₀]	Fluorene RN 86-73-7	200.0	837.	45.	187.	(1) [81KUD/KUD]	210.	880.
[C ₁₃ H ₁₀ O]	(C ₆ H ₅) ₂ CO RN 119-61-9	210.9	882.	12.	50.	(3) [78SAB/LAF2]	167.	698.
[C ₁₃ H ₁₃ P]	(C ₆ H ₅) ₂ (CH ₃)P RN 1486-28-8	230.3	963.5	44.	185.	Est	180.	751.5
[C ₁₃ H ₁₆ N ₂]	N,N,N'-Trimethyl-1,8-naphthalenediamine RN 20723-57-0	235.6	986.	-52.	-217.	Est	78.	327.
[C ₁₃ H ₂₁ N]	2,6-Di-t-butylpyridine RN 585-48-4	233.4	976.	-19.	-81.	Est	113.	472.5
[C ₁₃ H ₂₁ N]	2,4-Di-t-butylpyridine RN 29939-31-9	231.4**	968.**	-19.	-79.	Est	115.	483.
[C ₁₃ H ₂₅ N]	2,6-Di-t-butylpiperidine RN xxxxx	234.3	980.	-76.	-317.	Est	56.	233.
[C ₁₃ H ₂₅ N]	out-6H-1-Azabicyclo[4.4.4]tetradecane RN xxxxx	214.3	896.	-11.	-47.	[81ALD/ARR]	140.	586.
[C ₁₄ H ₁₀]	Anthracene RN 120-12-7	207.0	866.	55.	230.	(1) [79KUD/KUD4]	214.	894.
[C ₁₄ H ₁₀]	Phenanthrene RN 85-01-8	198.7	831.	49.	207.	(1) [79KUD/KUD4]	216.	906.
[C ₁₄ H ₁₂]	(C ₆ H ₅) ₂ C=CH ₂ RN 530-48-3	211.9	887.	59.	246.	(4) [77PED/RYL]	212.5	889.
[C ₁₄ H ₁₄]	C ₆ H ₅ (CH ₂) ₂ C ₆ H ₅ RN 103-29-7	194.6	814.	34.	143.	(2) [77PED/RYL]	205.	859.
[C ₁₄ H ₁₈]	1,2,3,4,5,6,7,8-Octahydro-phenanthrene RN 5325-97-3	204.7	856.	-8.	-34.	(8) [77SHA/GOL]	153.	640.
[C ₁₄ H ₁₈]	1,2,3,4,5,6,7,8-Octahydro-anthracene RN 1079-71-6	202.6	848.	-9.	-37.	(3) [77PED/RYL]	154.	645.
[C ₁₄ H ₁₈ N ₂]	N,N,N',N'-Tetramethyl-1,8-naphthalenediamine RN 20734-58-1	241.8	1012.	63.	262.	Est	186.5	780.
[C ₁₄ H ₂₇ N]	1-Methyl-2,6-t-butylpiperidine RN xxxxx	239.2	1001.	-67.	-311.	Est	60.	250.
[C ₁₅ H ₁₂]	9-Methylanthracene RN 779-02-2	213.9	895.	48.	201.	Est	200.	836.
[C ₁₅ H ₁₂]	2-Methylanthracene RN 613-12-7	210.3	880.	45.	187.	Est	200.	837.
[C ₁₅ H ₁₈]	1,4-Dimethyl-7-isopropylazulene RN 489-84-9	233.	975.	33.	139.	Est	165.	694.
[C ₁₆ H ₁₀]	Pyrene RN 129-00-0	206.1	862.	52.	216.	(1) [79KUD/KUD2]	211.	884.
[C ₁₆ H ₁₀]	Fluoranthene RN 206-44-0	199.3	834.	69.	289.	(1) [81KIM/KIM]	235.	985.
[C ₁₆ H ₁₆]	(4-CH ₃ C ₆ H ₄) ₂ C=CH ₂ RN xxxxx	215.4	901.	43.	180.	Est	193.	809.
[C ₁₆ H ₁₈]	C ₆ H ₅ (CH ₂) ₄ C ₆ H ₅ RN 1083-56-3	195.9	820.	24.	100.	Est	194.	810.
[C ₁₈ H ₁₂]	Tetracene RN 92-24-0	217.8	911.	67.	284.	(1) [79KUD/KUD2]	216.	903.
[C ₁₈ H ₁₂]	Chrysene RN 218-01-9	201.6	843.	63.	263.	(1) [79KUD/KUD2]	227.	949.5
[C ₁₈ H ₁₂]	Triphenylene RN 217-59-4	198.5	830.5	64.5	270.	(1) [79KUD/KUD2]	232.	969.5
[C ₁₈ H ₁₅ P]	(C ₆ H ₅) ₃ P RN 603-35-0	~230.	~962.	78.	328.	(21) [79STE]	214.	896.
[C ₂₀ H ₁₂]	Perylene RN 198-55-0	211.4	884.	74.	308.	(4) [77PED/RYL]	228.	953.5

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species—Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[C ₂₂ H ₁₂]	1,12-Benzoperylene RN 191-24-2	208.5	872.	72.	302.	[77STE/GOL]	229.	960.
[C ₂₂ H ₁₄]	Picene RN 213-46-7	203.4	851.	78.	326.	Est	240.	1005.
[C ₂₄ H ₁₂]	Coronene RN 191-07-1	205.0	858.	77.	323.	[77STE/GOL]	238.	995.
[Cl]	Cl RN 22537-15-1	123.6	517.	29.1	122.	[82/TN270]	271.8	1137.
[ClH]	HCl RN 7647-01-0	134.8	564.	-22.	-92.	[82/TN270]	209.	874.
[F]	F RN 14762-94-8	81.0	339.	19.	79.	[82/TN270]	>303.3	1270.
[FH]	HF RN 7664-39-3	117.	489.5	-65.	-271.	[82/TN270]	184.	770.
[F ₂ O ₂ S]	F ₂ SO ₂ RN 2699-79-8	159.0	665.	-181.	-759. (8)	[82JANAF]	25.	106.
[F ₃ N]	NF ₃ RN 7783-54-2	144	604	-30.	-125.	[82/TN270]	192.	802.5
[F ₃ OP]	OPF ₃ RN 13478-20-1	167.8	702.	-289.	-1211.	[82/TN270]	-91.5	-383.
[F ₃ P]	PF ₃ RN 7783-55-3	166.5	697.	-220.	-919.	[82/TN270]	-20.	-85.5
[HI]	HI RN 10034-85-2	150.	628.	6.	26.	[82/TN270]	222.	928.
[HNO ₃]	HNO ₃ See References to Table 1: 75FEH/HOW							
[HO ₂]	HO ₂ RN 3170-83-0	~158.	~661.	3.	11. (4)	[82BAU/COX]	210.	880.
[H ₂]	H ₂ RN 1333-74-0	101.3	424.	0.	0.	DEF	264.	1106.
[H ₂ O]	H ₂ O RN 7732-18-5	166.5	697.	-58.	-242.	[82/TN270]	141.	591.
[H ₂ O ₂]	H ₂ O ₂ RN 7722-84-1	162.	678.	-32.5	-136.	[82BAU/COX]	171.	716.
[H ₂ O ₄ S]	H ₂ SO ₄ RN 7664-93-9	~169	~707	-176.	-735. (8)	[82JANAF]	21.	88.
[H ₂ S]	H ₂ S RN 7783-06-4	170.2	712.	-5.	-21.	[82/TN270]	190.	797.
[H ₂ Se]	H ₂ Se RN 7783-07-5	171.3	717.	7.	30.	[82/TN270]	201.5	843.
[H ₂ N]	NH ₂ RN 15194-15-7	187	782	44.	185. (5)	[82MCM/GOL]	223.	935.
[H ₃ N]	NH ₃ RN 7664-41-7	204.0	853.5	-11.	-46.	[82/TN270]	151.	630.5
[H ₃ P]	PH ₃ RN 7803-51-2	188.6	789.	1.	5.	[82/TN270]	178.	746.
[H ₄ N ₂]	H ₂ NNH ₂ RN 302-01-2	204.7	856.	23.	95.	[82/TN270]	184.	770.
[H ₄ Si]	SiH ₄ RN 7803-62-5	~155	~648	8.	35.	[81BEL/PER]	219.	916.5
[He]	He RN 7440-59-7	42.5	178.	0.	0.	DEF	323.	1352.
[I]	I RN 14362-44-8	145.4	608.	25.5	107.	[82/TN270]	246.	1029.
[Kr]	Kr RN 7439-90-9	101.6	425.	0.	0.	DEF	264.	1105.
[Mg]	Mg See References to Table 1: 77PO/POR							
[Mg ₂]	Mg ₂ RN 29904-79-8	~219	916.					
[NO]	NO RN 10102-43-9	~127	~531	21.5	90.	[82BAU/COX]	260.	1089.
[N ₂]	N ₂ RN 7727-37-9	118.2	494.5	0.	0.	DEF	247.5	1035.5
[N ₂ O]	N ₂ O RN 10024-97-2	136.5	571.	20.	82.	[82/TN270]	249.	1041.
[Ne]	Ne RN 7440-01-9	48.1	201.	0.	0.	DEF	318.	1329.

Table 2. Proton affinities and heats of formation of molecules and corresponding protonated species--Continued

Formula	Compound (M)	Proton Affinity		$\Delta_f H(M)$		Reference	$\Delta_f H(MH^+)$	
		kcal/mol	kJ/mol	kcal/mol	kJ/mol		kcal/mol	kJ/mol
[O]	O RN 17778-80-2	116.3	487.	59.5	249.	[82/TN270]	309.	1293.
[OSi]	SiO See References to Table 1: 81FAH/FEH							
[O ₂]	O ₂ RN 7782-44-7	100.9	422.	0.	0.	DEF	265.	1108.
[O ₂ S]	SO ₂ RN 7446-09-5	161.6	676.	-71.	-297.	[82/TN270]	133.	557.
[O ₃ S]	SO ₃ RN 7446-11-9	138	577	-95.	-396.	[82/TN270]	133.	557.
[S]	S RN 7704-34-9	158.3	662.	67.	279.	[82/TN270]	274.	1147.
[Xe]	Xe RN 7440-63-3	118.6	496.	0.	0.	DEF	247.	1034.
[Zn]	Zn RN 7440-66-6	156	653	31.	131.	[82/TN270]	241.	1008.

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A reference of the form *00ABC/DEF means that a condensed phase heat of formation from 77PED/RYL has been used with a heat of vaporization or sublimation from the designated reference.

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