A unified view to Brønsted acidity scales: do we need solvated protons?†

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The most comprehensive solvent acidity scale spanning 28 orders of magnitude of acidity was measured in the low-polarity solvent 1,2-dichloroethane (DCE). Its experimental core is linked to the unified acidity scale (pHabs) in an unprecedented and generalized approach only based on experimental values. This enables future measurements of acid strengths and acidity adjustments in low polarity solvents. The scale was cross-validated computationally. The purely experimental and computational data agree very well. The DCE scale includes 87 buffer systems with pHabs values between −13.0 and +15.4, i.e. similar to water at hypothetical and extreme pH values of −13.0 to +15.4. Unusually, such high acidities in DCE are not realized via solvated protons, but rather through strongly acidic molecules able to directly donate their proton, even to weak bases dissolved in the solution. Thus, in all examined cases, not a single solvated proton is present in one liter of DCE.

Introduction

The proton is undoubtedly one of the most important particles in the universe. The most prominent phenomenon associated with it – acidity – is so important to nature that living organisms even have a sense for it. Acid-base reactions are ubiquitous, but are mainly investigated in polar and/or protic solvents containing ionizable protons at concentrations that allow for a direct measurement, e.g. by conductivity. Similarly, superacid chemistry is mainly done in highly polar, protic media like HF/SbF5 mixtures or the legendary “magic acid” HSO3F/SbF5. This overall polarity of a solvent medium physically arises from the hydrogen bond donor and acceptor capacity, polarizability and dipole moment of its molecules and the dipole density that allow for stabilizing interactions with the dissolved ions, thus lowering their chemical potential (partial molar free energy, i.e. rate of change of the free energy of the system with respect to the change in the number of molecules of the respective species that are added to the system). If the medium is non-polar, the solvation of ions, including H⁺, is weak and their chemical potential is high. Thus, paradoxically, the highest acidities should be achievable in an environment that is as non-polar as possible. In agreement with this, numerous e.g. acid-catalyzed – chemical reactions are carried out both in the lab as well as on an industrial scale in non- to weakly polar aprotic solvents with relative dielectric constants εr roughly between 2 and 14 (such as alkanes, substituted arenes, ethers or haloalkanes). Azeotropic dehydration in non-polar toluene with toluene-sulfonic acid present a popular example. Thus, acid chemistry in non-polar and non-basic solvents has been used in practice for decades. However, wouldn’t it be fundamentally interesting to learn about the medium acidity of this and other popular mixtures in quantitative terms, in order to rigorously compare reaction conditions in one solvent to another?

Towards this general goal of the thermodynamically sound evaluation of acidity, we have now derived a general approach to establish unified acidities in solvent media only on the basis of experimental values. This fundamental development is applied here to the exemplarily selected low-polarity solvent 1,2-dichloroethane (DCE, εr = 10.36 (ref. 7)). Due to its low basicity, but sufficient polarity to dissolve polar and ionic compounds at measurable concentrations/activities, DCE is a suitable solvent for studying acids and also superacids. The scale covers 28 orders of magnitude of acidity and is linked to the unified acidity scale (pHabs scale). It is based on our earlier work on the high acidities in DCE (stronger acids than picric acid, spanning about 15 orders of magnitude). Finally, we support the experimental results by a cross-validation based on quantum-chemical calculations with consideration of solvation effects by the SMD, rCCC, and COSMO-RS models.

Definitions and concepts

Acidity in non-aqueous media, and especially the comparison of acidity across media or even phase boundaries, is not common...
knowledge and quantitative views remain sparse. Therefore, we start with essential definitions and concepts.

**pH and the chemical potential of the proton**

pH is the most popular measure for medium acidity. In its standard definition, pH$_s$ = $-\log[a(H^+, S)]$, it is a measure for the solvated proton’s chemical potential $\mu_{abs}(H^+, S)$ in the solvent/medium S. The molecular acidity of an acid HA in a solvent S is described by the equilibrium (1),

$$HA_{(solv.)} \leftrightarrow A_{(solv.)}^- + H_{(solv.)}^+ \hspace{1cm} (1)$$

and the available activity $a(H^+, S)$ of solvated protons $H_{(solv.)}^+$ formally determines the equilibrium pH$_s$ value in S, i.e. its acidity in medium S (and limited to medium S). Decreasing the pH of a solution by one unit increases the proton chemical potential and thus the solution’s acidity by $RT \ln 10 = 5.71$ kJ mol$^{-1}$ at standard conditions ($25^\circ$C, 1 bar). The intrinsic molecular acidity of HA in S is quantitatively given by its acidity constant $K_{a,S}$ (eqn (1)), used as the negative decadic logarithm $pK_{a,S}$:

$$K_{a,S} = \frac{a(H^+, S) \cdot a(A^- , S)}{a(HA, S)} \hspace{1cm} (2)$$

Importantly, the pH$_s$ values as defined above – using reference states in medium S – are bound to the medium S and cannot be used to compare acidities in different solvents/media.

**Unified acidity scale**

Following the seminal work of Bartmess,$^4$ we introduced the ideal proton gas at one bar as a unified – and medium-independent – reference state for acidity and as a thermodynamic zero point.$^3$ The chemical potential of this reference state “proton gas” is set to $\mu_{abs}(H^+, g) = 0$ kJ mol$^{-1}$ and pH$_{abs} = 0$. The independence of this reference state from any medium allows for a unified comparison of acidities in terms of chemical potentials or the corresponding pH$_{abs}$ values in different media on an absolute basis. For example, water with a pH$_{water}$ of 0 has a pH$_{abs}$ of 193.5 (with the published Gibbs hydration energy of the gaseous proton ($\Delta_{solv}^0 G^\circ(H^+, H_2O) = -1105$ kJ mol$^{-1}$)$^{14-16}$). In order to express pH$_{abs}$ in a more familiar way, it is useful to shift it by 193.5 units, to obtain so-called pH$_{abs}^{193.5}$ values (pH$_{abs}^{193.5} = pH_{abs} - 193.5$)$^{17}$ which are a direct continuation of aqueous pH values and identical to them in water. This means that the chemical potential of the proton in any solvent/medium with pH$_{abs}^{193.5}$ 5 is the same as pH$_{water}$ 5.

**Acidity measurements in low polarity media**

Acidity measurements in low polarity media become increasingly difficult and only limited examples are known.$^a$ An acidity scale was even measured in the very non-polar n-heptane,$^{18}$ but it is obvious that protons solvated by heptane itself are not present in solution, as protonated n-heptane would decompose with the loss of H$_2$ or an alkane and rearrange instantly to a more stable branched tertiary carbenium ion. With the commonly used notion that acidity relies on dissolved ions, it needs acids with a pK$_{a,S}$ in a given solvent of less than 12 to obtain micro-molar ion concentrations that can be easily measured. In low polarity media, the choice of such acids is very limited. Only extreme acids are able to generate ions in low polarity media to a measurable/isolable extent. A drastic example is H[HCBr$_3$F$_{11}$] that even protonates the very non-polar medium liquid carbon dioxide giving the salt H(CO$_2$)$_2$[HCBr$_3$F$_{11}$]$.^{19}$ Since low polarity solvents weakly solvate ions, the extent of ion–ion interactions and aggregation increases drastically in such media. Therefore, effects like ion-pairing and others have to be evaluated and correction schemes have to be applied to obtain the “real acidity” data (cf. ESI Sections 2 and 3†). In addition, traces of basic impurities like water compete with the solvent base for the proton and therefore have a dramatic effect on the acid dissociation, e.g. by formation of (also further solvated) H$_2$O$^+$. This effect is evaluated in a later section.

**Superacidity**

Superacidity refers to the highest acidity and can be used in relation to media or molecules. Thus, we differentiate between two principal types of superacidity:

**Medium superacidity.** Medium superacidity was initially defined by Gillespie as the acidity of a medium that is “stronger than sulfuric acid”.$^{20}$ In the context of the unified acidity scale, a superacidic medium is one where the chemical potential of the proton lies higher than that in 100% sulfuric acid. Although the Hammett$^{21}$ function $H_0$ is most frequently used to assess superacidity,$^{22}$ it does not directly express the proton’s chemical potential and thus should not be considered a thermodynamic acidity scale (see ref. 5 and 10).$^\S$ In the unified acidity scale, the absolute chemical potential of the proton in neutral liquid H$_2$SO$_4$ was assessed as the threshold for superacidity: if a medium S has $\mu_{abs}(H^+, S) < \mu_{abs}(H^+, H_2SO_4 (l))$ or $-975$ kJ mol$^{-1}$, it is superacidic.$^5,19$ This corresponds to a pH$_{abs}^{193.5}$ below $-22.4$. $^\|$ 

**Molecular superacidity.** Molecular superacidity refers to an acidic molecule/molecular ion, which, in a given solvent/medium S, is a stronger acid than sulfuric acid, i.e. it has a lower pK$_{a,S}$ value in this solvent/medium than sulfuric acid. In the gas phase,$^{23}$ a superacidic molecule or molecular ion has a lower gas phase acidity (GA) value than the gaseous H$_2$SO$_4$ molecule.

The equilibrium acidity scale in DCE

Buffer solutions are a superior means to measure the self-consistent equilibrium acidity scale in DCE, both for reasons of the measurable concentrations of ions as well as the stability of the photochemical potential. The buffers used herein, i.e. mixtures of acids and bases, keep the acidity at an approximately constant value and allow for the reliable and reproducible measurement of the chemical potential. In the simplest case, a 1 : 1 mixture of an acid with a salt of its conjugate base buffers at the buffer point pH$_S = pK_{a,S}$. To construct a self-consistent ladder of relative acidities, protonation equilibria between a large number of acid pairs (including all acids on the scale) have to be evaluated$^{24}$ in the solvent S according to equilibrium (3),
HA₁(solv.) + A²⁻(solv.) ⇌ A⁻₁(solv.) + HA₂(solv.) \hspace{1cm} (3)

where HA₁ and HA₂ are the two acids. Their relative acidity difference \( \Delta pK_{a,S} \) in solvent S can therefore be calculated as follows (4):

\[
\log K = \Delta pK_{a,S} = pK_{a,S}(HA_1) - pK_{a,S}(HA_2)
= \log \frac{a(A_1^-, S) \cdot a(HA_2, S)}{a(HA_1, S) \cdot a(A_2^-, S)} \hspace{1cm} (4)
\]

However, the direct measurement in DCE gives values slightly differing from \( \Delta pK_{a,S} \) (eqn (4)) due to ion-pairing effects. Those are denoted as \( \Delta pK_p \) (\( p \) stands for ion-pairing). In this work, the acidity scale\(^a\) in DCE is presented, composed of a total of 226 \( \Delta pK_p \) measurements between 87 acids. For the construction of the scale, the sum of the squares of the differences between the experimental \( \Delta pK_p \) values and the differences between the assigned \( pK_{ip,rel} \) values was minimized. Taking picric acid as a reference acid by arbitrarily assigning a value of 0.0 to its \( pK_{ip,rel} \) value gave a self-consistent \( pK_{ip,rel} \) ladder (Table 2).\(^b\) It should be noted that in principle every acid could be taken as the reference acid without affecting the accuracy of the procedure. Using picric acid is rather traditional (it is also used as the reference acid in acetonitrile\(^a\)). To obtain the \( \Delta pK_{a,DCE} \) values, all \( pK_{ip,rel} \) values were corrected (by up to 0.51 units, see ESI, Section 3\(^†\)) for the logarithmic difference of the ion-pair dissociation constants \( \Delta pK_d \) of the acids under study using the Fuoss model as described in the methods section and the ESI (Section 2\(^†\)).\(^d\) Thus, one obtains the corrected thermodynamic \( pK_{a,DCE} \) values with respect to picric acid (Table 2). The compiled self-consistent acidity scale in Table 2 spans 28.4 orders of magnitude. The overall reliability of the logarithmic acidity constant values was checked using the consistency standard deviation.\(^e\) For both \( pK_{ip,rel} \) and \( pK_{a,rel} \) values, it is 0.04 log units or 0.23 kJ mol\(^{-1}\), which can be considered very good for a low polarity solvent.

### Anchoring the DCE scale to pH\(_{H_2O}^{abs}\) values

The construction of a \( pH_{abs}^{H_2O} \) scale hitherto required in all cases the Gibbs solvation energy for at least one ion, for example the hydration energetics of the gaseous proton \( (\Delta_{olv}G(\text{H}^+, \text{H}_2\text{O}) = -1105 \text{ kJ mol}^{-1}) \).\(^{13-16}\) This value is not directly accessible by experimental methods, and was obtained by the extrapolation of the hydration thermodynamics of gaseous ion–water clusters to the bulk solvent. According to the authors, it is tainted with an error bar of at least ±8 kJ mol\(^{-1}\) or ±1.4 pH units! Advantageously, in the \( pH_{abs}^{H_2O} \) scale proposed herein, large single ion Gibbs solvation energies can be removed or replaced by single ion Gibbs transfer energies, which are typically lower by 1–2 orders of magnitude. This reduces the overall uncertainty in the chemical potential assessment. Still, for partitioning the chemical potential of dissolved ionic compounds into single ion values, so-called extrathermodynamic assumptions have to be introduced. The most used assumption by far is the so-called “TATB assumption”.\(^f\) It assumes that when transferring the salt tetraphenylarsonium-tetraphenylborate (TATB) from one medium to the other, the chemical potential difference of both ions is about the same between the solvents and the Gibbs transfer energy of each ion just amounts to half of the measurable sum.\(^g\) Although this assumption is disputed,\(^h\) any error or difference between these reference ions will systematically add to the single ion quantities, so that the overall structure of the \( pH_{abs}^{H_2O} \) scale will remain intact.

### General approach

Neither the medium \( pK_{a,DCE} \) of acids nor \( pH_{DCE} \) values are experimentally accessible (see below). Despite this, we can construct a thermodynamically consistent \( pH_{abs}^{H_2O} \) scale using single ion Gibbs transfer energies from water to DCE via suitable Born–Fajans–Haber cycles (BFHCs, see ESI Section 6 for details\(^†\)). The only acid in Table 2 for which sufficient experimental data for such a connection to aqueous pH exists is HCl. Fortunately, the Gibbs solvation energies of single ions cancel out in this calculation or can be replaced by one single ion Gibbs transfer energy, namely that of the chloride ion. According to these calculations, detailed in the ESI in Section 5,\(^†\) a 1 : 1 HCl/Cl\(^-\) buffer mixture in DCE has an acidity that corresponds to pH 2.5 in water. With the knowledge that HCl has a \( pK_{a,rel} \) of 0.2 vs. picric acid (Table 2), we obtain the simply applicable universal formula (5) that is valid for all our measured acids:

\[
pH_{abs}^{H_2O} = 2.3 + pK_{a,rel} + \log \frac{a(A^-, \text{DCE})}{a(HA, \text{DCE})} \hspace{1cm} (5)
\]

Using formula (5), we obtain a measured \( pH_{abs}^{H_2O} \) range in DCE between +15.4 for the weakest acid 9-COOMe-fluorene and −13.0 for the strongest acid CN-TCNP in 1 : 1 (neutral acid : acid anion) buffer solutions. Such 1 : 1 buffer solutions correspond to the so-called buffer point (BP) and the corresponding \( pH_{abs}^{H_2O} \) can be termed as \( pH_{abs}^{BP} \) and calculated via eqn (6).

\[
pH_{abs}^{H_2O}(\text{BP}) = 2.3 + pK_{a,rel} \hspace{1cm} (6)
\]

The \( pH_{abs}^{H_2O}(\text{BP}) \) values for all of the acids are presented in Table 2. Using the acids from Table 2, DCE solutions of well-defined unified acidity, i.e. \( pH_{abs}^{BP} \) values, can be prepared on the basis of our data together with eqn (5) and (6) by any first-year chemist. For higher ion concentrations, a correction scheme for non-ideality (Debye–Hückel effects) is deposited in the ESI, Section 7.\(^†\)

### Cross validation: quantum-chemical solvation models vs. experimental data

Several terms sum up to the overall Gibbs solvation energy as calculated by quantum chemistry. For small ions and molecules, the electrostatic part is very sensitive to the chosen calotte radii or isodensity surface and dominates the overall solvation thermodynamics. For large ions and molecules, the van-der-Waals interaction as well as the cavity energy becomes increasingly important. Thus, here we decided to use the
medium sized HNTf$_2$/NTf$_2^-$ system for model building. An evaluation of the favorable performance of this system by comparing calculated (rCCC model) and experimental pK$_a$s of HNTf$_2$ in MeCN or DMSO was published. For validation, the pK$_{a,rel}$ (HNTf$_2$, DCE) of $-12.0$ from Table 2 was set into eqn (6). We obtain $p_{H^+}^{H_2O}(BP) = 2.3 + (-12.0) = -9.7$ for HNTf$_2$ in DCE at the buffer point BP (i.e. $pH_2 = pK_{a,5}$). As described above, the main error source in this value is the uncertainty of $\Delta_a$G$^\circ$ (Cl$^-$, DCE $\rightarrow$ H$_2$O), obtained with the TATB assumption. Eqn (7)

$$p_{H^+}^{H_2O}(BP) = \Delta_{solv} G^\circ (\text{HNTf}_2, \text{DCE}) + \Delta_{solv} G^\circ (\text{NTf}_2^-, \text{DCE}) + GA(\text{HNTf}_2) + \Delta_{solv} G^\circ (H^+, H_2O)$$

(7)

can be used for a cross validation with the solvation models. For this, we augmented our published calculated rCCC$^{19}$ and SMD$^{20}$ data set with COSMO-RS$^{11,12}$ solvation calculations (see ESI Section 14 for details$^\ddagger$). By combining the experimental $\Delta_{solv} G^\circ$ (H$^+$, H$_2$O) ($-1105$ kJ mol$^{-1}$) and the experimental gas phase acidity ($1199$ kJ mol$^{-1}$)$^\dagger$ of HNTf$_2$ with the quantum-chemically calculated Gibbs solvation energies of HNTf$_2$ and NTf$_2^-$, we obtained the validation data in Table 1.

Pleasingly and supporting our experimental findings, the calculated $p_{H^+}^{H_2O}(BP)$ values between $-6.7$ and $-10.7$ collected in Table 1, and obtained with the different quantum-chemical solvation models, agree within $-1.0/3.0$ pH units to the experimental one at $-9.7$. This corresponds to $-5.7/17.1$ kJ mol$^{-1}$ at standard conditions. It should be noted that, in addition to the error in the calculated Gibbs solvation energies, the $\Delta_{solv} G^\circ$ (H$^+$, H$_2$O) value has an estimated error bar$^{14,15}$ of $\pm 8$ kJ mol$^{-1}$ or $\pm 1.4$ pH units. In the end, it cannot be decided whether the experimental or one of the quantum chemically calculated $p_{H^+}^{H_2O}(BP)$ values is more accurate.

**Does one need solvated protons for protonation of bases?**

One may be surprised that neither our experimental $p_{H^+}^{H_2O}$ values (Tables 1 and 2) nor our $p_{H^+}^{H_2O}$ calculations based on quantum-chemical solvation models (Table 1) require knowledge of the medium $pH_{DCE}$ or any information about the solution thermodynamics of the solvated proton in DCE. The reason is that the proton bound in neutral HNTf$_2$ is in equilibrium with the solvated proton in DCE. Thus, in chemical equilibrium, the protonated by the neutral, non-dissociated acid present – here HNTf$_2$(DCE) – and not by the hardly existing solvated protons!

Is it possible to obtain true medium pK$_a$ values in DCE?

In order to transfer the obtained relative pK$_{a,rel}$ values, with respect to picric acid, to medium pK$_{a,DCE}$ values, we need the reliable medium pK$_{a,DCE}$ of at least one of the acids we measured, as an anchor point. Our calculation of the pK$_{a,DCE}$ value of HNTf$_2$ in DCE gave rather robust values all rounding to $33$, irrespective of the solvation model used (range: $32.6$–$33.1$, Table 1). Even taking the lowest pK$_{a,DCE}$ of $32.6$, this would imply that in a 0.01 M solution of HNTf$_2$ in pure DCE only $5 \times 10^{-18}$ mol acid per liter is dissociated. This is unmeasurable! An additional problem arises from the low basicity of DCE: tiny traces of more basic impurities, especially water, have an enormous influence on acid dissociation and make medium pK$_{a,DCE}$ determinations (nearly) impossible. Even well dried DCE (around 0.15 ppm) still contains around $10^{-2}$ mol L$^{-1}$ of water. A rough SMD calculation gave a pK$_{a,DCE}$ value of $13.6$ for the H$_2$O$^+$ cation in DCE (see ESI Sections 7 and 13 for details$^\ddagger$). According to the law of mass action this means that even at a $2.5 \times 10^{-14}$ mol L$^{-1}$ (!) concentration of water, 50% of all solvated protons would be attached to water molecules and not to DCE. Even in buffered solutions, we see no chance for directly measuring pK$_{a,DCE}$ or pH$_{DCE}$ values in this case. Therefore, and in order to obtain estimates of true medium pK$_{a,DCE}$ values (eqn (2)), the scale was reconstructed with the corrected pK$_{a,rel}$ values and the computationally assessed medium pK$_{a,DCE}$ value of HNTf$_2$. Since all values in Table 1 round to $33$, we used this value as an anchor point (entry 72 in Table 2).

**The DCE acidity scale: explanation and discussion of the entries in Table 2**

The columns pK$_{ip,rel}$, pK$_{a,DCE}$ and $p_{H^+}^{H_2O}$(BP) of Table 2 illustrate the different levels of complexity in data treatment and the corresponding depth of information that can be obtained. In short:

- pK$_{ip,rel}$ – accurately measured experimental and strictly ion-pair (molecular) acidities of compounds relative to picric acid in DCE. The relative values are reliable, but their absolute
Table 2  Acidity scale in 1,2-dichloroethane; see the text for in-depth explanations of the data

| No | Acid                                      | $pK_{A,1}$ | $pK_{A,2}$ | $pK_{A,3}$ | $pH_{1/2}^{\text{DCE}}$ (BP) | Directly measured $\Delta pK_{A}$ values in DCE |
|----|-------------------------------------------|------------|------------|------------|-----------------------------|----------------------------------|
| 1  | 9-COO- Me-fluorene                        | 13.2       | 13.1       | 58.1       | 15.4                        | 0.24                             |
| 2  | (4- Me- C$_6$H$_5$)CHCN                   | 12.9       | 12.8       | 57.8       | 15.1                        | 0.03                             |
| 3  | (4- Me- C$_6$H$_5$)(C$_6$H$_5$)CHCN       | 11.9       | 11.7       | 56.7       | 14.0                        | 0.03                             |
| 4  | 9-CN-fluorene                             | 11.0       | 10.9       | 55.9       | 13.2                        | 0.02                             |
| 5  | (4-H- C$_6$H$_5$)(C$_6$H$_5$)CHCN         | 10.8       | 10.7       | 55.7       | 13.0                        | 0.02                             |
| 6  | (3-CF$_3$- C$_6$H$_5$)C$_6$F$_5$CBCN      | 9.9        | 9.7        | 54.7       | 12.0                        | 0.04                             |
| 7  | (2-C$_6$F$_5$)(C$_6$H$_5$)C$_6$F$_5$CBCN  | 9.5        | 9.4        | 54.4       | 11.7                        | 0.02                             |
| 8  | 9-(C$_6$F$_5$-octafluoro)fluorene         | 9.0        | 8.8        | 53.8       | 11.1                        | 0.02                             |
| 9  | (C$_6$F$_5$)CHCN                          | 8.6        | 8.4        | 53.4       | 10.7                        | 0.02                             |
| 10 | (4-CF$_3$- C$_6$H$_5$)(C$_6$H$_5$)CHCN   | 7.5        | 7.3        | 52.3       | 9.6                         | 0.04                             |
| 11 | (C$_6$F$_5$)CHCN                          | 7.5        | 7.5        | 52.5       | 9.8                         | 0.02                             |
| 12 | (3-CF$_3$- C$_6$H$_5$)CHCH(C$_6$F$_5$)    | 7.3        | 7.2        | 52.2       | 9.5                         | 0.02                             |
| 13 | (2-C$_6$F$_5$)CH(C$_6$F$_5$)COOB         | 7.1        | 7.0        | 52.0       | 9.3                         | 0.02                             |
| 14 | (4-CF$_3$- C$_6$H$_5$)C$_6$F$_5$COOB      | 5.7        | 5.6        | 56.6       | 7.9                         | 0.02                             |
| 15 | (4-NC$_3$)(C$_6$F$_5$)CHCN               | 5.7        | 5.5        | 50.5       | 7.8                         | 0.02                             |
| 16 | (4-NC$_3$)(C$_6$F$_5$)CHCN               | 5.7        | 5.5        | 50.5       | 7.8                         | 0.02                             |
| 17 | 3-CF$_3$- C$_6$H$_5$C$_6$F$_5$CN$_2$     | 4.0        | 4.0        | 49.0       | 6.3                         | 0.02                             |
| 18 | (C$_6$F$_5$)C$_6$H$_5$COOB               | 3.0        | 2.8        | 47.8       | 5.1                         | 0.02                             |
| 19 | 4-Cl-C$_6$F$_5$CHCN                      | 2.5        | 2.5        | 47.5       | 4.8                         | 0.02                             |
| 20 | (4-NC$_3$)(C$_6$F$_5$)CHCN               | 2.4        | 2.3        | 47.3       | 4.6                         | 0.02                             |
| 21 | 2-C$_6$F$_5$CH(C$_6$F$_5$)CN$_2$         | 1.4        | 1.3        | 46.5       | 3.6                         | 0.02                             |
| 22 | Bromothymol blue                          | 1.4        | 1.1        | 46.1       | 3.4                         | 0.02                             |
| 23 | Brucine red                               | 0.5        | 0.2        | 45.2       | 2.5                         | 0.02                             |
| 24 | Picric acid                               | 0.0        | 0.0        | 45.0       | 2.3                         | 0.02                             |
| 25 | HCl                                       | -0.4       | 0.2        | 45.2       | 2.5                         | 0.02                             |
| 26 | 2,3,4,6-(CF$_3$)$_4$C$_6$CH(C$_6$F$_5$)CN$_2$ | -0.6  | -0.6       | 44.9       | 1.6                         | 0.02                             |
| 27 | 2,6-(CF$_3$)$_2$C$_6$H$_5$CH(C$_6$F$_5$)CN$_2$ | -0.6  | -0.6       | 44.2       | 1.5                         | 0.02                             |
| 28 | 2,3,4,6-(CF$_3$)$_4$C$_6$H$_5$CH(C$_6$F$_5$)CN$_2$ | -0.6  | -0.6       | 44.2       | 1.5                         | 0.02                             |
| 29 | 2,6-(CF$_3$)$_2$C$_6$H$_5$CH(C$_6$F$_5$)CN$_2$ | -0.6  | -0.6       | 44.1       | 1.4                         | 0.02                             |
| 30 | 4-CF$_3$- C$_6$H$_5$SO$_2$NH$_2$          | -1.5       | -1.7       | 43.3       | 0.6                         | 0.02                             |
| 31 | 2-CF$_3$- C$_6$H$_5$SO$_2$NH$_2$          | -1.7       | -1.4       | 43.6       | 0.9                         | 0.02                             |
| 32 | 4-CF$_3$- C$_6$H$_5$SO$_2$NH$_2$          | -1.7       | -1.4       | 43.6       | 0.9                         | 0.02                             |
| 33 | 2-CF$_3$- C$_6$H$_5$SO$_2$NH$_2$          | -1.7       | -1.4       | 43.6       | 0.9                         | 0.02                             |
| 34 | 4-CF$_3$- C$_6$H$_5$SO$_2$NH$_2$          | -1.7       | -1.4       | 43.6       | 0.9                         | 0.02                             |
| 35 | 3-NO$_2$-4-CF$_3$-C$_6$H$_5$SO$_2$NH$_2$  | -4.1       | -4.3       | 40.7       | -2.0                        | 0.02                             |
| 36 | 3-NO$_2$-4-CF$_3$-C$_6$H$_5$SO$_2$NH$_2$  | -4.5       | -4.6       | 40.4       | -2.3                        | 0.02                             |
| 37 | HBr                                       | -4.9       | -4.4       | 40.6       | -2.1                        | 0.02                             |
| 38 | 4-NO$_2$- C$_6$H$_5$SO$_2$C$_6$F$_5$CN$_2$ | -5.1       | -5.1       | 39.9       | -2.8                        | 0.02                             |
| 39 | 2,4,6-(SO$_3$F)$_2$phenol                 | -5.9       | -6.0       | 39.0       | -3.7                        | 0.02                             |
| 40 | 2,4,6-H$_2$phenol                         | -6.4       | -6.6       | 38.4       | -4.3                        | 0.02                             |
magnitudes are arbitrary. The relative values are robust within the medium, as are approximations used in other nonpolar media, and depend less on impurities than the absolute values and are thus useful for comparing acids, e.g. to rationalize synthesis conditions or design electrochemical cells within DCE.

Table 2 (Contd.)

| No. | Acid | $pK_{a,rel}$ | pH $H_2O$ | Directly measured $\Delta pK_{a,p}$ values in DCE |
|-----|------|-------------|-----------|---------------------------------------------|
| 41  | CH(CN)$_2$ | 6.6 | -4.4 | 38.6 | -4.1 |
| 42  | 4-C$_6$H$_4$SO$_2$NH$_2$ | 6.9 | -7.0 | 38.0 | -4.7 |
| 43  | NH$_2$-TCNFP | 6.8 | -6.8 | 38.2 | -4.5 |
| 44  | 2,5-Tricyanocyclopentadiene | 7.0 | -7.0 | 38.0 | -4.7 |
| 45  | Pentacyanophenol | 7.6 | -7.5 | 37.5 | -5.2 |
| 46  | 4-C$_6$H$_4$SO$_3$(+NTNHSO$_3$C$_6$H$_4$-4-Cl) | 7.6 | -7.9 | 37.1 | -5.6 |
| 47  | HI | 7.7 | -7.3 | 37.7 | -5.0 |
| 48  | $\text{NO}_2$-C$_6$H$_4$SO$_3$NH$_2$ | 7.8 | -7.9 | 37.1 | -5.6 |
| 49  | Me-TCNFP | 8.6 | -8.6 | 36.4 | -6.3 |
| 50  | 3,4-(Me$_3$)C$_6$H$_4$-TCNPF | 8.7 | -8.9 | 36.1 | -6.6 |
| 51  | 4-MeO-C$_6$H$_4$-TCNFP | 8.7 | -8.9 | 36.1 | -6.6 |
| 52  | C$_6$N$_2$(+CN)$_2$OH | 8.8 | -8.7 | 36.3 | -6.4 |
| 53  | 4-C$_6$H$_4$SO$_3$(+NTNHSO$_3$C$_6$H$_4$-NO$_2$) | 8.8 | -9.1 | 35.9 | -6.8 |
| 54  | 4,4'-NO$_2$-C$_6$H$_4$SO$_2$OH | 8.9 | -9.0 | 36.0 | -6.7 |
| 55  | $\text{CF}_3$CH(=C$\equiv$C$\equiv$CH) | 9.0 | -9.1 | 35.9 | -6.8 |
| 56  | $\text{HBr}$($\text{CN})(\text{CF}_3)$ | 9.3 | -9.3 | 35.7 | -7.0 |
| 57  | $\text{Ph}$-TCNFP | 9.4 | -9.5 | 35.5 | -7.2 |
| 58  | $\text{HBF}_4$ | 10.3 | -10.0 | 35.0 | -7.7 |
| 59  | $\text{FSO}_3$OH | 10.5 | -10.3 | 34.7 | -8.0 |
| 60  | 3-$\text{CF}_3$CH$_2$-TCNFP | 10.5 | -10.7 | 34.3 | -8.4 |
| 61  | H-TCNFP | 10.7 | -10.7 | 34.3 | -8.4 |
| 62  | $\text{[C}_6$H$_4$O$_3$(+NTN)$_3$]OH | 11.1 | -11.4 | 33.6 | -9.1 |
| 63  | $\text{[(CF}_3)$-2$\text{PO}_4$]OH | 11.3 | -11.5 | 33.5 | -9.2 |
| 64  | 2,4,6$(\text{NO}_2$)-C$_6$H$_4$SO$_3$OH | 11.3 | -11.4 | 33.6 | -9.1 |
| 65  | $\text{[C}_6$N$_2$(+CN)$_2$C$_6$H$_4$]OH | 11.4 | -11.5 | 33.5 | -9.2 |
| 66  | $\text{TIOH}$ | 11.4 | -11.3 | 33.7 | -9.0 |
| 67  | $\text{C}_6$H$_4$SO$_3$(+NTNHSO$_3$C$_6$H$_4$-NO$_2$) | 11.5 | -11.7 | 33.3 | -9.4 |
| 68  | $\text{TIC}(\text{CN})_2$ | 11.6 | -11.5 | 33.5 | -9.2 |
| 69  | Br-TCNFP | 11.8 | -11.8 | 33.2 | -9.5 |
| 70  | $\text{[C}_6$N$_2$(+CN)$_2$C$_6$H$_4$]OH | 11.8 | -12.0 | 33.0 | -9.7 |
| 71  | 3,5-$\text{CF}_3$-C$_6$H$_4$-TCNFP | 11.8 | -12.0 | 33.0 | -9.7 |
| 72  | $\text{TIH}_2$ | 11.9 | -12.0 | 33.0 | -9.7 |
| 73  | 4-C$_6$H$_4$SO$_3$(+NTNHSO$_3$C$_6$H$_4$-NO$_2$) | 12.1 | -12.3 | 32.7 | -10.0 |
| 74  | $\text{Cl}$-TCNFP | 12.1 | -12.1 | 32.9 | -9.8 |
| 75  | $\text{C}_6$F$_3$SO$_3$NH$_2$ | 12.1 | -12.3 | 32.7 | -10.0 |
| 76  | $\text{C}_6$F$_3$SO$_3$NH$_2$ | 12.2 | -12.4 | 32.6 | -10.1 |
| 77  | $\text{CS}$-CH$_2$-TCNFP | 12.3 | -12.4 | 32.6 | -10.1 |
| 78  | $\text{C}_6$F$_3$SO$_3$NH$_2$ | 12.3 | -12.4 | 32.6 | -10.1 |
| 79  | $\text{CF}_3$-TCNFP | 12.7 | -12.8 | 32.2 | -10.5 |
| 80  | $\text{HClO}_4$ | 13.0 | -12.8 | 32.0 | -10.5 |
| 81  | $\text{CF}_3$CF$_2$SO$_3$NH$_2$ | 13.1 | -13.1 | 31.9 | -10.8 |
| 82  | 4-NO$_2$-C$_6$H$_4$SO$_3$(+NTNHSO$_3$C$_6$H$_4$-NO$_2$) | 13.1 | -13.3 | 31.7 | -11.0 |
| 83  | $\text{HBr}$ | 13.3 | -13.3 | 31.7 | -11.0 |
| 84  | $\text{FSO}_3$CH | 13.6 | -13.6 | 31.4 | -11.3 |
| 85  | $\text{TIC}(\text{CN})$ | 14.0 | -16.0 | 30.0 | -12.7 |
| 86  | 2,4,6-Tricyanocyclopentadiene | 15.1 | -15.1 | 29.9 | -12.8 |
| 87  | $\text{CN}$-TCNFP | 15.3 | -15.3 | 29.7 | -13.0 |

$^a$ Estimates of absolute acidities in terms of aqueous pH of 1 : 1 HA/A$\text{^-}$ buffer solution in DCE of the respective acid. $^b$ $pK_{a,rel}$ value of picric acid is arbitrarily set to 0. $^c$ Tos represents the 4-Me-C$_6$H$_4$SO$_3$-group. $^d$ Tf represents the CF$_3$SO$_2$-group. $^e$ X-TCNP represents 2-X-1,1,3,3-tetracyanopropene. $^f$ Reference acid for the $pK_{a,DCE}$ values with a computational $pK_{a,DCE}$ value of 33.
derived from $pK_{a,rel}$ by anchoring to a robust computational value (here: $pK_{a, DCE}(HNITf_2) = 33$). The values are dependent on the medium and thus are strictly non-comparable between different media.

- $pH_{abs}^{H_2O}(BP)$ – medium acidity (as opposed to molecular acidity) of a 1:1 buffer solution expressed in relation to the aqueous pH scale. It is derived experimentally from the $pK_{ip,rel}$ values in Table 2 at the buffer point (BP), corrected to relative $pK_{a,rel}$ and anchored to the $pH_{abs}^{H_2O}$ scale via the experimental procedure delineated above. The values correspond to the thermodynamic proton activity in water at the same pH values, and are comparable between different media.

**Do acids exist that allow for direct $pK_a$ determination in DCE?**

In the course of direct $pK_a$ measurements, when increasing the acidity of the solution, traces of $H_2O$, rather than DCE molecules, will be protonated. With the SMD-calculated medium $pK_{a,DCE}[H_2O]^+$ of 13.6, 0.01 M of an acid HA (as a function of $pK_{a,DCE}[HA]$) and in DCE with $10^{-5}$ M water contamination (0.14 ppm), only acids with $pK_{a,DCE}$ values below 6.7 are suitable for direct medium $pK_{a,DCE}$ measurements. This is 23 orders of magnitude more acidic than the strongest acid CN-TCNP included within Table 2 (entry 87; $pK_{a,DCE}$ 29.7). However, do acids with such $pK_a$ values exist in DCE? Halogenated derivatives of closo-1-carba-undecaborane acids may have such acidities.**

Our computational estimates of their medium $pK_{a,DCE}$ values in DCE are $-0.1$ for $H_3[B_12_{12}]$ and $-1.6$ for $H[CB_{11}F_{12}]$, which means that they should be fully dissociated in DCE solutions! The strongest acid prepared to date is $H[CB_{11}F_{11}]^{32}$ which is only weaker by 21 $k$mol$^{-1}$ than $H[CB_{11}F_{12}]$, and thus it is also expected to be sufficiently dissociated in DCE.**

Can we reach superacidity in DCE?

**Medium superacidity.** From the $pK_{a,rel}$ of sulfuric acid ($-2.2$), we can calculate a $pH_{abs}^{H_2O}(BP)$ of 0.1 for a sulfuric acid buffer in DCE, which is very far off the $-22.4$ we suggest for neutral sulfuric acid as a medium. The main reason is that a conjugate acid anion (here: $[HSO_4]^{-}$) is much less solvated and thus much more basic in DCE than in protic solvents. Even our strongest acid buffer ($pH_{abs}^{H_2O}(BP) = -13.0$), including the molecular superacid CN-TCNP, is more than 9 orders of magnitude away from showing medium superacidity as defined above. But, with the above delineated calculated $pK_{a,DCE}$ values of the carborane acids, it is clear that medium superacidity may be reached in DCE, yet only with special and difficult to prepare acids, and is awaiting experimental realization.

**Molecular superacidity.** By contrast, 55 entries in Table 2 are molecular superacids in DCE holding a lower medium $pK_{a,DCE}$ value than sulfuric acid.

Solvent effects on acidity: a comparison to the gas phase and MeCN

Many acids studied in this work were investigated in acetonitrile (MeCN) and in the gas phase (Table S1 in the ESI†). $pK_{a,DCE}$ and $pK_{a,MeCN}$ are very well correlated (Fig. S3, ESI†), as described by (9):

$$pK_{a,DCE} = 1.08 pK_{a,MeCN} + 33.0, \ s(slope) = 0.02; \ s(intercept) = 0.2; \ n = 44; \ R^2 = 0.992; \ S = 0.6 \quad (9)$$

This indirectly supports the quality of the results and provides a convenient tool to predict the acidities of strong acids in MeCN, which cannot be directly measured in that solvent. As expected, DCE is about 8\% more differentiating than MeCN. The intercept at 33 is just the difference between the standard Gibbs solvation energies of the proton in the two media, *i.e.* in MeCN (1058 $kJ\ mol^{-1}$) and DCE (869 $kJ\ mol^{-1}$). This difference of 189 $kJ\ mol^{-1}$ converted into the log-scale by division through 5.71 $kJ\ mol^{-1}$ gives 33.

**Why are the DCE and GA values poorly correlated?**

Given the inertness of DCE as a solvent, one could expect a good correlation between the acidities in DCE and the gas phase. The reality is very different. The correlation across all acid families is next to non-existent. It thus turns out that even such a low-polarity and inert solvent as DCE is by its influence on acid ionization much more similar to polar aprotic solvents (MeCN) than to the gas phase. Better correlations are obtained when acidities within families are compared (see ESI, Fig. S4†). The poor correlation can be rationalized from a thermodynamic cycle (S-BFHC 3 in ESI†) for the protonation equilibrium between two acids. Thus, the relation between medium $pK_{a,s}$ values and GA values can be derived as:

$$\Delta pK_a = pK_a(HA_2) - pK_a(HA_1)$$

$$= \frac{\Delta GA + \Delta \Delta_{solv}G'[anions] - \Delta \Delta_{solv}G'[acids]}{RT \ln 10} \quad (10)$$

A linear correlation would require that the Gibbs solvation energy difference between a certain acid and its anion is identical for all acids. Obviously, this is not at all the case.

**Conclusions**

Using acids from Table 2, buffer solutions of a well-defined composition can be prepared spanning an acidity range of over 28 pH units (or 160 $kJ\ mol^{-1}$), which is double the pH window of water. Considering an acid/base catalyzed reaction, this means an acceleration or slow-down factor of $10^{28}$. This corresponds to the difference between one millisecond and the age of the universe! It is as yet unclear if medium $pK_{a,DCE}$ and $pH_{DCE}$ values can ever be directly measured in DCE. Despite this, experimental $pH_{abs}^{H_2O}$ values were established in this work that describe the solvated proton’s thermodynamics without any knowledge of the proton’s
specific solvation and its activity in DCE. The quality of the derived $pK_{a,\text{H}_2\text{O}}$ values mainly depends on the quality of the extrathermodynamic assumption (in our case the TATB assumption) and the accuracy of the so obtained single ion transfer thermodynamics. This approach is general and can be transferred to any solvent $S$, given that at least one acid is known for which the transfer energies from water to $S$ exist for all particles. The general relation between aqueous acidities and the range of our buffer acidities measured is shown in Fig. 1. One can easily see that with our used buffer systems, the protolytic window of water is by far exceeded, if compared on the unified acidity scale. However, according to our assessed medium $pK_{a,\text{DCE}}$ values, even in our most acidic buffer system CN-TCNP, the proton concentration of its buffer in DCE of $10^{-29.7}$ mol L$^{-1}$ is much less than one proton per liter. The latter would correspond to $1.6 \times 10^{-24}$ mol L$^{-1}$. This clearly shows that for measurable acid thermodynamics, solvated protons do not need to exist in the medium. Rather, the bound proton in the solvated neutral acid, being in equilibrium with the ionic solvated proton, has the same chemical potential and accounts for the protonation event. However, especially for non-polar media and in unbuffered solutions, impurities may determine the $pH$. Thus, chemical reactions influenced by acidity may proceed in an unpredictable way, if not run in a carefully selected buffer, for example selected from the 87 systems collected in Table 2. The good correlations between the $pK_{a,\text{DCE}}$ values and the corresponding $pK_a$ values in MeCN, as well as in heptane and DMSO (Fig. S7 and S8 in the ESI, respectively†), suggest that the values in Table 2 can, in principle, be reliably transferred to other organic solvents.

**Quantum chemical calculations**

Quantum chemical calculations on the SMD model were done with the same procedure as in ref. 30. rCC calculations were done with the same procedure as in ref. 10. COSMO-RS input files (.cosmo and .energy) were created with the COSMO module of the Turbomole program system according to the "BP_TZVPD_FINE_C30_1501.ctd" formalism. The COSMOTHERM program was used to calculate COSMO-RS Gibbs solvation energies.

**Conflicts of interest**

There are no conflicts to declare.

**List of abbreviations/compendium**

- **Unified acidity**
  - $pK_{a,\text{S}}$: Negative decadic logarithm of the medium acidity constant in the solvent $S$
  - $pK_{a,\text{DCE}}$: Negative decadic logarithm of the medium acidity constant in the solvent DCE
  - $pK_{a,\text{H}_2\text{O}}$: Negative decadic logarithm of the medium acidity constant in the solvent $\text{H}_2\text{O}$
  - $pK_{ip,rel}$: Negative decadic logarithm of the relative (to picric acid) acidity constant of the acid–titrant base ion pair (here the solvent is DCE, unless stated otherwise)
  - $pK_{a,rel}$: Directly measured $pK_{ip,rel}$ corrected for ion-pairing effects according to the Fuoss-model to a true relative (to picric acid) acidity constant (here the solvent is DCE, unless stated otherwise)

- **Activity**
  - $\alpha(\text{H}^+, \text{S})$: Activity of the solvated proton in solvent/medium $S$
  - $\alpha(\text{H}^+, \text{DCE})$: Activity of the solvated proton in solvent DCE
  - $\alpha(\text{H}^+, \text{H}_2\text{O})$: Activity of the solvated proton in solvent $\text{H}_2\text{O}$

- **Other**
  - $\alpha(\text{H}^+, \text{DCE}) = 10^{14}$ mol L$^{-1}$
  - $\alpha(\text{H}^+, \text{H}_2\text{O}) = 10^{29.7}$ mol L$^{-1}$
  - $\alpha(\text{H}^+, \text{H}_2\text{O}) = 10^{-14}$ mol L$^{-1}$
  - $\phi(\text{H}_2\text{O}) = 0$
  - $\phi(\text{H}_2\text{O}) = +14$
  - $\phi(\text{DCE}) = 58.1$
  - $\phi(\text{H}_2\text{O}) = +15.4$
  - $\phi(\text{DCE}) = 29.7$
  - $\phi(\text{H}_2\text{O}) = -13.0$
  - $\phi(\text{DCE}) = 29.7$
  - $\phi(\text{H}_2\text{O}) = -13.0$
  - $\phi(\text{DCE}) = 29.7$
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  - $\phi(\text{DCE}) = 29.7$
  - $\phi(\text{H}_2\text{O}) = -13.0$
  - $\phi(\text{DCE}) = 29.7$
  - $\phi(\text{H}_2\text{O}) = -13.0$
  - $\phi(\text{DCE}) = 29.7$
S

The liquid medium the acidity is investigated in. It may be a molecular solvent like DCE or a strong acid itself.

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Notes and references

‡ We exclusively use the term “acidity” for proton acidity according to Arrhenius, Bronsted and Lowry.**

§ From principal thermodynamic considerations, the \( H_0 \) curve follows \( dH_0 = -RT \ln (10) \cdot (d[H]) - d(BH^+) + d(B) \) and is “contaminated” by the indicator base system’s chemical potential change.

† “Neutral” means that the \( pH_{H_2SO_4} \) is half of the \( pK \) of autoprotolysis. It is commonly addressed as “100% H\( _2SO_4 \”).

** However, a further problem arises from the fact that even the slightly weaker carborane acids like H[\( \text{HCB}_{11}\text{Cl}_{11} \)] and H[\( \text{HCB}_{11}\text{Br}_{6}\text{H}_{5} \)] as well as the H\( _2[\text{B}_{12}\text{X}_{12} \) \( (X = \text{Cl, Br}) \) acids** are known to decompose many solvents including dichloromethane” and thus most probably also DCE by elimination of HCl and formation of carbocations.

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