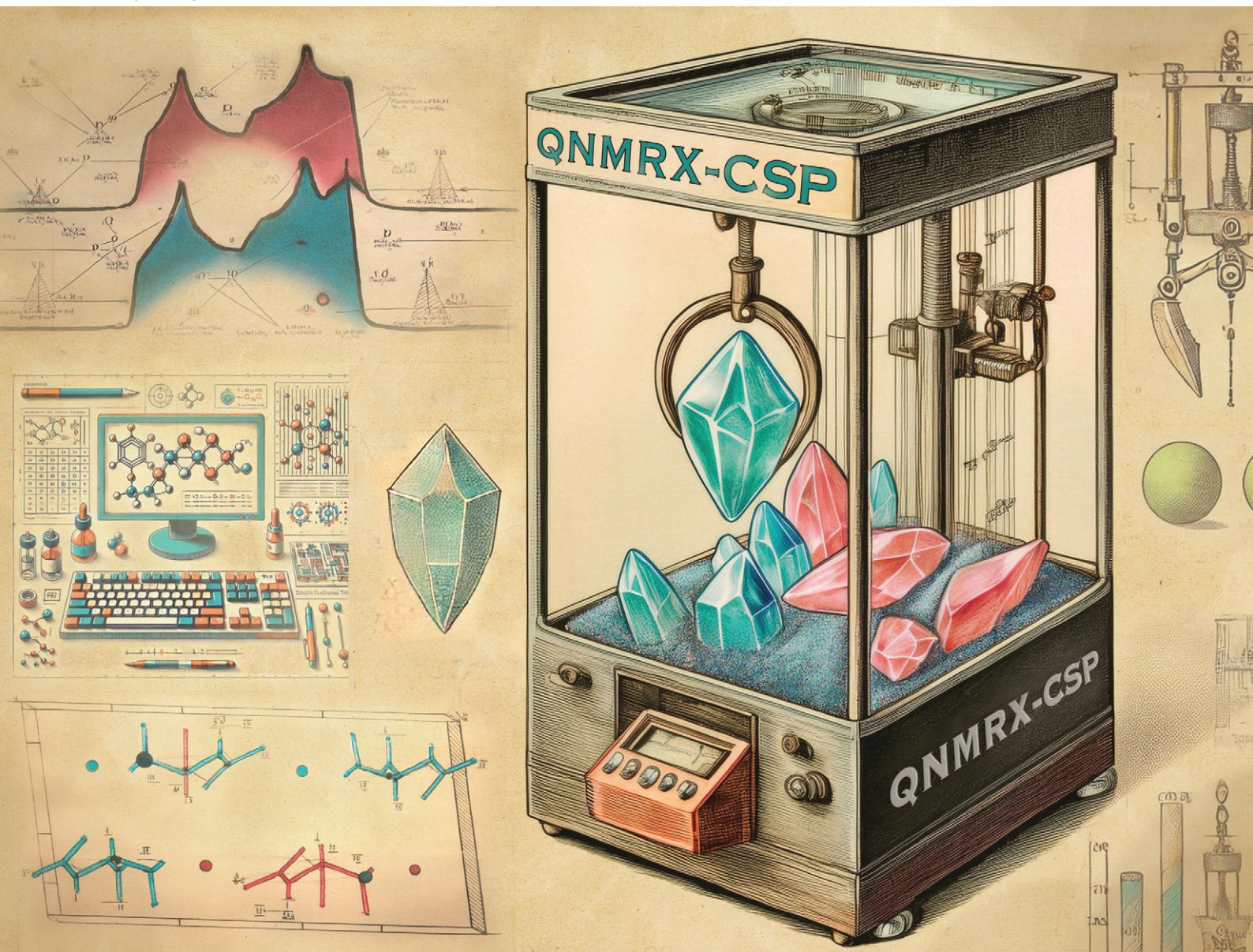


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Quadrupolar NMR crystallography guided crystal structure prediction (QNMRX-CSP)[†]

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We describe a new quadrupolar NMR crystallography guided crystal structure prediction (QNMRX-CSP) protocol for the prediction and refinement of crystal structures, including its design, benchmarking, and application to seven organic HCl salts. Five HCl salts with a limited number of low-energy conformations were chosen as model systems for benchmarking: betaine HCl, glycine HCl, D-alanine HCl, guanidine HCl, and aminoguanidine HCl; two were chosen for blind tests: *N,N'*-dimethylglycine HCl and metformin HCl. The QNMRX-CSP protocol uses experimental ³⁵Cl solid-state NMR (SSNMR) spectra and X-ray diffraction (XRD) data in tandem with Monte Carlo simulated annealing and dispersion-corrected plane-wave density functional theory (DFT-D2*) calculations. The protocol comprises three modules: (i) the assignment of motion groups, (ii) a Monte Carlo simulated annealing algorithm for generating tens of thousands of candidate structures, and (iii) DFT-D2* geometry optimizations of structural models and concomitant computation of ³⁵Cl EFG tensors. Key benchmarked metrics are used for retaining the best candidate structures, including unit cell parameters, static lattice energies, and EFG distances (I_{EFG}). The protocol is shown to generate structural models that are excellent matches with experimental crystal structures that have been DFT-D2* geometry optimized, as validated by crystallographic *R*-factors (*R*) and root-mean squared distances (RMSDs) of atomic positions that are well below 10% and 0.2 Å, respectively. Finally, consideration is given to the use of the QNMRX-CSP protocol as a standalone technique or in concert with other NMRX methods and Rietveld refinements, and possible applications employing other quadrupolar nuclei (*i.e.*, ¹⁴N and ¹⁷O).

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1. Introduction

Crystal structure prediction (CSP) methods are increasingly used to calculate the structures and properties of known materials, as well as for the discovery of new materials.^{1–7} CSP methods can be grouped into two general but complementary areas: (i) first principles calculations for structural prediction with limited or no experimental input and (ii) computational mining of databases such as the Cambridge Structural Database,^{8,9} Crystallography Open Database,¹⁰ and Inorganic Crystal Structure Database.¹¹ CSP is critical for establishing structure–property–function relationships in materials; as such, the Cambridge Crystallographic Data Center⁹ (CCDC)

has conducted a series of collaborative “blind tests” in which they solicited researchers to use CSP methods for the prediction and refinement of crystal structures.^{12–17}

CSP methods are applied extensively to organic compounds^{18–23} and active pharmaceutical ingredients (APIs)^{24,25} to aid in screening of their potential solid forms (*i.e.*, polymorphs, salts, hydrates, solvates, and cocrystals)^{26–29} and prediction of properties.^{30,31} CSP methods are not without their limitations, since they can be computationally expensive in the absence of key structural information, such as the unit cell parameters, the number of crystallographically distinct molecules (*Z'*), and/or the identities of multiple molecular conformations or electronic configurations (*e.g.*, conformers, tautomers, zwitterions, *etc.*). Furthermore, the differences in energy among similar structures tend to be small and can depend on the computational methods, posing difficulties in evaluating and validating structures based solely on their energies.^{32–34} In view of this, CSP methods can benefit from data derived from powder X-ray diffraction (pXRD) (*i.e.*, unit cell parameters, space group, *Z'*) and solid-state NMR (SSNMR) spectroscopy (*i.e.*, *Z'*, molecular conformations, dynamics, disorder, internuclear distance measurements, and bonding).

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In this regard, NMR crystallography (NMRX), which uses SSNMR data to aid in the prediction, validation, and/or refinement of crystal structures, has emerged as a fruitful area of research, complementing and enhancing CSP investigations.^{35–41} In the most simple form of NMRX, SSNMR has long been applied to aid in identifying the space group and/or number of crystallographically distinct sites in the unit cell.^{35,42–44} By contrast, more modern NMRX studies have featured the measurements and/or first principles calculations of ¹H chemical shifts, homonuclear dipolar couplings, and spin diffusion rates to solve crystal structures of small organic molecules,^{45–50} or in other instances, ²⁹Si chemical shifts and homonuclear dipolar couplings to determine structures of zeolites and silicate frameworks.^{51–53}

More recently, contemporary NMRX-CSP methods have demonstrated impressive results for molecular organic crystals and pharmaceutical solids,^{54–60} usually combining computational methods and experimental data that usually include: (i) density function theory (DFT) or other *ab initio* calculations for geometry optimization and calculation of NMR parameters; (ii) SSNMR methods for the accurate measurement of chemical shifts; and (iii) the acquisition of high-quality pXRD data, for purposes of fingerprinting, structural validation, and/or Rietveld refinement.^{61–63} DFT calculations of magnetic shielding parameters, which largely involve comparison to ¹³C, ¹H, or ¹⁵N isotropic chemical shifts,^{64–73} and to a lesser degree, chemical shift tensors,^{74–79} can be computationally demanding, depending on the choice of functional and basis set. Furthermore, modeling the influences of long-range inter- and intramolecular interactions (*e.g.*, hydrogen bonding, π - π stacking, *etc.*) on NMR interaction tensors can be approached with methodologies involving periodic boundary conditions, cluster-based calculations, or fragment-based calculations. GIPAW calculations are widely used for the calculation of NMR tensor parameters from models of periodic solids, but are limited in the type of model chemistry that can be used efficiently (*e.g.*, calculations using hybrid functionals are computationally expensive);^{72,80–82} in contrast, calculations using clusters and/or fragments to model the extended lattice structure permit the exploration of different classes of functionals in an efficient manner, allowing for increasingly accurate calculations of chemical shifts.^{64,79} Finally, since the successful prediction of chemical shifts has generally come with high computational expense, Emsley *et al.* and other researchers have developed machine-learning methods for rapidly predicting ¹H, ¹³C, ¹⁵N, and ¹⁷O chemical shifts, with applications in NMRX-CSP.^{83–88}

Quadrupolar nuclides (*i.e.*, nuclear spins $I > \frac{1}{2}$) have been utilized in NMRX studies to a much lesser degree,^{89–109} though we note there are numerous studies that employ quadrupolar SSNMR, XRD, and DFT methods that are not explicitly described as NMRX. To the best of our knowledge, there are no NMRX-CSP studies that feature the combined use of crystal packing algorithms, DFT calculations of electric field gradient (EFG) tensors, and quadrupolar NMR data for organic solids. This is somewhat surprising, given the many

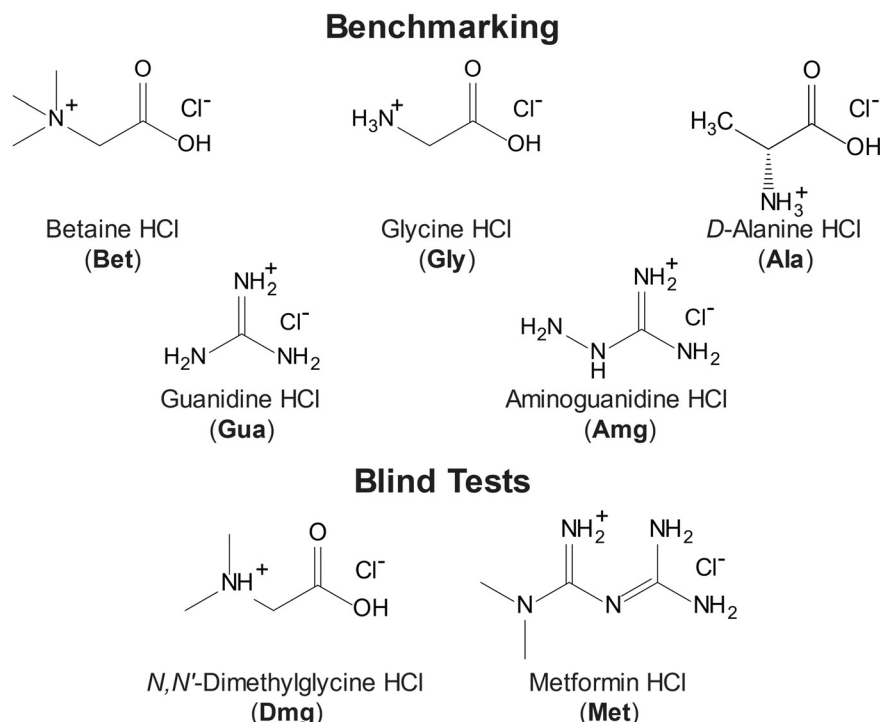
quadrupolar nuclides of elements of great import to organic solids (*i.e.*, ¹⁴N, ¹⁷O, ²³Na, ³⁵Cl, *etc.*) and the exquisite sensitivity of EFGs at the nucleus to the local structural environment (this is especially true of the longer-range, weaker intermolecular interactions associated with crystal packing, which often have only minor influences on chemical shifts). Many studies by our group and others have shown the great utility of the relationships between EFG tensors and atomic environments in API solid forms^{110–119} for purposes of spectral fingerprinting,^{120,121} identifying impurities,¹²² observing molecular-level dynamics,^{123–127} and quantification,^{128–130} which are critical first steps towards applications in NMRX.

Crucial in quadrupolar NMRX methods is the computation of EFG tensors that match well with experiment. In organic solids, the positions of hydrogen atoms are well known to influence ¹⁴N, ¹⁷O, and ³⁵Cl EFG tensors.^{111,131,132} We have found that DFT methods employing a dispersion correction (DFT-D2*) for the accurate refinement of atomic coordinates, particularly hydrogen atom positions, generally result in excellent agreement between calculated and measured EFG tensors.^{133–135} Notably, ³⁵Cl EFG tensors of chloride ions, which are prevalent in the many HCl salts of APIs (nearly half of known solid forms of APIs),¹³⁶ are very sensitive to the numbers, arrangements, and types of hydrogen bonds (under the broadest definition of H...Cl bonds of ≤ 2.6 Å,^{137–139} it is those of *ca.* 2.2 Å or less that most dramatically impact ³⁵Cl EFG tensors).⁹⁷ Hence, with the recent successes of DFT-D2* methodologies, SSNMR of quadrupolar nuclei can start to play an important role in modern NMRX-CSP studies of organic solids.

In this work, we present a new quadrupolar NMR crystallography guided crystal structure prediction (QNMRX-CSP) protocol that uses SSNMR and XRD data, along with a Monte Carlo simulated annealing algorithm and plane-wave DFT-D2* calculations, and its application to organic HCl salts (Scheme 1), with an emphasis on the role of ³⁵Cl EFG tensors. The QNMRX CSP protocol (Scheme 2) is comprised of three modules, each of which features clearly delineated steps, and key metrics for retaining the best candidate structures (see §2.2), which include unit cell parameters, static lattice energies (E_{lat} and E_{cutoff}), and the EFG distances (Γ_{EFG}). Final candidate structures are validated using crystallographic *R*-factors (*R*) and comparison of atomic positions *via* calculations of root-mean squared distances (RMSDs).

We discuss (i) the design of the QNMRX-CSP protocol; (ii) the restriction of the space group and the benchmarking of metrics for selecting candidate structures using structural models of five HCl salts of organic molecules (*i.e.*, betaine HCl, glycine HCl, D-alanine HCl, guanidine HCl, and aminoguanidine HCl; Scheme 1), which have known crystal structures and ³⁵Cl EFG tensors; and (iii) four different stages of calculations, including three benchmarking stages (Scheme 2: S1, S2, and S3, indicated by yellow, green, and blue arrows, respectively) and one stage for blind tests and/or predictions of unknown structures (Scheme 2: S4, red arrow). A detailed walkthrough of benchmarking is presented for the





Scheme 1 Molecular diagrams of the seven HCl salts of organic molecules. Five structural models are used for benchmarking calculations (betaine HCl (Bet), glycine HCl (Gly), D-alanine HCl (Ala), guanidine HCl (Gua), and aminoguanidine HCl (Amg)) and two are used in blind test cases (*N,N'*-dimethylglycine HCl (Dmg) and metformin HCl (Met)).

example of betaine HCl, and blind tests for the predictions of the structures of *N,N'*-dimethylglycine HCl and metformin HCl are considered. Finally, we discuss the use of the QNMRX-CSP protocol for molecules of greater size and flexibility, their implementation with other quadrupolar nuclides, and their general application as both standalone and complementary methods for refinement, prediction, and discovery of new crystal structures.

2. Methods

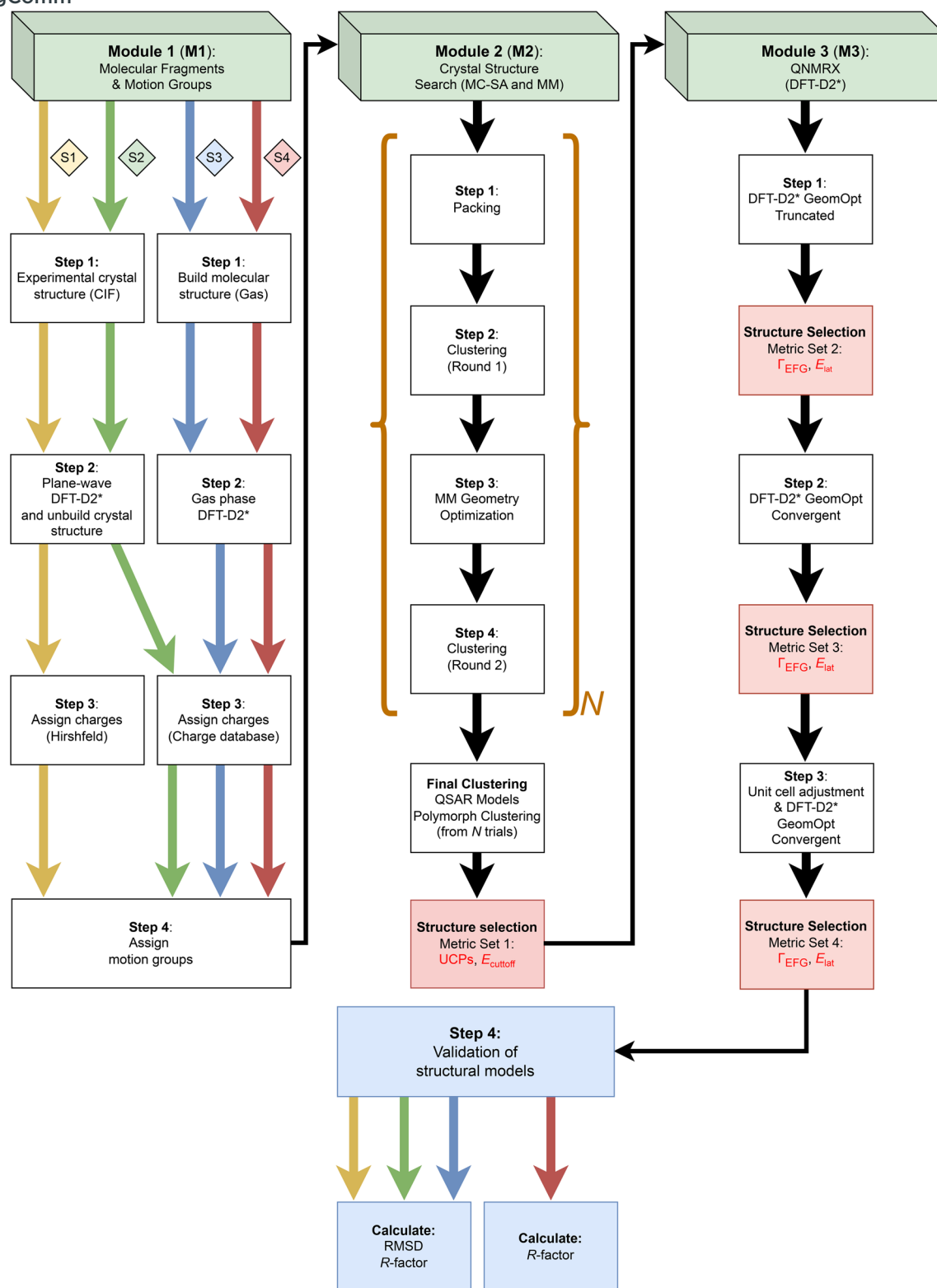
2.1 Computational details

Overview. The following tools were used within BIOVIA Materials Studio 2020 R3:¹⁴⁰ polymorph was used to assign motion groups and generate candidate crystal structures, quantitative structure activity relationship models polymorph clustering was used to cluster multiple polymorph trials, and CASTEP was used to run plane-wave DFT calculations⁸² on either structural models generated from polymorph or obtained from the CSD. See Supplement S1 in the ESI† for information on workstations.

Polymorph. The polymorph prediction routine includes a Monte Carlo simulated annealing algorithm to search a phase space and generate candidate crystal structures. All space groups can be searched during the prediction routine; however, we chose to search space groups of the crystal structures from which the structural models were developed (§3.1). The polymorph prediction routine (steps 1–4 in M2, the Crystal Structure Search module) consists of four steps: (i) packing, (ii)

a first round of clustering, (iii) geometry optimization, and (iv) a second round of clustering, which are all performed using the default “ultra-fine quality” settings. In the packing step (step 1), the maximum number of candidate structures to generate is set to 10 000, with maximum and minimum temperatures of 1.5×10^5 K and 300 K, respectively, heating and cooling factors of 0.025 and 0.0005, respectively, and a minimum move factor of 1.0×10^{-10} . Heating during the packing step is carried out until either the maximum temperature is reached or a minimum of 14 consecutive structures are accepted (*i.e.*, 14 structures, each lower in energy than the previous), at which point cooling begins. A preliminary geometry optimization is included in the packing step (*vide infra*). For the first clustering step (step 2), similar structures are grouped together and the lowest energy representative is retained based on the forcefield type (in this case, the Dreiding force field)¹⁴¹ using the following default set of parameters: a radial distribution cut-off of 7.0 Å (maximum interatomic distance cutoff for electrostatic interactions between pairs of atoms), a tolerance of 0.13 (the minimum similarity measure value for a structure to be included in a cluster, this value ranges between 0 and 1, with 1 being an exact match) and 140 bins (the maximum number of structures that are evaluated in a cluster). The geometry optimization step (step 3) is performed using the following convergence thresholds: 2×10^{-5} kcal mol⁻¹ for energy, 10^{-3} kcal mol⁻¹ Å⁻¹ for forces, 10^{-3} GPa for stress, and 10^{-5} Å for structural displacement. During this step, the positions of the motion groups and unit cell parameters are allowed to vary over 500 iterations (the relative positions of the atoms within the motion groups are kept





Scheme 2 A flowchart of the QNMRX-CSP protocol. The protocol comprises three modules (M1, M2, and M3) indicated by the green boxes at the top of the figure, with each step indicated by a white box, benchmarked metrics used for selecting the best candidate structures indicated by red boxes, and structural validation steps indicated by blue boxes. The black arrows indicate progression through the protocol regardless of the stage, whereas the coloured arrows indicate the stage of the calculation (S1 yellow, S2 green, S3 blue, and S4 red) involved in benchmarking and blind tests. The polymorph prediction routine is indicated by the brown brackets and is repeated N times. Details are given in §3.2.

constant). To calculate the energies of the structural models, an atom-based summation method for all interaction energies (*i.e.*,

electrostatic, Van der Waals, and hydrogen bonding) involving the Dreiding forcefield was applied.¹²⁴ Hirshfeld charges used



in this step were obtained from a protocol described in the ESI† (see Table S1 and Supplement S2 for information on the Charge Database). Finally, the second clustering step (step 4) is performed with the same parameters used in the first clustering step, retaining the lowest energy structures. The polymorph prediction routine is repeated over N trials (in this study, N varies between 10 and 20, *vide infra*).

QSAR models polymorph clustering. After performing N trials, the QSAR models polymorph clustering module¹⁴² was used to group any similar structures across different trials (*e.g.*, trials 1 through 10), retaining the lowest energy structures and discarding the rest. The parameters used are identical to those involved in the clustering steps outlined in §2.1. The final structures from the Crystal Structure Search module (M2) are selected on the basis of unit cell parameters that best match experiment (§3.2) and the lowest static lattice energies (§2.2).

CASTEP. Plane-wave DFT-D2* geometry optimizations^{133,134,143} employed the low-memory BFGS scheme,¹⁴⁴ with structural refinements proceeding for five cycles (*i.e.*, herein designated as the truncated BFGS scheme) or to convergence. Maximum values for structural convergence were set as follows: energy change (5×10^{-6} eV per atom), displacement (5×10^{-4} Å per atom), and Cartesian force (10^{-2} eV Å⁻¹). The RPBE functional was used for all calculations, along with ultrasoft pseudopotentials generated on-the-fly,⁷² a plane-wave cutoff energy of 800 eV, evaluation of integrals over the Brillouin zone using a Monkhorst-Pack grid with a k -point spacing of 0.05 Å⁻¹, and semi-empirical dispersion corrections using a modified version of Grimme's two-body atomic-pairwise model.¹⁴³ Following all geometry optimizations, ³⁵Cl EFG tensors were computed using a nuclear quadrupole moment of $Q(^{35}\text{Cl}) = -8.112 \text{ fm}^2$.¹⁴⁵

2.2 Metrics used in QNMRX-CSP

Several key metrics are used to assess agreement between experimentally determined structures and NMR parameters with their counterparts obtained from calculations.

Powder diffraction. Diffraction patterns from known crystal structures or candidate structures were simulated using the powder pattern module within the Mercury program.¹⁴⁶ Simulations included a Cu K α radiation source with a wavelength of 1.54056 Å, a 2θ range of 5–50° with a step size of 0.02°, Bragg-Brentano geometry, and a full-width half maximum peak width of $(0.1)\theta$. The crystallographic R -factors (R), which can be used as a measure of disagreement between observed (F_o) and calculated (F_c) signal amplitudes, are given by:

$$R = \frac{\sum |F_o - F_c|}{\sum |F_o|} \times 100\% \quad (1)$$

Crystal packing similarity. The crystal packing similarity function¹⁴⁷ in the CSD-Materials module within Mercury¹⁴⁶ is used to compare crystal structures and calculate root-mean squared distances (RMSDs) of the atomic positions relative to a reference structure. Default options are used, including a

packing shell size of 15 molecules with geometric tolerances of 20% for distances and 20° for angles.

Static lattice energy. The static lattice energies (E_{lat}) of structural models are used both after the M2 step 3 geometry optimization (E_{cutoff}) and each M3 DFT-D2* geometry optimization. In M2 step 3, extensive benchmarking calculations across all model systems determined that those structural models with values of E_{lat} that are higher than E_{low} by a factor of $0.135 \cdot |E_{\text{low}}|$ are retained. Since E_{lat} values are negative, an equation defining the value of E_{cutoff} for structural models retained by this metric is:

$$E_{\text{cutoff}} \leq E_{\text{low}} - 0.135 \cdot E_{\text{low}} \quad (2)$$

In M3 steps 1–3, all structures are retained with static lattice energies falling under the lowest value of E_{lat} determined from benchmarking calculations (§3.2).

EFG distance. The degree of similarity between calculated and experimental EFG tensors with principal components V_{kk} , $k = 1, 2, 3$, which are defined such that $|V_{33}| \geq |V_{22}| \geq |V_{11}|$, is evaluated using the EFG distance (Γ_{EFG}) metric,¹³⁴ which is analogous to the chemical shift distance introduced by Grant *et al.*¹⁴⁸ Γ_{EFG} is defined as:

$$\Gamma_{\text{EFG}} = \left(\frac{1}{15} [3\Delta_{11}^2 + 3\Delta_{22}^2 + 3\Delta_{33}^2 + 2\Delta_{11}\Delta_{22} + 2\Delta_{22}\Delta_{33} + 2\Delta_{11}\Delta_{33}] \right)^{1/2} \quad (3)$$

$$\Delta_{kk} = |V_{kk}^{\text{calc}} - V_{kk}^{\text{exp}}| \quad (4)$$

N.B.: because the sign of C_Q (and V_{33}) cannot be determined experimentally, only the absolute magnitudes of the values of V_{kk} are used in this expression.

3. Results and discussion

The development, testing, and application of the QNMRX-CSP protocol involved refinement of the steps within each module in terms of their position, function, and computational expense, as well as benchmarking key metrics over several computational stages that are used to select structural candidates that best match with experiment. We describe: (i) in §3.1 our rationale and methodology for choosing certain simple organic hydrochloride salts as model systems (Scheme 1); (ii) in §3.2 the three modules (M1, M2, and M3) that constitute the protocol, including the individual steps in each module (Scheme 2); (iii) in §3.3 the four stages of calculations (S1, S2, S3, and S4, as indicated by the yellow, green, blue, and red arrows, respectively, in Scheme 2) involved in benchmarking the metrics *via* comparisons to known structures (S1, S2, and S3) or for “blind” tests (S4), with the goal of enabling the prediction of hitherto unknown structures; (iv) in §3.4 a walkthrough of the application of the QNMRX-CSP protocol (in S1) to predict a structural model of betaine HCl



and presentation of similar data for other model systems; and (v) in §3.5 the application of QNMRX-CSP to two blind tests (in S4) to solve the structures of *N,N*-dimethylglycine HCl and metformin HCl.

3.1 Choice of model systems

We highlight five organic HCl salts (Scheme 1) that were used to benchmark the QNMRX-CSP protocol through three stages (S1–S3), including Gly, Bet, Ala, Gua, and Amg, with Dmg and Met used in S4 for “blind” tests. These systems were selected using the following criteria: (i) they are crystalline solids of with a limited number of low-energy conformations; (ii) they have known crystal structures (see Table 1, *N.B.*: there are two polymorphs for Bet and Met, which were not attempted and do not have their crystal structure information listed);^{149,150} (iii) they feature a single chemical formula unit per asymmetric unit (*i.e.*, $Z' = 1$);¹⁵¹ and (iv) they have accurately measured ³⁵Cl EFG tensors (see Table 2) that are in good agreement with those obtained from structural models refined using the DFT-D2* method.^{133,134} Although these molecules are expected to have varying, small degrees of conformational flexibility, their structural models are treated as rigid bodies during the Monte Carlo simulated annealing routines in M2, but are allowed to vary conformationally during the plane-wave DFT-D2* geometry optimizations.

3.2 Quadrupolar NMR crystallography guided crystal structure prediction

The QNMRX-CSP protocol (Scheme 2) includes three modules: (i) M1: molecular fragments and motion groups; (ii) M2: Crystal Structure Search; and M3: quadrupolar NMR crystallography (QNMRX). Each module was designed to include well delineated steps, some of which use key metrics, determined by numerous benchmarking trials, to retain candidate structures. We reiterate that the numerical values of these metrics were benchmarked on several structural models of HCl salts, which are used as upper limits for structural retention. They include unit cell parameters (*i.e.*, a , b , c , α , β , and γ), static lattice energies (*i.e.*, E_{cutoff} and E_{lat} , §2.2 for definitions), and comparisons of experimental and

Table 2 Experimentally measured EFG tensor and chemical shift parameters for organic HCl salts

Compound	C_Q^a (MHz)	η_Q^b	δ_{iso}^c (ppm)	Ref.
Bet	5.95(8)	0.1(2)	95(3)	128
Gly	6.42(5)	0.61(3)	101(5)	113
Ala	6.4(1)	0.75(6)	106(5)	118
Gua	2.65(8)	0.78(4)	101(4)	128
Amg	2.0(2)	0.76(4)	50(3)	111
Dmg	4.74(6)	0.74(2)	103(5)	128
Met	2.44(4)	0.61(5)	60(2)	129

The experimental uncertainty in the last digit(s) for each value is indicated in parentheses. ^a $C_Q = eQV_{33}/h$, where the principal components of the EFG tensor are defined such that $|V_{33}| \geq |V_{22}| \geq |V_{11}|$. The sign of C_Q cannot be determined from the experimental ³⁵Cl NMR spectra, but can be determined from first principles calculations. ^b $\eta_Q = (V_{11} - V_{22})/V_{33}$. ^c $\delta_{\text{iso}} = (\delta_{11} + \delta_{22} + \delta_{33})/3$.

calculated ³⁵Cl EFG tensors, as described by EFG distances (Γ_{EFG} , §2.2 for definitions).

The QNMRX-CSP protocol proceeds as follows: M1 provides the molecular fragments and motion groups, where molecular fragments are defined as chemically sensible groupings of atoms into molecules and/or ions with independent formula units, and motion groups are defined as independent formula units that are allowed to move during M2. We note that there may be multiple motion groups in a molecular fragment; however, for the benchmarking cases described herein, the molecular fragments and motion groups are identical for all intents and purposes. In M2, these molecular fragments are input into a Monte Carlo simulated annealing algorithm, which generates thousands of candidate structures with distinct unit cells and packing arrangements. Structural models are retained based on unit cell parameters and E_{cutoff} . In M3, a series of three distinct plane-wave DFT geometry optimizations are applied, along with calculations of ³⁵Cl EFG tensors and E_{lat} , all of which are used as metrics for retention of structures. Finally, the structural models are validated based on comparison to pXRD patterns by calculating R and RMSDs (see §2.2 for definitions).

We now turn our attention to the details of these modules, beginning with benchmarking calculations, which feature comparison of computationally derived structural models

Table 1 Crystallographic information for organic HCl salts

Compound ^a	Space group	a (Å)	b (Å)	c (Å)	α (°)	β (°)	γ (°)	Volume (Å ³)	Z^b
Betaine HCl	$P2_1/c$	7.428	9.108	11.550	90.00	96.71	90.00	776.054	4
Glycine HCl	$P2_1/c$	7.117	5.234	13.745	90.00	97.25	90.00	507.913	4
D-Alanine HCl	$P2_12_12_1$	7.148	17.590	5.202	90.00	90.00	90.00	654.065	4
Guanidine HCl	$Pbcn$	7.614	9.015	13.051	90.00	90.00	90.00	895.941	8
Aminoguanidine HCl	$P2_1/n$	6.038	7.809	11.103	90.00	90.00	77.07	510.241	4
<i>N,N</i> -Dimethylglycine HCl	$P2_1/c$	7.061	7.194	13.932	90.00	98.88	90.00	699.219	4
Metformin HCl	$P2_1/c$	7.923	13.894	7.923	90.00	114.49	90.00	793.797	4

^a Crystallographic structure database reference codes for crystal structures glycine HCl (GLYHCL), betaine HCl (BETANC01), D-alanine HCl (ALAHCL), guanidine HCl (GUANIDC01), aminoguanidine HCl (AMGUAC02), *N,N*-dimethylglycine (BUTNIN), and metformin HCl (JAMRIY01).

^b All structures have $Z' = 1$. See main text for discussion.



and NMR parameters with known experimental data (*i.e.*, a QNMRX-CSP computation in S1, following the pathway indicated by the yellow arrows in Scheme 1). Here, the objective is to start with a known molecular fragment and execute the QNMRX-CSP protocol while considering the functions of the individual steps and benchmarking of metrics, in order to obtain one or more structural models for validation.

Module 1: molecular fragments and motion groups. The objective of M1 is to obtain suitable molecular fragments with appropriate atomic charges and well-defined motion groups. First, for purposes of benchmarking calculations in S1, a crystal structure is obtained from the CSD and a structural model based on the CIF is subjected to a DFT-D2* geometry optimization. Next, the crystal structure is unbuilt to obtain the molecular fragments (*i.e.*, the chloride anion and organic cation) to which the calculated Hirshfeld charges are applied (Tables S2–S8†). In the initial stages of developing our QNMRX-CSP protocol, the assignments of charges using both Hirshfeld¹⁵² and Mulliken¹⁵³ population analysis were explored (Table S9†); we proceeded with Hirshfeld charges, since this led to valid structural solutions with greater frequency. Finally, the chloride anion and organic cation are set as independent motion groups.

Module 2: Crystal Structure Search. M2 generates thousands of candidate structures with distinct unit cells and arrangements of molecular fragments using Monte Carlo simulated annealing, with the fragments obtained from M1 as a starting point. We chose the polymorph prediction routine as implemented in BIOVIA Materials Studio,¹⁴⁰ which is reported in CSP studies^{154–159} and CCDC blind tests of organic molecules.^{12–17}

The polymorph prediction routine in M2 consists of four steps: packing, a first round of clustering, geometry optimization, and a second round of clustering, where (i) packing randomly generates crystal structures and minimizes their energies with unit cell and geometry optimizations; (ii) a first round of clustering aggregates similar structures into groups and retains the lowest energy representatives; (iii) geometry optimizations minimize the energies of the remaining structures, and (iv) a second round of clustering removes any remaining duplicate structures (§2.1). For a given structural model, this routine is executed *N* times (the value of *N* is subjective, and largely determined by the retention of structural models that pass structural validation in M3, *vide infra*).

There are several parameters in M2 that can be adjusted. First, the algorithm in M2 can be executed with multiple clustering and geometry optimization steps, as demonstrated by Park *et al.*, where a routine consisting of six individual steps was used (*i.e.*, packing, clustering, geometry optimization, clustering, geometry optimization, and clustering) to generate candidate crystal structures of a contorted-hexabenzocoronene.¹⁵⁹ We found the extra steps of a second geometry optimization and third round of clustering unnecessary. Second, each step has its own settings, which strikes a balance between computational cost and confidence

in sampling all possible candidate structures.^{154–159} Third, the choice of force field and summation method for interaction energies is critically important. For organic molecules, two reliable classes of force fields are Dreiding¹⁴¹ and variations of COMPASS (*i.e.*, I, II, and III),^{160,161} which were used in previous CCDC blind tests.^{12–17} The summation methods for these force fields can be chosen as atom-based, group-based, Ewald, or PPPM.^{162–164} In this work, we used the Dreiding force field with an atom-based summation, as this combination provides the best balance between quality and computational cost. Finally, we selected the space group(s) of interest. It is common practice to search the nine most common space groups that occur for *ca.* 95% of organic crystalline solids, including *P*₂/c, *P*₁, *P*₂12₁2₁, *P*₂1, *C*2/c, *P*bca, *P*nma, *P*na2₁, and *P*bcn (of these, the first 5 comprise 75% of known structures).¹⁶⁵ Herein, the choice of space group in M2 is restricted to that of the reported crystal structure (*i.e.*, *P*₂/c, *P*₂12₁2₁, or *P*bcn) in our benchmarking calculations and blind tests as this greatly reduces the computational expense and enable us to validate our metrics of static lattice energies and EFG tensors for structural selection and not space group selection. *N.B.*: in the case of an unknown structure, indexing of a high-quality pXRD pattern could provide information on the space group; however, it is possible to explore different space groups, albeit at increased computational expense (this is beyond the scope of the current work).

On average, one trial of the polymorph prediction routine in M2 (*i.e.*, one iteration of packing, clustering, geometry optimization, and clustering—indicated by the square brackets in Scheme 2) for a small organic HCl model generates between *ca.* 3000 and 5000 candidate structures. The total number of trials ranges between *N* = 10 and 20.

After these trials, the first set of benchmarked metrics involving *E*_{cutoff} (*i.e.*, structures are retained that fall within 13.5% of the lowest energy structure) and unit cell parameters (*i.e.*, *a*, *b*, and *c*, and the unique angle β , which are within $\pm 20\%$ of those of the known crystal structure with α and γ fixed at 90° for the space groups herein), are applied. One final clustering cycle was performed to remove any remaining duplicate structures between trials.

Module 3: quadrupolar NMR crystallography. M3 consists of four steps. Steps 1, 2, and 3 entail three consecutive DFT-D2* geometry optimizations (under different conditions, *vide infra*) and concomitant calculations of ³⁵Cl EFG tensors. The metrics, Γ_{EFG} and *E*_{lat}, were progressively adjusted downwards with each consecutive calculation, for selecting the best candidate structures. In step 4, the structural models are validated by calculating the *R* and positional RMSD values (*N.B.*: this type of structural validation is only made in benchmarking calculations when comparison to a refined crystal structure is possible; for blind tests or unknown structures, only the *R* is calculated for comparison against experimentally derived pXRD data).

The details of the DFT-D2* geometry optimization steps and metrics used in structural selection in M3 are as follows: first, DFT-D2* truncated BFGS geometry optimizations (see



§2.1) are performed and the ^{35}Cl EFG tensors are calculated. These rapidly eliminate many unreasonable structural models without the need for fully-converged geometry optimizations. Benchmarked metrics of $\Gamma_{\text{EFG}} \leq 0.7$ MHz and $E_{\text{lat}} \leq 135$ kJ mol $^{-1}$ are used to retain candidate structures (typically, 10% or less are retained). Second, convergent DFT-D2* geometry optimizations are performed and ^{35}Cl EFG tensors are calculated, and $\Gamma_{\text{EFG}} \leq 0.49$ MHz and $E_{\text{lat}} \leq 50$ kJ mol $^{-1}$ are used to retain the best candidates. Finally, the remaining structural models have their unit cell parameters adjusted to match those of the known crystal structures, followed by a convergent DFT-D2* geometry optimization and calculation of ^{35}Cl EFG tensors. At this point, the benchmarked metrics are the most stringent, with $\Gamma_{\text{EFG}} \leq 0.49$ MHz and $E_{\text{lat}} \leq 1$ kJ mol $^{-1}$. *N.B.*: the adjustment of the unit cell is not possible unless there are unit cell parameters available from a known crystal structure (in benchmarking) or from indexed pXRD data (in blind tests and for unknown structures), which is a limitation of the protocol at this point. This limitation arises, in part, from the tendency of DFT-D2* geometry optimizations to overestimate the unit cell volumes;^{166,167} this may be rectified in future studies, as routines for higher-order dispersion corrections become available in the CASTEP platform (*e.g.*, D3 and D3(BJ)).^{168–170}

In step 4, comparisons of R (experimental *vs.* theoretical pXRD patterns) and RMSDs of atomic positions (when known structures are available) are used to validate the final structural models. In protein and pharmaceutical crystallography, refined structures that are deemed acceptable have $R \leq 25\%$ and $R \leq 10\%$, respectively.^{171–174} In recent CCDC blind tests, submitted structures are reported as being valid with RMSDs ≤ 0.80 Å; however, this large RMSD is dependent on the size and flexibility of the molecule, with the best structures typically ≤ 0.20 Å.¹⁷ In this work, the candidate structural models that passed our final set of metrics consistently yielded values of $R \leq 9.2\%$ and RMSD ≤ 0.12 Å, with most well below the thresholds of the abovementioned standards (see Tables S10–S14†).

M3 is the most computationally demanding module in the QNMRX-CSP protocol; as such, we have explored numerous measures to reduce computational times while not sacrificing the selection of suitable candidate structures. The first DFT-D2* geometry optimization and calculation of EFG tensors is computationally inexpensive; however, the second and third DFT-D2* geometry optimizations and EFG tensor calculations take considerably more time (*N.B.*: these calculations are run in parallel to reduce computational time). The net computational time depends on several factors: (i) the size of the unit cell; (ii) the numbers and types of atoms in the molecular units; (iii) increased numbers of motion groups (including separate molecular fragments and/or conformational flexibility); (iv) the number of candidate structures remaining after the application of a given metric; and (v) computational power. The structural models we have chosen for this study are ideal in the sense that they have relatively few atoms and small unit cells; however, some

initial studies in our group on systems of increasing size and complexity are indicative of success, albeit with longer net computational times (this is beyond the scope of the current work). Of the factors above, an increased number of motion groups presents interesting problems that must be dealt with in both M1 and M3. It is possible that careful selection of multiple low-energy conformers in M1 could greatly accelerate this protocol for calculations on structures of increasing complexity and degrees of motional freedom. These aspects are given preliminary exploration in the discussion of blind tests (§3.5).

3.3 QNMRX-CSP benchmarking stages

The QNMRX-CSP protocol and its three modules described in §3.2 are used for purposes of designing and refining the protocol and benchmarking its metrics. As discussed above, this initial benchmarking stages is referred to as S1 (Scheme 2, yellow arrows). To progress toward blind or *de novo* CSP, three more stages were explored, including two other benchmarking stages, S2 and S3, and a series of blind tests in S4. These stages each start with different types of structural models and/or assignments of Hirshfeld charges, as described below.

Stage 1: known crystal structure and calculated Hirshfeld charges. S1 (Scheme 2, yellow arrows, §3.2), serves as the initial test of the QNMRX-CSP protocol, where the goal is to predict a structure using geometry-optimized fragments and concomitant Hirshfeld charges with the known crystal structure as the starting point.

Stage 2: known crystal structure and charge database. S2 (Scheme 2, green arrows) is identical to S1 in all respects, except for the use of a database of Hirshfeld charges to make charge assignments on the molecular fragments. The motivation for this benchmarking stage is (i) to test the QNMRX-CSP protocol with pre-determined Hirshfeld charges; and (ii) to ensure that when QNMRX-CSP is applied to unknown structures, for which no computationally derived Hirshfeld charges are available, that a reliable database of charges can be used in M1 calculations. Details on the development of the database of Hirshfeld charges, are found in the ESI,† Supplement S1, and Table S1.

Stage 3: gas-phase structural model and Hirshfeld charge database. In S3 (Scheme 2, blue arrows), molecular fragments are not taken from crystal structures, but rather, derived from gas-phase geometry optimizations. First, a structural model of the organic cation is built (net charge of +1), confined within the center of a $(15 \times 15 \times 15)$ Å³ unit cell with $P1$ symmetry, and subjected to a convergent DFT-D2* geometry optimization. This new geometry-optimized organic cation is assigned Hirshfeld charges as described in S2. This organic cation and a chloride ion are subsequently used in calculations in the M1 module. S3 is still considered a benchmarking stage, since (i) structural validation in M3 includes comparison of both R and RMSD values, the latter of which depends on knowledge of the crystal structure; and



(ii) the results of these calculations, which closely resemble the approach to be used for blind tests and unknown structures, were used to refine the QNMRX-CSP protocol and its associated metrics.

Stage 4: blind tests and unknown structures. S4 (Scheme 2, red arrows) is identical in almost every respect to S3; however, it is reserved for the application to systems with unknown crystal structures. This includes blind tests, in which the structure is known to a third party who validates the final structures *via* comparison of RMSDs (*i.e.*, the best structural candidates are submitted to the third party, with only *R* used for final structural validation in step 4). Furthermore, in S4, the unit cell parameters and EFG tensors are derived solely from experiment, and the final candidate structures are validated using benchmarked values of Γ_{EFG} and *R* derived from S1 to S3 (*N.B.*: after the successful prediction of the crystal structure in a S4 blind test, it is possible to use the resulting data for overall refinement of the QNMRX-CSP protocol and improvement of metrics).

3.4 QNMRX-CSP of structural models: a walkthrough for betaine HCl

A walkthrough of the application of QNMRX-CSP in S1 is described below using Bet as the example (Fig. 1–3). The numbers of structural models generated by M2 and M3 steps 1–3, along with the structural candidates selected by application of the benchmarked metrics, are summarized in Table 3.

In M1 (Fig. 1), step 1 consists of obtaining the known crystal structure (*i.e.*, CIF) of Bet (BETANC01) from the CSD. In step 2, a DFT-D2* geometry optimization is conducted on a structural model derived from the CIF to refine the atomic coordinates and calculate the Hirshfeld charges. In step 3,

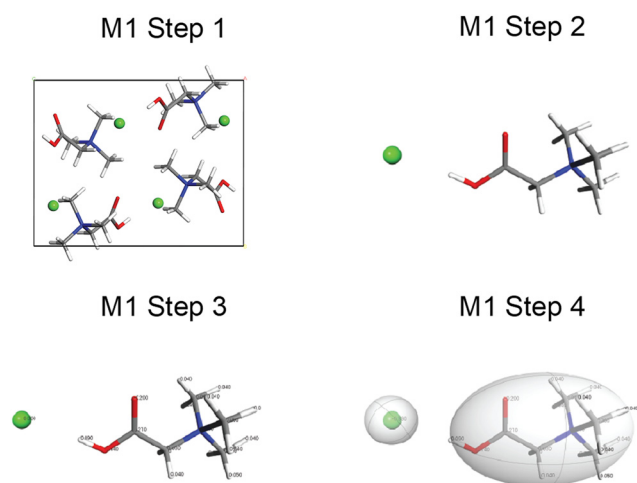


Fig. 1 Schematic diagrams for the walkthrough of the QNMRX-CSP benchmarking protocol for betaine HCl in Module 1, stage 1 (molecular fragments and motion groups). M1 step 1: obtain a known crystal structure (BETANC01). M1 step 2: perform a DFT-D2* geometry optimization and unbuild the crystal structure. M1 step 3: assign the Hirshfeld charges to the atoms. M1 step 4: assign the motion groups.

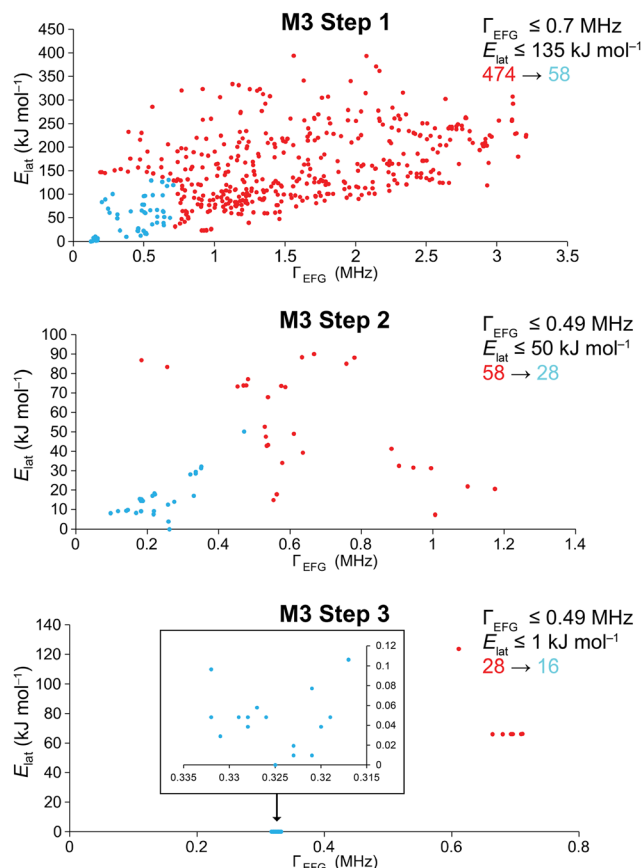


Fig. 2 Scatter plots of E_{lat} vs. Γ_{EFG} for the walkthrough of the QNMRX-CSP protocol in Module 3, stage 1, steps 1–3 (QNMRX) for betaine HCl: red and blue points denote discarded and retained candidate structures, respectively. The numbers of structures before (red) and after (blue) the application of benchmarked metrics are shown to the right. Shown in the inset of the scatter plot in M3 step 3 are the structures that have $E_{\text{lat}} \leq 1 \text{ kJ mol}^{-1}$.

the Hirshfeld atomic charges are assigned to atoms (Scheme SA and Table S2†) on the organic cation and chloride anion. In step 4, the organic cation and chloride anion are assigned as motion groups.

In M2, the motion groups of Bet are subjected to $N = 10$ trials of the four steps of the polymorph prediction routine, which generates 49 451 structures (Table 3). The first set of benchmarked metrics are applied at this point, eliminating the majority of structural candidates; the 474 structural candidates that are retained (*ca.* 1% of the total) are passed to M3.

In M3 steps 1–3 (Fig. 2), DFT-D2* geometry optimizations and increasingly stringent metrics are used to retain the best candidate structures, progressing from 474 → 58 → 28 → 16 (Table 3). In M3 step 4, these final candidates are validated *via* their *R* and RMSDs, which range from 1.1% to 2.7% and 0.002 to 0.119 Å, respectively. We note that $R \leq 3\%$, in line with *R* reported for numerous structures deposited in the CSD, in which *ca.* 90% of structures have $R \leq 10\%$.^{171–174} Two of the sixteen best structural candidates for Bet, 9-632 and 10-103 (*i.e.*, the 632nd and 103rd structure generated



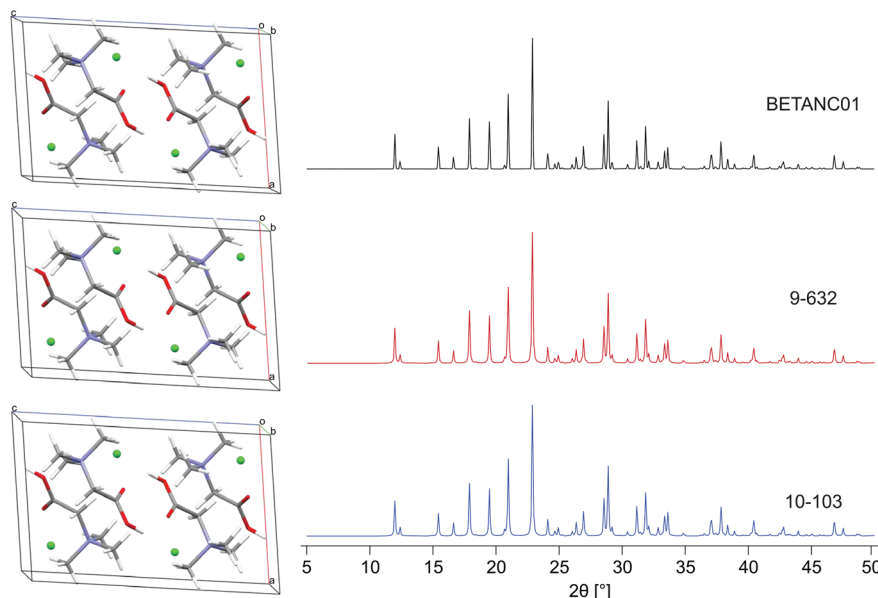


Fig. 3 A comparison of the DFT-D2* geometry-optimized structural model of betaine HCl derived from its known crystal structure (BETANC01) with two (from a set of 16) validated structural models from S1 benchmarking trials, 9-632 and 10-103 with a $\Gamma_{\text{EFG}} = 0.321$ and 0.317 MHz, $E_{\text{lat}} = 0.010$ and 0.106 kJ mol $^{-1}$, $R = 1.135$ and 1.737% , and RMSD = 0.003 and 0.113 Å, respectively.

Table 3 The numbers of structural models retained from the QNMRX-CSP protocol in stage 1 as applied in benchmarking calculations on betaine HCl

QNMRX-CSP protocol ^a	Number of structures	
	Initial	Retained
M2: $\pm 20\%$ $a, b, c, \alpha, \beta, \gamma$; $E_{\text{cutoff}} \leq 13.5\%$	49 451	→ 474
M3 step 1: $\Gamma_{\text{EFG}} \leq 0.70$ MHz; $E_{\text{lat}} \leq 135$ kJ mol $^{-1}$	474	→ 51
M3 step 2: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 50$ kJ mol $^{-1}$	51	→ 28
M3 step 3: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 1$ kJ mol $^{-1}$	28	→ 16

^a Benchmarked metrics for structure selection in modules M2 and M3 are listed in the leftmost column, including the uncertainty in unit cell parameters, a, b, c, α, β , and γ ; and threshold values for the static lattice energies, E_{cutoff} and E_{lat} ; and the EFG distance, Γ_{EFG} (see text for more details).

during the polymorph prediction routine from trials 9 and 10), are compared to the reported structure refined at the DFT-D2* level in Fig. 3. The final Γ_{EFG} , E_{lat} , R , and RMSD values for the validated structures are in Table 4.

The QNMRX-CSP protocol in S1 was similarly applied to Gly, Amg, Ala, and Gua, with validated structural models shown in Fig. S1–S12.† All benchmarking calculations in stages 1, 2, and 3 resulted in successful predictions of the known crystal structure, with all culminating in the selection of valid structural models (Table 5), thereby demonstrating the robustness of QNMRX-CSP for these systems. Visual comparisons and structural validations *via* comparison of the RMSDs and R are shown in Fig. S13–S17 and Tables S10–S14,† respectively.

Crucially, part of the benchmarking process also involved the exploration of structural models with only one of Γ_{EFG} or E_{lat} falling below their threshold values, in order to gauge the importance of having both of these metrics for structural retention. In numerous instances, structural models produced by M3 step 2 that had Γ_{EFG} and/or E_{lat} values above the thresholds were tested in M3 step 3 to see

if convergence and agreement with experiment could be obtained. The vast majority of the structural models of this type failed on both of these counts (examples for Bet are given in Table S15 and Supplement S3†), confirming the importance of both Γ_{EFG} or E_{lat} for retaining the structural models that agree best with experiment.

3.5 QNMRX-CSP stage 4: blind tests

After thorough benchmarking, the QNMRX-CSP protocol was executed in blind tests (*i.e.*, S4 calculations, Scheme 2, red arrows) on two structural models, Dmg and Met (Scheme 1). In the case of this work, the experimental scXRD structures were concealed by a third party, and we were provided with the following: (i) a molecular diagram, (ii) crystallographic information (*i.e.*, space group and unit cell parameters), (iii) experimental EFG tensor parameters, and (iv) pXRD diffraction data.

***N,N*-Dimethylglycine HCl.** In this first blind test involving Dmg, two individuals executed QNMRX-CSP in S4 (Scheme 2, red arrows) following identical procedures, with



Table 4 Validated structural models of betaine HCl from the benchmarking of the QNMRX-CSP protocol in stage 1

Structural model ^a	Γ_{EFG} (MHz) ^b	E_{lat} (kJ mol ⁻¹) ^c	R (%) ^d	RMSD (Å) ^e
2-320	0.325	0.000	1.387	0.112
2-486	0.323	0.010	1.461	0.113
9-632	0.321	0.010	1.135	0.003
9-158	0.323	0.019	1.936	0.111
2-330	0.331	0.029	2.253	0.110
2-494	0.32	0.039	1.549	0.112
4-272	0.328	0.039	2.017	0.002
2-313	0.328	0.048	2.035	0.119
4-46	0.326	0.048	2.059	0.003
7-75	0.332	0.048	2.534	0.002
10-105	0.329	0.048	2.024	0.114
8-152	0.319	0.048	1.765	0.114
10-47	0.327	0.058	1.843	0.113
5-98	0.321	0.077	1.952	0.004
2-326	0.332	0.096	1.955	0.113
10-103	0.317	0.106	1.737	0.113

^a The structural model notation is defined as the trial number-structure number. ^b Γ_{EFG} is the EFG distance; see §2.5 and eqn (2) and (3) for further information. ^c E_{lat} is the static lattice energy of the structural model, normalized to that of the lowest energy structure, which is assigned a value of $E_{\text{lat}} = 0$ kJ mol⁻¹. ^d R is the R -factor, $R = \sum |F_o - F_c| / \sum |F_o| \times 100\%$. ^e RMSD is the root-mean squared distance, which is a measure of the distance between corresponding atomic positions and bond angles from the reported crystal structure and candidate structural model(s).

the goal of independently testing the protocol and to see if both runs converge to similar candidate structures. In M1, the motion groups generated by the individuals (Fig. S18A†) led to similar cationic conformers (denoted as Dmg1 and Dmg2). In M2, the motion groups of Dmg1 and Dmg2 were subjected to a total of $N = 20$ trials each of the polymorph prediction routine (steps 1–4), resulting in totals of 96 606 and 100 040 structures, respectively (Table 6). After applying the first set of benchmarked metrics, 204 and 421 candidate structures, respectively, are passed to M3. In M3, truncated and convergent DFT-D2* geometry optimizations and

selection of the best candidate structures based on benchmarked metrics (steps 1–3, Table 6), both led to 2 candidate structures (for a total of 4) that were validated using R (stage 4, Table 7), which are effectively all identical to one another. QNMRX-CSP was successful in correctly predicting the crystal structure of Dmg in this first blind test Fig. 4, as evidenced by the RMSDs and R -factors that are well within the uncertainties acceptable for structural models obtained from Rietveld refinements, validation of crystal structures using pXRD data, and NMR crystallographic refinements.^{171–174}

Table 5 The numbers of structures retained from the QNMRX-CSP protocol in S1, S2, and S3, as applied in benchmarking calculations on the five structural models

Structural model	QNMRX-CSP protocol	M2 ^a		M3 ^b Step 1	M3 ^b Step 2	M3 ^b Step 3
		Initial	Selected	Selected	Selected	Selected
Bet	S1	49 451	474	51	28	16
	S2	51 855	239	41	24	10
	S3	59 323	276	40	25	12
Gly	S1	55 172	70	9	4	2
	S2	59 348	123	24	15	6
	S3	102 860	70	21	3	1
Ala	S1	42 440	305	63	22	10
	S2	49 045	99	19	5	3
	S3	38 427	259	55	17	4
Gua	S1	50 496	386	81	41	19
	S2	53 659	398	96	48	47
	S3	43 937	149	19	6	6
Amg	S1	61 414	334	63	23	3
	S2	58 380	86	20	8	1
	S3	104 280	84	21	7	1

^a Benchmarked metrics for structure selection in M2 include $\pm 20\%$ a , b , c , α , β , and γ and $E_{\text{cutoff}} \leq 13.5\%$. ^b Benchmarked metrics for structure selection in M3 include $\Gamma_{\text{EFG}} \leq 0.70$ MHz and $E_{\text{lat}} \leq 135$ kJ mol⁻¹ in step 1, $\Gamma_{\text{EFG}} \leq 0.49$ MHz and $E_{\text{lat}} \leq 50$ kJ mol⁻¹ in step 2, and $\Gamma_{\text{EFG}} \leq 0.49$ MHz and $E_{\text{lat}} \leq 1$ kJ mol⁻¹ in step 3.



Table 6 The numbers of structures retained from the QNMRX-CSP protocol in stage 4 as applied in two independent blind tests for two starting conformers of *N,N'*-dimethylglycine HCl, Dmg1 and Dmg2 and metformin HCl, Met1 and Met2

Conformer	QNMRX-CSP protocol ^a	Number of structures	
		Initial	Retained
Dmg1	M2: $\pm 20\%$ <i>a</i> , <i>b</i> , <i>c</i> , α , β , γ ; $E_{\text{cutoff}} \leq 13.5\%$	96 606	204
	M3 step 1: $\Gamma_{\text{EFG}} \leq 0.70$ MHz; $E_{\text{lat}} \leq 135$ kJ mol ⁻¹	204	31
	M3 step 2: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 50$ kJ mol ⁻¹	31	4
	M3 step 3: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 1$ kJ mol ⁻¹	4	2
Dmg2	M2: $\pm 20\%$ <i>a</i> , <i>b</i> , <i>c</i> , α , β , γ ; $E_{\text{cutoff}} \leq 13.5\%$	100 040	421
	M3 step 1: $\Gamma_{\text{EFG}} \leq 0.70$ MHz; $E_{\text{lat}} \leq 135$ kJ mol ⁻¹	421	72
	M3 step 2: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 50$ kJ mol ⁻¹	72	9
	M3 step 3: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 1$ kJ mol ⁻¹	9	2
Met1	M2: $\pm 20\%$ <i>a</i> , <i>b</i> , <i>c</i> , α , β , γ ; $E_{\text{cutoff}} \leq 13.5\%$	54 278	175
	M3 step 1: $\Gamma_{\text{EFG}} \leq 0.70$ MHz; $E_{\text{lat}} \leq 135$ kJ mol ⁻¹	175	26
	M3 step 2: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 50$ kJ mol ⁻¹	26	5
	M3 step 3: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 1$ kJ mol ⁻¹	5	0
Met2	M2: $\pm 20\%$ <i>a</i> , <i>b</i> , <i>c</i> , α , β , γ ; $E_{\text{cutoff}} \leq 13.5\%$	59 420	212
	M3 step 1: $\Gamma_{\text{EFG}} \leq 0.70$ MHz; $E_{\text{lat}} \leq 135$ kJ mol ⁻¹	212	55
	M3 step 2: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 50$ kJ mol ⁻¹	55	7
	M3 step 3: $\Gamma_{\text{EFG}} \leq 0.49$ MHz; $E_{\text{lat}} \leq 1$ kJ mol ⁻¹	7	4

^a See the footnotes in Table 3 for definitions.**Table 7** Validated structural models of *N,N'*-dimethylglycine HCl and metformin HCl in two independent blind tests of the QNMRX-CSP protocol in stage 4

Conformer	Structural model ^a	Γ_{EFG} (MHz) ^b	E_{lat} (kJ mol ⁻¹) ^c	<i>R</i> (%) ^d	RMSD (Å) ^e
Dmg1	17-199	0.116	0.056	1.189	0.013
Dmg1	18-730	0.083	0.015	0.655	0.007
Dmg2	11-1746	0.080	0	0.484	0.004
Dmg2	14-1079	0.090	0.059	0.197	0.015
Met2	7-14	0.067	0	0.086	0.007
Met2	8-24	0.056	0.019	1.075	0.009
Met2	1-15	0.060	0.019	1.42	0.011
Met2	3-19	0.059	0.067	0.962	0.007

^a The structural model notation is defined as the trial number-structure number. ^b Γ_{EFG} is the EFG distance; see §2.2 and eqn (3) and (4) for further information. ^c E_{lat} is the static lattice energy of the structural model, normalized to that of the lowest energy structure, which is assigned a value of $E_{\text{lat}} = 0$ kJ mol⁻¹. ^d *R* is the *R*-factor, $R = \sum |F_o - F_c| / \sum |F_o| \times 100\%$. ^e RMSD is the root-mean squared distance, which is a measure of the distance between corresponding atomic positions and bond angles from the reported crystal structure and candidate structural model(s).

Metformin HCl. Two considerations were made prior to the application of the QNMRX-CSP protocol in S4 to Met. First, the protonation site is unclear, since there are two possibilities: a primary ketimine or secondary ketimine (Scheme 3). Hence, initial DFT-D2* gas-phase geometry optimizations were performed on both molecular fragments, where it was found that the primary ketimine structure is 50 kJ mol⁻¹ lower in energy than the secondary ketimine; therefore, only the primary ketimine was subjected to QNMRX-CSP. Second, there are two possible low-energy conformers of the primary ketimine, Met1 and Met2; gas-phase geometry optimizations reveal that Met1 has an energy *ca.* 3 kJ mol⁻¹ lower than that of Met2.

The remainder of the QNMRX-CSP S4 protocol was carried out using the two conformers of Met1 and Met2 as starting points. In M1 steps 3 and 4, the Hirshfeld charges were applied to the atoms and motion groups were selected (Fig. S18B†). In M2, 10 trials of the polymorph prediction routine

were performed for each conformer, leading to 54 278 and 59 420 structural models, with benchmarked metrics yielding 175 and 212 structural candidates for Met1 and Met2, respectively. In M3, steps 1–3 reduced the total number of suitable candidate structures to 0 and 4 for Met1 and Met2, respectively (Table 6). The final structural candidates were given to a third party, who confirmed their validity. Two of the four validated structures are compared to the known structure in Fig. 5 and a summary of the final Γ_{EFG} , E_{lat} , *R*, and RMSD values are given in Table 7.

The two blind tests resulted in final structural models matching the reported crystal structures (with $R \leq 1.5\%$ and $\text{RMSD} \leq 0.5$ Å), demonstrating that QNMRX-CSP can yield structures comparable to those obtained from diffraction methods. This is very promising for future predictions of unknown crystal structures of HCl salts of pharmaceuticals and other organic molecules. A complete summary of the culmination of the results for both



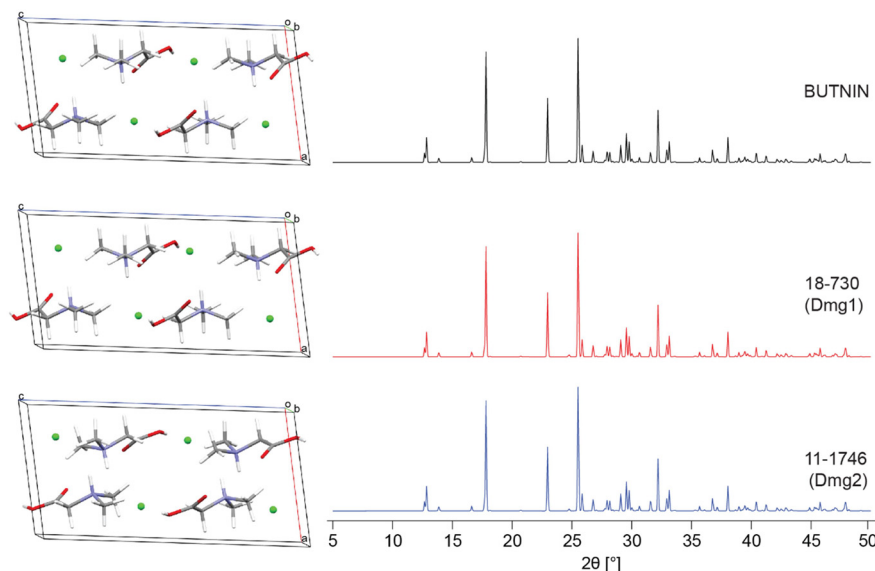


Fig. 4 A comparison of the DFT-D2* geometry-optimized structural model of *N,N'*-dimethylglycine HCl derived from its known crystal structure (BUTNIN) with the validated structural models from S4 blind tests, 18-730 (Dmg1) and 11-1746 (Dmg2), with a $\Gamma_{\text{EFG}} = 0.083$ and 0.080 MHz, $E_{\text{lat}} = 0.015$ and 0 kJ mol⁻¹, $R = 0.655$ and 0.484% , and $\text{RMSD} = 0.007$ and 0.004 Å, respectively.

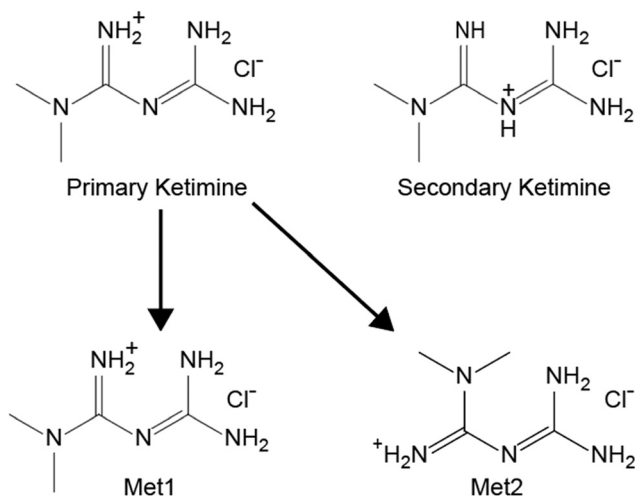
benchmarking across three Stages (*i.e.*, S1, S2, and S3) and the two blind tests representing S4 that are discussed (*vide supra*) can be found in Table 8.

4. Conclusions

We have designed a novel protocol, QNMRX-CSP, for the prediction of crystal structures that utilizes quadrupolar parameters determined from SSNMR data (*i.e.*, ³⁵Cl EFG tensors) and unit cell parameters obtained from

diffraction methods. The QNMRX-CSP protocol was benchmarked using reported ³⁵Cl EFG tensors and scXRD data from five HCl salts of small organic molecules, *via* a process of assigning motion groups, Monte Carlo simulated annealing for crystal packing, and DFT-D2* calculations for structural refinement. Key metrics for selecting the best candidate structures were determined from this process and applied in two blind tests, from which the structures of *N,N'*-dimethylglycine HCl and metformin HCl were predicted. For all systems, the QNMRX-CSP protocol predicted structural models that compare well with experiment, featuring average RMSDs < 0.2 Å and $R < 10\%$ (Table 8).^{171–174}

The QNMRX-CSP protocol, in this nascent stage, has only targeted the use of ³⁵Cl EFG tensor parameters and a limited set of structures; however, there are a plethora of interesting frontiers for its extension. First, it is possible that the protocol may function with other quadrupolar nuclei commonly found in APIs and other organic molecules, such as ¹⁴N and/or ¹⁷O, since calculations of their EFG tensors from structural models obtained from DFT-D2* geometry optimizations have shown excellent agreement with experiment.^{133–135} Unlike chloride ion sites, the oxygen and nitrogen atoms are almost always involved in covalent bonding schemes, which narrows the potential range of EFG tensors that can be investigated for different structural moieties; however, hydrogen bonding is known to dramatically impact these EFG tensors,^{99,111,131} suggesting possible use of ¹⁴N and/or ¹⁷O NMRX-CSP protocols as either a standalone or complementary methodology. Second, the two blind tests, along with ongoing work in our laboratory (beyond the scope of the current work), suggest that this protocol may be useful



Scheme 3 Molecular diagrams of the two protonation sites for metformin HCl (top), the primary ketimine (left) and the secondary ketimine (right). Gas phase calculations reveal the former to have an energy ca. 50 kJ mol⁻¹ lower than that of the latter, only the two conformers of the primary ketimine (bottom) were subjected to the full QNMRX-CSP protocol in S4.



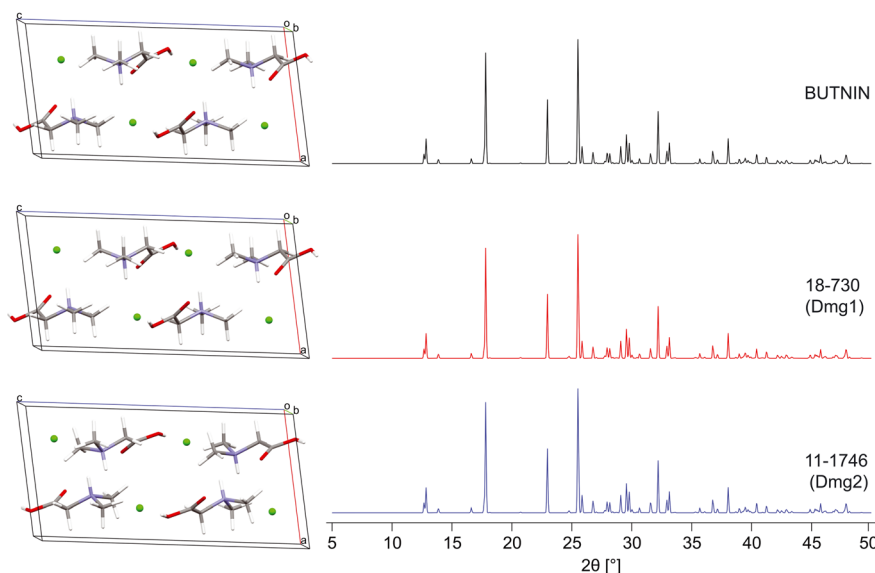


Fig. 5 A comparison of the DFT-D2* geometry-optimized structural model of metformin HCl derived from its known crystal structure (JAMRIY01) with two (from the set of 4) validated structural models from S4 blind tests, 3-19 and 7-14 with a $\Gamma_{\text{EFG}} = 0.059$ and 0.067 MHz, $E_{\text{lat}} = 0.067$ and 0 kJ mol⁻¹, $R = 0.962$ and 0.086% , and RMSD = 0.007 and 0.007 Å, respectively.

Table 8 Summary of key statistics during M2 and M3 and average of final metrics for validated benchmarked and blind tests structures for the QNMRX-CSP protocol

Compound	Stage	Number of trials (N)	Initial number of structures ^a	Final number of structures ^b	Average Γ_{EFG} (MHz)	Average E_{lat} (kJ mol ⁻¹)	Average R (%)	Average RMSD (Å)
Bet	1	10	49 451	16	0.325	0.045	1.853	0.079
	2	10	51 855	10	0.328	0.033	1.643	0.057
	3	10	59 323	12	0.326	0.037	1.693	0.085
Gly	1	10	55 172	2	0.462	0.015	1.855	0.020
	2	10	59 348	6	0.444	0.010	1.478	0.019
	3	20	102 860	1	0.455	0.000	2.173	0.021
Ala	1	10	42 440	10	0.200	0.022	1.175	0.010
	2	10	49 045	3	0.202	0.015	1.689	0.013
	3	10	38 427	4	0.224	0.131	9.10	0.065
Gua	1	10	50 496	19	0.423	0.414	4.476	0.041
	2	10	53 659	47	0.427	0.053	1.968	0.013
	3	10	43 937	6	0.424	0.030	1.693	0.009
Amg	1	10	61 414	3	0.304	0.003	1.156	0.006
	2	10	58 380	1	0.301	0.000	1.060	0.006
	3	20	104 280	1	0.317	0.000	1.271	0.009
Dmg	4	40 ^c	196 646 ^d	4	0.093	0.033	0.631	0.010
Met	4	20 ^e	113 698 ^f	4	0.061	0.026	0.886	0.009

^a Number of structures after N trials of M2. ^b Number of structures after M3 step 3. ^c Total number of trials across Dmg1 and Dmg2 (see §3.5).

^d Sum of structures from Dmg1 and Dmg2 after N trials of M2 (see §3.5). ^e Total number of trials across Met1 and Met2 (see §3.5). ^f Sum of structures from Met1 and Met2 after N trials of M2 (see §3.5).

for the prediction of structures of increasing complexity (*i.e.*, an increased number of motion groups and/or more torsional flexibility in the organic cations). Third, as the sizes of the organic cations are increased, it is possible that the QNMRX-CSP protocol, with EFG tensors alone, may not provide valid structural predictions; however, if the protocol is used in tandem with current NMRX-CSP methods (*e.g.*, involving either the first principles calculations of chemical shifts or their prediction with machine learning algorithms like ShiftML),^{83–87} it is possible that valid structural models may be obtained in

greater frequency, accuracy, rapidity than by the independent application of only one of these methods.

The QNMRX-CSP protocol, in its current form, is limited due to the reliance upon unit cell parameters determined from diffraction methods (this is especially noted in Module 2 where unit cell parameters are a metric, and in Module 3 step 3, where the unit cell parameters of the candidate structural models are adjusted to match those of the reported crystal structure). We believe as methods for dispersion corrections (*e.g.*, DFT-D3*, DFT-D3/BJ)^{168–170} continue to develop and the treatment of unit cell expansion during geometry



optimizations¹⁵⁵ becomes more reliable, their implementation in software packages like CASTEP may be able to better predict the unit cell parameters, thereby reducing dependence on diffraction data. Another pathway for improving and expanding the QNMRX-CSP protocol is the implementation of crystal packing algorithms other than the Monte Carlo simulated annealing routine in Materials Studio/Polymorph; fortunately, there are several options available.^{175,176} This would also expand the use of the protocol to a wider range of researchers with access to different software packages. Furthermore, the calculation of anisotropic displacement parameters must also be a future objective, in order to provide estimates of uncertainties in the atomic positions, akin to those obtained from diffraction studies (preliminary work has already been reported from NMRX-CSP studies reliant upon chemical shifts).^{177,178}

Finally, the QNMRX-CSP protocol was designed with three key points in mind: (1) the predicted and refined structures are obtained after extensive benchmarking, taking into account efficiency and reproducibility; (2) the modules, steps, stages, and metrics are carefully delineated, so that those who are interested in applying the protocol for solving unknown structures have a reliable path forward; and (3) the protocol may be of enormous value in situations where single-crystal structures are not available, since it may be possible to index pXRD data and obtain potential sets of unit cell parameters and space groups, which can be narrowed down using complementary NMR data and QNMRX methods, and ultimately be of use in aiding Rietveld analyses.

Data availability

The data sets supporting this article have been uploaded as part of the ESI.† The CASTEP data manager (CDM) is an executable application developed in-house to automate the execution and processing of truncated and converged DFT-D2* calculations on candidate structures generated through the polymorph prediction routine and is available on the web <https://github.com/rschurko>.

Author contributions

A. A. Peach, C. H. Fleischer III, K. Levin, and J. E. Sanchez participated in the execution of the QNMRX-CSP protocol on benchmarked systems. A. A. Peach and C. H. Fleischer III participated in the blind tests of the QNMRX-CSP protocol. K. Levin developed the CASTEP data manager used for queueing and running calculations in parallel. A. A. Peach, C. H. Fleischer III, K. Levin, S. T. Holmes, and R. W. Schurko contributed to the design of the QNMRX-CSP protocol. A. A. Peach, C. H. Fleischer III, S. T. Holmes, and R. W. Schurko contributed to the writing and editing of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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