The Exploration of Chemical Reaction Networks

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Abstract
Modern computational chemistry has reached a stage at which massive exploration into chemical reaction space with unprecedented resolution with respect to the number of potentially relevant molecular structures has become possible. Various algorithmic advances have shown that such structural screenings must and can be automated and routinely carried out. This will replace the standard approach of manually studying a selected and restricted number of molecular structures for a chemical mechanism. The complexity of the task has led to many different approaches. However, all of them address the same general target, namely to produce a complete atomistic picture of the kinetics of a chemical process. It is the purpose of this overview to categorize the problems that are to be targeted and to identify the principle components and challenges of automated exploration machines so that the various existing approaches and future developments can be compared based on well-defined conceptual principles.
1. INTRODUCTION

Any chemical process may be decomposed in terms of a network of elementary steps. The exact knowledge of all elementary steps, including intermediates, transition structures, and products allows for kinetic modelling and the prediction of concentration fluxes through the network. Due to advances in the field of theoretical and computational chemistry in the past decades and the ever increasing computational power of modern hardware, it has now become feasible to explore chemistry on a broad scale, i.e., tackling the vast dimension of chemical reaction space. In general, the truly exhaustive exploration of some chemical process cannot be guaranteed, but the current state of theoretical and computational chemistry allows for the generation of algorithms that march into an enormously larger fraction of this space than what would be accessible by manual exploration.

Chemical reaction space exploration comprises methods that generate knowledge about a chemical mechanism through atomistic modeling. These methods can be closely tied to data-driven approaches, which attempt to induce reliable results about potential reactivity from existing experimental knowledge in the chemical literature. Reactive pathways are then inferred from rules and models generated on the basis of existing data, such as reactions published in patents or curated databases. These approaches were initially devised to exploit expert generated rules and have recently seen an impressive revival owing to the rise of machine learning techniques. Machine learning facilitates automating and abstracting the rule generation based on amounts of data that were unprocessable before. This data-driven approach can be considered a powerful means to generate ideas about chemical reactivity for new systems similar (according to some measure) to reactants studied in the literature. By contrast, chemical reaction mechanism exploration based on quantum-chemical first-principles methods, which we consider in this work, provides an option, next to experimental synthesis, to probe the validity of the ideas derived from data-driven inference, provided that depth, reliability, and accuracy of the quantum-chemical exploration can be guaranteed. One aim of this overview is to work out the criteria that need to be considered and fulfilled in order to make quantum-chemical mechanism exploration a reliable peer to data-driven reactivity deduction.

In a recent review we discussed in detail algorithms for chemical reaction space exploration. They may be grouped into three classes: i) those that aim at a complete exploration of a given potential energy surface, ii) those that trade breadth for depth by relying on structure hopping and chemical heuristics, and finally, iii) those that exploit human intuition to tame the combinatorial explosion of structures in vast networks involving numerous possible reactants and pathways through, e.g., steering by interactive quantum mechanics. As further reviews on the topic can be found in Refs. we only mention some key methodological work in the field, highlighting that a variety of algorithms and concepts has already been devised. They comprise graph-based approaches, first-principles heuristics that extract rules from the conceptual interpretation of the electronic wave function, chemical heuristics using first-principles calculations, and stochastic approaches. It is not surprising that the global extension of local search and sampling methods has delivered an even broader range of exploration algorithms. Examples are those that exploit artificial forces, growing string methods, exploratory ab-initio molecular dynamics, meta-dynamics, and other enhanced sampling methods. Since exhaustive overviews on the topic have already been provided, this work will focus on comparability and missing links needed to make the different approaches
comparable. Because of the large number of concepts and algorithmic procedures introduced so far, it is rather difficult to have a balanced comparison of the different exploration schemes on the same footing. This becomes even more difficult when considering that exploration algorithms may be designed to serve specific purposes (e.g., gas-phase versus solution chemistry, restrictions to specific compound classes or computational methods, and so forth).

There have been some attempts in the literature to compare existing algorithms at the example of specific target problems. However, such comparisons are difficult. A trivial hurdle turns out to be the parameters and thresholds that control an exploration algorithm, which may be chosen in a non-optimal way, making a direct comparison less conclusive. Moreover, the performance of a some algorithm may also depend on the choice of a specific task. For this reason, it is desirable to have a set of criteria at hand, which allow one to arrive at informed and balanced conclusions about a specific computational approach.

For this reason, we adopt a meta-conceptual perspective aiming at the definition of common general concepts and requirements. First, we discuss and categorize the range of possible mechanistic targets. To keep this task well defined and controllable, we focus on the identification of elementary reaction steps and hardly touch upon the natural extension and combination with subsequent kinetic modeling. For work relating to kinetic modeling in the context of reaction networks we would like to point the interested reader to Refs. 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64.

2. CATEGORIZING MECHANISTIC SEARCHES

The central paradigm of reaction mechanism exploration is the idea that a chemical process under consideration can be mapped onto a (transformation) network of elementary reaction steps connecting reactants and stable intermediates through transition-state structures. As starting point, we may adopt the simplest concepts as they emerge from Eyring’s absolute rate theory (65) (or Kramers’ theory (66, 67)). For the sake of clarity, this may be considered sufficiently fundamental as a network of elementary reaction steps can be arbitrarily refined through subsequent calculations that provide data required for more advanced rate theories.

We consider all parts of chemical (reaction) space relevant for a problem as one huge network that encodes a chemical function. Considering the fact that an exploration process will have some starting point and might already target at a specific end point, we define three principle exploration types: a forward exploration with an open end (FOE), a backwards exploration with an open start (BOS), and a start to end exploration with known start and end (STE) (see Figure 1 for a graphical representation of these exploration types).

Asking ‘How does compound A react with compound B?’ manifests a prototypical forward exploration example with a potentially open end. The compounds A and B constitute a known start. We must then assume that any stable intermediate of the detailed mechanism may, in principle, again react with the starting compounds A and B, or with any other intermediate provided that it is sufficiently long-lived. By contrast, the task ‘How to synthesize compound C?’ constitutes an inverse problem (68, 69, 70), which will be difficult to solve as the number and type of reagents in forward direction is not known in the beginning. Hence, this backwards exploration of open-start type will in general require additional information on potential reagents and catalysts that can promote the forward reaction. The machine-learning based retrosynthesis algorithms mentioned in the Introduction.
Three principal exploration types. Top: forward open-end (FOE) exploration; middle: backward open-start (BOS) exploration; bottom: start-to-end (STE) exploration. Starting material (initial reactants) are given on the left-hand side and are typically of small molecular weight. Common reagents (including ubiquitous molecules in the environment) are depicted separately to highlight their multi-purpose application. A network can grow exponentially in forward explorations, because all newly emerging stable intermediates or side products need to be considered as new reactants.
tion can provide valuable hints on which reagents to choose. If starting and end points of
an exploration are for some reason known and fixed, an exploration algorithm may identify
the viable network of connecting elementary steps and then contribute to the question of
what is the (according to some measure) best pathway and how to promote it.

Due to the fact that the final product(s) of a chemical process may take part in subsequent reactions and that chemical reactions are in general reversible, the labels ‘start’ and ‘end’ can become somewhat arbitrary. Still, the definition of the three types discussed so far represent typical classes of exploration problems.

Although all reagents that are part of the network are treated in the same way, it can be advantageous to single out specific compounds due to their ubiquitous or universal nature as reactants such as, for example, H₂O, O₂, small elimination products (e.g., HBr), (solvated) protons and so on. In this way, screening tasks such as those for a reaction’s robustness towards functionalized molecules [71] can be simplified. Further conceptual steps, such as work-up, reactant separation, and sequential reaction steps can be easily encoded into a network by restricting the connectivity of certain nodes. Note, however, that a specific reference to an actual environment of a reaction is not made in the labels introduced so far. Implicit and explicit solvation, as well as metal surfaces, protein environments, or other environments will initially lead to different, though related networks.

The setting defined so far comprises a broad range of scenarios from simple reaction mechanisms in the gas phase (STE without additional common reagents) to auto-catalyzed reactions with side reactions (FOE with additional common reagents), to probing a retrosynthetic proposal for synthesis (STE with various restrictions and constraints) to screening of an optimum catalyst for a specific target transformation (STE with different catalysts as common reagents). Classifying reaction network exploration protocols according to the three types introduced above allows us to categorize the different algorithmic and conceptual approaches towards the exploration challenge and to compare them on the same basis. Considering the open nature of FOE-type explorations, it is obvious that an actual algorithm will require some adjustable termination determination of the exploration. For the termination of FOE and BOS explorations, bounds on molecular weights are simple criteria and in the case of STE explorations the number of reactions between start and end could be limited. Furthermore, constraints based on kinetic modelling at a certain temperature and other external constraints can be introduced.

3. GENERAL NOMENCLATURE FOR REACTION NETWORKS

Since notions such as ‘system’, ‘molecule’, ‘compound’, ‘reagent’ and ‘structure’ are often used interchangeably, we are advised to introduce and define the essential concepts. To define the components of a reaction network, we first settle on the notion of a ‘molecular structure’:

Molecular structure A single arrangement of atomic nuclei on a Born–Oppenheimer potential energy surface shall be denoted a molecular structure. As a consequence, it is represented by a set of Cartesian coordinates for the nuclei. Depending on the interpretation in terms of a kinetic theory, we will y, in general, be interested in structures with specific properties, most importantly those with vanishing geometry gradients that highlight stable intermediates and first-order transition states.

Properties Each such molecular structure may then be assigned some property that relates to an electronic structure defined by the number of electrons and spin assigned, its
state of excitation (typically the ground state) and any derived molecular property (such as the electronic energy or total dipole moment). Here, we may also allow for conceptual properties such as partial charges and local spin that rely on some chosen decomposition scheme.

With these basic terms we may proceed and introduce a 'compound' as

**Compound** A set of molecular structures with the same nuclear composition and connectivity in terms of chemical bonds denotes a chemical compound. The bonding pattern that is crucial for the assignment to the set may either be fixed based on heuristic rules or, preferably, be determined through a bonding analysis (e.g., in terms of Mayer bond orders).

While such a bonding analysis may often be unique (and particularly helpful to detect and assign molecular structures as different conformers to the same compound), it cannot be excluded that certain structures end up in a gray zone (for instance, if bond orders are very low). This may then create ambiguities on the conceptual level of 'compounds', but not on the more fundamental level of the molecular 'structures' (characterized by a vanishing gradient).

Whereas we consider 'structures' essentially as static for quantum-chemical single-point energy calculations, the concept can be extended to molecular dynamics simulations. Then, however, clustering algorithms will be required to assign sets of structures to a node, which is then a generalization of the 'compound' concept. Transitions between such compounds derived from clustering trajectory data can provide access to kinetic information through Markov state modelling. A single-point-calculation approach will require the explicit generation of all relevant conformers (for example, with the program package RDKit), whereas molecular dynamics simulations then generate compounds through clustering.

In order to generate relations between structures and eventually compounds, transformations from one node in the network to another one are to be uncovered. In general, one will first strive to identify all relevant elementary steps that connect molecular structures and then later generate a reaction network out of these elementary steps that transform compounds into one another. For transformations we will use the following terms:

**Elementary (Reaction) Step** One (or more) structures may be connected with one (or more) different structures via a single transition state representing an elementary step that can be resolved as a sequence of structures along a corresponding minimum energy pathway.

**Reaction** A chemical transformation of one (or more) compounds into one (or more) different compounds represents a reaction that can be distilled from at least one, but in general more than one elementary steps.

As an example, Figure 2 shows a small reaction network built from the four types of nodes defined above.

Any algorithm tackling a chemical question posed in terms of changing molecular structures should explore the transformation steps of these structures and eventually generate compounds and reactions. Hence, chemical reactivity and function will then be encoded in terms of the elementary steps connecting structures as nodes of a network. However, we may introduce a few more general concepts that allow for comparison between compounds.
and reactions. These generalizations are important to cover concepts such as 'residue' that assign a spectator role to certain parts of a molecule, while they are indispensable in an explicit quantum chemical description, which, in general, requires detailed knowledge about all electrons and nuclei involved. As such, these generalizations represent the first step towards mechanistic interpretation, because the network of structures and their connecting elementary steps is agnostic with respect to the assignment of any role in the chemical function under consideration. Moreover, generalizations can be invoked to assign networks that are very different on the level of atoms and elementary particles to the same class. Typical examples are generalizations of molecular function to be fulfilled by a comparatively wide range of compounds, such as are 'oxidizing agent' or 'hydride source'. For these reasons, we introduce three additional terms:

State of a Compound Reducing or oxidizing a compound (and hence, all of its structures), changing the degree of excitation or the spin state, or protonation lead to different molecular structures in a quantum chemical description and would therefore produce different nodes in a reaction network from which new elementary steps lead to further structures. The close relationship to quantum mechanical concepts and prototypical reactions (protonation, electron transfer) make them amenable to automated identification and assignment in quantum chemical procedures.

Reagent A group of compounds that shares the same role in a network form the category of a 'reagent' that is specific for that role. Typically reagent are 'oxidant', 'reductant', 'acid', 'base', and 'catalyst'.

Purpose The concept of 'reagent' allows for comparison and a higher degree of abstraction of a network, which for the network itself is its 'purpose'. 'Purpose' is the superordinate concept to 'reaction'. A typical 'purpose' in reaction chemistry is 'synthesis' or 'catalysis', but the term may as well refer to any molecular functions encoded in terms of structural changes such as 'molecular motor' or 'mechanochemical device'.

To assign an oxidation, charge, spin, excitation, protonation state to a compound not only attributes chemical meaning to a network of structures, it also has consequences for
network data handling, storage, analysis, and presentation. Given the very fundamental nature of these states also allows for their monitoring during exploration.\(^{(82)}\) Moreover, note that some compounds may be part of different sets of ‘reagent’ in different contexts. Early stage exploration algorithms will not feature these last two types of nodes and how they encode ‘purpose’, because of the bottom-up nature of reaction network exploration based on the first-principles of quantum mechanics.

Given two or more explored networks, these concepts define connections between them, which can and should be exploited by exploration algorithms. One type of connection is node sharing, i.e., the fact that a compound (and hence, its structures) can occur also in another network. Calculated data on such parts of a network can be re-used and call for a centralized database storage (the QC\textsc{archive} project by MolSSI\(^{(83)}\) is one example for a technical realization that could be exploited by network exploration algorithms). Another type of connection is contextually joining reaction networks of structures that are very different on a quantum chemical level through the assignment of the same purpose. A typical example is a generic catalytic cycle, which on a realization level can put networks with different catalysts or with differently substituted substrates into the same context. As a result, one could even derive functional networks in which modulating effects of molecular structure are hidden in an environment description of a quantum-chemical embedding approach.

Deliberately, we have left some terms unspecified in order to allow for some vagueness in the context of network exploration. This is convenient as it facilitates comparability of different algorithms and does not interfere with the core concepts introduced above. For instance, we have not specified the term ‘system’ and may use it as a stand-in for the way possible elementary steps are actually set up and then explored in an algorithm.

4. FEATURES OF REACTION NETWORK EXPLORATION ALGORITHMS

4.1. General Foci

The main results of a reaction network exploration will be extracted at the level of compounds and reactions. Context-based abstractions into reagents and purpose then allow for arranging and highlighting the vast amounts of data. Splitting up reactions and compounds into structures and elementary steps allows for understanding the fine details of a particular mechanisms at the level of the physical dynamics of the involved molecular species.

At the level of compounds and reactions, it is reasonable to distinguish between two dimensions: the breadth of a network and its depth. Here, breadth refers to the amount of reactions and compounds that have been incorporated into the network, whereas depth denotes the amount of structures and elementary steps discovered for each of the compounds and reactions. Naturally, both will be difficult to determine in absolute terms as this would require complete knowledge about a chemical process, which is actually the target of an exploration algorithm.

A lack of depth will likely yield qualitatively wrong kinetics, and hence, it will predict wrong distributions of reactants across the networks once these are modeled. Depth fidelity also comprises the accuracy of the chosen computational methods which manifests itself in errors on barrier heights. A lack of network breadth will yield qualitatively wrong results for the total kinetics as it would imply that important side reactions have been missed.

In order to describe a chemical process as reliably as possible, both dimensions have to be pushed to their limits and it will be important to find algorithm-intrinsic measures
that hint toward their saturation. Therefore, it is instructive to define two distinct types of completeness, one for the breadth, the graph fidelity, and one for the depth, the node fidelity.

Naturally, different applications can require a different focus with respect to these directions. Consider the following two examples:

- The calculation of the feasibility of a single reaction cascade to gain insights into its reaction mechanism by exploring a multitude of conformers and pathways requires slow growth of the network breadth and very accurate exploration of its depth.
- The exploration of an entirely unknown set of reactions of given compounds may favor a quick growth, with an a posteriori automatic refinement scheme to increase depth and breadth for 'interesting' parts of the network. A low accuracy in terms of depth for unaccessible and therefore uninteresting parts of the network will be sufficient and often mandatory for the sake of feasibility.

As should have become apparent by now, there exists a multitude of requirements for a generally applicable exploration tool. It shall be able to switch between the different exploration modes defined in Section 2 and also arrive at a sufficient accuracy. To this end, we define a more complete set of key challenges for exploration algorithms in the next section that allows one to classify and assess algorithmic developments regarding their scope and capabilities.

4.2. Challenges

In order to later compare and eventually rank algorithms that explore chemical space, computationally it is imperative to define their goals. These targets presented in the next subsections are driven by the following general challenges:

Validation Challenge The exquisite details that exploration algorithms can generate for any chemical process raise the question of reliability as, in general, no or very little experimental or theoretical reference data will be available. As a consequence, uncertainty quantification will become a crucial part of the whole exploration process.

Operating with Huge Amounts of Raw Data Automated exploration algorithms will be most useful for cases that require thousands or millions of structure searches and optimizations. As a consequence, huge amounts of data will be produced whose manual inspection is not at all possible. As a consequence, automated exploration algorithms must be very stable, handle all data in a fully automated and integrated way, and automatically draw the operator’s attention to critical situations such as convergence failures that cannot be resolved automatically.

Minimal Expectations on the Operator Side The software design that implements an exploration algorithm must take into account that all its parameters, thresholds, screws, and bolts cannot be fully understood on the application side. In other words, the efficiency and reliability of the algorithm should depend as little as possible on knowledge about the intricate effects of changing some of its parameters. Naturally, the default values for them should be expected to be stable and applicable to a wide range of problems and situations.

Unknown Degree of Incompleteness of Generated Data For any reasonably complicated case, it cannot be rigorously proven that the exploration of the corresponding
reaction network of interest has been completely explored. As a consequence, it will be difficult to construct sufficiently complex benchmark cases against which exploration depth and breadth can be measured, and most likely, these will only emerge in a joint effort of various approaches over time.

4.3. Targets

Within this basic setting, we may now formulate goals and targets of exploration algorithms. Some of these goals have been tackled already, but all of them remain to be improved upon in some ways. It is therefore decisive to have this list at hand for future developments in the field.

4.3.1. General Applicability and Stability. A key goal for the exploration of reaction space must be flexibility with respect to the class of molecules and their environments that can be considered. An algorithmic restriction towards a specific compound class, reaction environment, or state of aggregation can be a severe limitation in the discovery process as it would severely limit the domain of exploration and exclude potentially decisive reagents, solvents, and so on. An ultimately useful protocol must be able to accommodate any potentially relevant molecular scenario, even if it is truly hard to map in a virtual screening process. Examples can be found in structures with multiple transition metals in a protein environment or with molecules on metal surfaces.

Moreover, it must be ensured that the algorithms employed are as stable as possible because an exploration protocol may lead them to regions of configuration space that pose problems for them. For instance, orbital convergence will be a key issue – solution methods for complete failure of convergence(86) or for convergence to wrong solutions(87) are registered.

It will, in general, be necessary to operate with a wide range of approaches, spanning fast and less accurate as well as slow and accurate energy assignment protocols, from detailed structure construction and search to advanced sampling through molecular dynamics and Monte Carlo methods. Therefore, the requirements for exploration algorithms in terms of general applicability are immense.

4.3.2. Intrinsic Constraint Monitoring and Adaptation of the Exploration Algorithm. Obviously, any algorithm will involve choices that eventually limit its applicability. For instance, exploring reaction space based on individual evaluations of the stationary Schrodinger equation emphasizes the role of the electronic energy, which will be reasonable as long as it determines the major part of the relevant energy differences. When the exploration then enters a regime of a potential energy surface that is rugged, then this needs to be substituted by a proper molecular dynamics or Monte Carlo sampling approach – and vice versa. The quantum chemically explored reaction network can then be supplemented with proper information from some enhanced sampling approach, possibly encoded in terms of a kinetic model built from this sampling (e.g., through Markov state models(78, 79, 51, 80)).

In order to recognize these limitations, it is crucial to determine the limits with algorithm-intrinsic means while exploring a network in order to achieve sufficient breadth and depth. In the example above, a large number of small barrier heights in some region of reaction space can be taken as a sufficient indication. As long as no fully satisfactory integrated software is available, interoperability of different implementations will be of key
importance.

4.3.3. Taming Conformational Explosion. With increasing size of molecular structures, conformations of a compound become increasingly important. For moderately sized compounds, conformers can be explicitly constructed and optimized. At some point, however, this will no longer be feasible, and again, sampling approaches will be needed. For truly large molecules, this will eventually become a cumbersome task as highlighted by the protein folding problem. The whole menace of conformational depth becomes even more severe when considering transition state structures, especially when embedded into a fluxional environment such as, for instance, water. An exploration based on a single conformation per compound could be a first step to generate an overview on the most interesting parts of a network of elementary steps, but later refinement will then be more mandatory.

4.3.4. Type of Energy Data Provided. It is obvious that ultimate energetical data assigned to nodes of a network should be free-energy data within a well defined thermodynamic ensemble. However, this ultimate goal is not easily accessible to arbitrary accuracy and a first step would therefore be a network based on electronic energies only (to be later refined by modelling contributions from the nuclear framework at some finite temperature). Given the fact that macroscopic constraints such as temperature, pressure, volume, particle number, and so forth can change, it would be desirable to store all data per structure from calculations, which allow one to evaluate free-energy data for changing external parameters at will. Arriving at free energies may be accomplished within very different models ranging from those that start with the standard rigid-rotor harmonic-oscillator particle-in-a-box model (see, e.g., Ref. 57), to continuum solvation approaches (see, e.g., Ref. 89), to additivity schemes (see, e.g., Ref. 90), and to explicit sampling approaches (see, e.g., Ref. 91, 92).

4.3.5. Environment Embedding. Exploring some chemical function or reactivity is most easily accomplished for reactants lacking any environment (i.e., for isolated species that may be considered to represent a gas-phase situation at low pressure). The inclusion of a suitable environment represents a major challenge that must be addressed by an exploration algorithm. Clearly, explicit molecular dynamics in a box of sufficient size under periodic boundary conditions is an attractive choice, but quickly limits the scope of an exploration to the Born-Oppenheimer surface that can be constructed for the elementary particles in that specific box. Changing reagents will not easily be possible. Hence, embedding schemes (for reviews see Refs. 93, 94, 95, 96) play an important role, possible combined in a multi-layer strategy that extends from explicit environment structures close to a reactant to structureless dielectric environments at large distance. Different embedding scenarios will lead to variations of similar networks that require specific care on the data-management size in order to allow for relevant chemical interpretation and conclusion.

4.3.6. Error and Uncertainty Diagnostics. Any model that produces raw data for a network exploration algorithm will rely on certain approximations. For instance, electronic structure models are likely to be based on fast semiempirical or density-functional theory methods, which can be affected by surprisingly large errors for specific structures (see Refs. 99, 100, 101 for examples). Without proper uncertainty quantification, explorations based on such data will be inherently unreliable. Error estimation therefore becomes key. Methods have been devised that point the way of how this can be achieved. 102, 57
4.3.7. Automated Error Reduction. Naturally, nonnegligible errors for certain structures detected during exploration require (automated) refinement by launching more accurate energy calculations. Typically, this will encompass starting reliable ab initio calculations. While this is feasible for single-reference methods such as explicitly correlated coupled cluster theories (105), which can typically be run as black boxes, this is not that straightforward for multi-configurational problems although significant progress has been made in this respect (106, 107, 108). Then, only feasibility consideration are an issue, but may be alleviated by embedding calculations (109, 110, 111, 112). If an exploration protocol allows for such calculations on demand, then the determined error should be propagated through the network in order to reduce the uncertainties for the approximate exploration method. As we have shown (85, 113), this is possible by means of machine learning. Such approaches are determined to play an important role in maximizing the accuracy whilst minimizing the computational cost of the network explorations.

4.3.8. Intuitive and Immersive Interaction and Visualization. As vast networks will contain too much data for a human to grasp, it is imperative that results of any exploration can be displayed in an accessible way and that conclusions can be drawn with the help of algorithms. This requires specific software for human-machine interaction as visual inspection of alphanumerical data will be unfeasible and pointless. Suitable graphical user interfaces are required that also free one from looking at alphanumerical raw data stored in some databank container. They may invoke new hardware (see, for instance, Refs. 114, 115, 116, 117) to intuitively experience these data in order to easily put focus on relevant aspects. Such hardware and software also allows for manual interference and control of the otherwise automated exploration process.

Obviously, interaction cannot mean manual editing of input files for a specific program and a manual database query and insertion, but rather a simple click on a node, opening a context-dependent minimal menu. In view of the definitions introduced above, it is possible to define layers for this purpose reflecting in the first layer structures and elementary steps, in the second compounds and reactions, and then on top of this one abstract layer that encodes reagents and purposes. Figure 2 shows one example of such a representation, also indicating the idea of switching between these layers.

It appears natural that interacting with such an interface and steering the whole exploration protocol would be natural (and for nonexperts most convenient) through language processing algorithms. Voice control of exploration is one option that has already been proposed as a valuable goal for computational chemistry (118, 119). Given the advances in the context of daily-life artificial-intelligence assistants and the availability of open-source language processing libraries (120, 121, 122), this is not at all difficult and even the set-up of an interpretation assistant, which can translate between chemical jargon and the precise ingredients required for quantum chemical calculations on a properly prepared atomistic model, will be rather straightforward.

4.3.9. Maximum Accessibility. While many quantum chemistry programs are reasonably easy to install, an exploration software will most likely involve a database, a user front-end, and a back-end that manages all calculations. Hence, installation and therefore accessibility are more complex by design. Both hardware and quantum chemical software packages
available at the outset of an exploration attempt will result in various setups with different computational capabilities. Given the recent rise of cloud computing and virtual machines an easy access to standardized configurations of the back-end accessible as images is likely a key feature that allows easy setup of the entire machinery for nonexperts. While proprietary software can be made accessible in this way, open-source initiatives are likely to form a more stable and user supported base. The latter are to be preferred in any case due to better reproducibility, reliability, and fidelity. With the plethora of settings already available for quantum chemical calculations and the profound impact they have on the resulting data(123) it will only be possible to reproduce and understand data if their production and processing can be inspected at the source-code level. Given the additional algorithmic layers needed in a reaction network exploration it is of key importance to document and explain possible settings. For the same reasons a comprehensive way of accessing and documenting the used options for a given exploration run is another key part of this goal.

4.3.10. Data Transferability. For efficient explorations it is highly desirable that already generated networks and the contained data can be reused and combined such that subsequent explorations do not need to recalculate them and instead can incorporate existing data seamlessly. Considering the vast amount of standard reactions and the resulting list of frequently used reagents and often occurring reactants, data integration will be highly efficient. However, integration of existing data will only leverage an exploration attempt if uncertainty quantification has rated and labeled the existing data. It is therefore desirable to generate a central library or database of highly accurate results that is continuously extended with data from local explorations, if that data has been generated with an accuracy above a certain threshold. A central database of chemical reaction space would also be a promising starting point for the application of meta-algorithms and machine learning models to exploit and learn from the reaction chemistry mapped out.

4.3.11. Enhanced Kinetic Modelling. Eventually, kinetic modelling will be required to study concentration fluxes through a network. As it is a priori not clear what the number and kind of elementary steps will be that constitute the network, rather general microkinetic solvers will be required, possibly tightly entangled with the exploration algorithm itself to enhance and guide the latter. (63) As a consequence, such solvers should be capable of dealing with vast time-scale ranges and varying degrees of molecularity (at least up to second order kinetics). Whereas special tools (124) and commercial solutions (125) are already available, it is obvious that reaction network exploration presents further challenges that demand more developments. Moreover, to accurately model the flux in reaction networks, theories beyond Eyring’s absolute rate theory will be needed. (126, 127, 128, 129, 130, 131) Clearly, also quantum tunneling must be considered (131). However, improvements on this level will represent a natural extension of a network of elementary steps as the software may automatically gather more information about the potential energy surface in the vicinity of the nodes in such a way that advanced rate theoretical expressions can be evaluated.

5. OPTIONS FOR COMPARISONS OF NETWORK EXPLORATION ALGORITHMS

Comparisons of existing algorithms that explore reaction networks have already been attempted. (46, 47) However, in view of the many aspects to be considered in such an at-
tempt and the fact that a balanced set of benchmark reaction networks resembling various scenarios of practical relevance would be required, demonstrate that such comparisons will not be easy, although certainly needed. It is important to assemble a set of criteria by which an exploration algorithm should be assessed and rated in order to highlight the different problem classes that need to be addressed. It will be necessary to create a multi-dimensional diagnostic to characterize a specific exploration protocol.

Ultimately, any such measure will have to be based on solid comparable data, and hence, benchmark data should be generated and collected in future work. Given the many different types of chemical processes that could be probed, an extendable set of smaller networks appears to be the best choice in practice. Using the three initially defined exploration modes (FOE, BOS, and STE; see Section 2) we may propose three main tests to be run for each benchmark network. In view of the fact that automated network exploration has a significant and important technical and implementational component, extensibility and access should be measured. Furthermore, the reproducibility in terms of the licensing of the source code should also be a factor. Assuming a scale from one to ten, a software that is closed source and requires a specific set of commercial software without any possible extension would be ranked zero. A fully open-source software with the possibility to interface any other quantum chemical code and a set of open-source default programs provided with the exploration tool to guarantee full functionality would rank as ten.

For a fine-grained comparison of exploration algorithms and implementations assessed at benchmark networks five key descriptors sketched in Figure 3 should be sufficient. The five descriptors first and foremost include the two types of reliability measures described above. The node fidelity (depth) is calculated by comparison of the energy spectra of the different nodes with the reference data. The network fidelity (breadth) is calculated as the amount of correctly identified compounds and reactions. Naturally, the expense of the calculations is also one of the descriptors. The computational cost could be measured similar to the work in Refs. (46, 47), which uses the number of gradient calculations as dominant measure. Due to the fact that the accuracy depends on the methods used and that gradient calculations with different methods can vary significantly in duration, the measure should also incorporate an average time of a gradient calculation on a certain hardware. In order to probe the transferability of networks, meaning the possibility to reuse previously generated network data, we may measure the acceleration of an exploration when starting from a subset of already existing nodes in the reference network and also supplementing the running exploration on the fly with structures surfacing during the exploration that are found to be already in a central database. Furthermore, nodes which are automatically determined to be similar (e.g., nodes that were calculated with different solvents as environment) can and should be used as starting guesses. In order to credit the fact that these software packages and algorithms should eventually be useful to non experts in the field of computational and theoretical chemistry, it is important to measure the user-friendliness (immersion ) and also the degree of full automation of a given protocol. The amount of human work required to steer and control an automated network exploration software could be taken as a measure for user-friendliness and for the level of automation achieved. Unfortunately, this may require extended users studies. Given the vastly different approaches to actually solve many of the problems discussed in this work, a challenge similar to the CCDC organic crystal structure prediction challenge(132) or the SAMPL challenge(133) could be a desirable testing ground for the field. It would allow for better comparisons for subtopics and also help focus the efforts on particular unsolved problems.
6. ASSESSMENT OF THE CURRENT STATUS

So far, we have discussed general concepts and goals for reaction network exploration algorithms and their implementation. To assess the current status of the field we will highlight some aspects according to the literature. First, it is noteworthy that most exploration methods employ existing quantum chemical software packages and models. They generally do not implement new electronic structure models in a closed-source form, which has advantages for availability, reliability, and reproducibility of the overall method. The rigorous estimation of the associated errors on structures and energies, their propagation and minimization during the exploration have been tackled only in a few instances.\(^{(57, 84, 58, 85)}\)

In terms of the scope of the chemistry that has been the subject of explorations attempts so far, most methods work more or less routinely for basic (gas-phase) organic chemistry, and some of the reported algorithms have been shown to work with transition metals\(^{(134, 43, 23, 28, 18, 20, 21, 40, 25)}\). The explorations reported so far have largely avoided explicit description of environments such as solvents or protein embedding. Most applications reported for exploration algorithms have been of the proof-of-principle type. It is therefore still a long way ahead of us until we have general implementations available.
that can be routinely employed in daily research. No study has reported data on a net-
work with, say, 1,000 confirmed compounds and 10,000 possible reactions. (Note that this
implies $\gg 1,000$ conformers to be generated and $\gg 10,000$ elementary steps to be calcu-
lated.) The challenges of the visualization and analysis of such an amount of data and the
transferability of sub-spaces of the network have only been touched upon\textsuperscript{23,24}. Overall,
however, important developments have been accomplished and key steps have been taken
(see Sec. \textsuperscript{1}). Considering three major requirements, i.e., a robust yet flexible exploration
algorithm, a scalable and extendable back-end for the actual computations, and kinetic
modelling relating the generated data to actual chemistry, we note that no single software
framework has been established so far that accomplishes these goals so that comparisons of
implementations on equal footing become feasible. We note, however, that we have set out
to provide such a program package\textsuperscript{135}.

7. CONCLUSIONS

In this work, we provided a general description for automated atomistic reaction network
exploration algorithms. Owing to the huge diversity and heterogeneity of the tasks, we
considered it necessary to define core concepts, targets, and challenges in order to make
this fast growing field accessible to assessment and validation and to identify weak spots
in existing schemes. Although a detailed rating of existing schemes has been beyond the
scope of this work for various reasons, of which the limited accessibility of implementations
is an important one, the present overview may still be very well suited as a general guideline
toward the application range and capabilities of existing algorithms.

We defined three basic exploration patterns for chemical reaction networks that de-
pend on the scientific purpose for which they are designed. In connection with a basic
nomenclature presented afterwards we then introduced two fidelity measures related to the
breadth and depth of a network. These are graph accuracy, i.e., the correct identification
and connectivity of nodes, and node accuracy, i.e., the correct description of the set of
structures that represent a compound. We continued to classify prototypical tasks for net-
work exploration algorithms. These include a high level of automation, error diagnostics
and automated uncertainty handling, network data transferability, and eventually kinetic
modelling. Finally we proposed a transferable comparative scheme of reaction space explo-
rati on software based on the concepts and targets elaborated on in this work. Key measures
will be: node and graph accuracy, software extensibility, automation, immersion, and user-
friendliness, data transferability, and of course, the computational cost associated with a
given algorithm/software.

We are certain that such a multi-dimensional ranking will become decisive in future work
on reaction network exploration algorithms as it can advance the field by clear categorization
of new exploration protocols in the context of existing ones. Detailed quantitative measures
can then be provided for existing and new schemes once balanced benchmark networks
become available that represent the variety of algorithmic features highlighted in this work.

| SUMMARY POINTS |
|----------------|
| 1. Predictive theoretical work on molecular reactivity and function will require in all but the simplest cases the study of a huge amount of molecular structures and their relation. |

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2. Automated procedures are mandatory for this task and recent developments have point into fruitful directions for such approaches.

3. Reaction-space exploration approaches must address a wide range of requirements ranging from stable exploration procedures for advanced atomistic modelling to uncertainty quantification for error assessment and cure to new visualization and immersion soft- and hardware to interact with complex networks of many thousand nodes.

4. The comparison of exploration algorithms and software is a multi-dimensional task that requires to carefully assess the pros and cons regarding their theoretical background, efficient and stable implementation, and overall software engineering.

FUTURE ISSUES

1. Future developments of algorithms for chemical (reaction) space exploration will require a very broad theoretical basis in order to cope with the many different scenarios that one may encounter in real-world molecular processes.

2. Such algorithmic developments will eventually demand a high level of sophistication of software engineering and integration in order to move on from advanced tools for specific problems to general tool boxes for explorations of molecular processes at the nanoscale.

3. Eventually, fully automated computational chemistry software will become a peer to a human operator with respect to molecular science in research and education if it can act autonomously on arguments and questions raised in natural language by the operator.

DISCLOSURE STATEMENT

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

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