Quantum materials with strong spin–orbit coupling: challenges and opportunities for materials chemists

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Spin–orbit coupling is a quantum effect that can give rise to exotic electronic and magnetic states in the compounds of the 4d and 5d transition metals. Exploratory synthesis, chemical tuning and structure–property characterisation of such compounds is an increasingly active area of research with both fundamental and application-related outlooks. However, these exotic states have an extremely sensitive dependence on the structure and bonding of the host materials, meaning that great care is required with regards to their chemistry, but this is not always considered sufficiently. This Perspective will give an accessible introduction to topical materials with strong spin–orbit coupling, their crystal chemistry, and challenges in their synthesis and characterisation that must be faced if their structure–property relationships are to be correctly determined. These are common to investigations of the same compounds in other fields and rely on the contributions of materials chemists.

1 Introduction

Whilst the properties of all materials originate in one way or another from quantum mechanics, the term ‘quantum materials’ has recently come to be used to collectively categorise those materials – such as superconductors, Mott insulators, and those with topologically-protected states, amongst many others – that have macroscopic properties which cannot be described simply through their constituent atoms and electrons behaving as classical (or at least ‘semi-classical’) particles.1–3 Materials exhibiting quantum magnetism (Fig. 1(a)), as a contrast to conventional magnetic materials, provide another – highly-topical – example. Atoms with unpaired electrons have a net magnetic moment, and in conventional magnets these can be considered to interact with one another according to the principles of standard ferromagnetic and antiferromagnetic exchange interactions to form ordered states. An increasing number of materials, however, are being discovered in which quantum-mechanical effects such as fluctuations, particle entanglement, wavefunction topology and spin–orbit coupling are significant enough that those conventional magnetic states are in competition with exotic ones that are now being revealed for the first time.4–6

(a) Quantum magnetism
(b) Photovoltaics
(c) Catalysis
(d) Energy storage

Fig. 1 Strong spin–orbit coupling underpins the electronic and magnetic properties of many topical materials, such as (a) the realisation of quantum spin liquids and other exotic states in compounds of the platinum group transition metals,7 and (b) determining the band gap of hybrid metal halide perovskites (shown are calculated band structures for MAPbI3).8 These materials also have a variety of functional properties – for example, the same iridates that have been heavily studied with regards to quantum magnetism also have exceptional (c) catalytic9 and (d) electrochemical10 properties. All of these are rooted in the structure and bonding of the material, so there is great potential for different communities of materials chemists to benefit from each others’ investigations. Figures reprinted/adapted with permission from (a) ref. 7 Nature, 2018; (b) ref. 8, (c) ref. 9 and (d) ref. 10 American Chemical Society, 2013, 2019, 2017.
The quantum effect of interest in the materials that we will focus on in this Perspective is spin–orbit coupling. This is the interaction between the spin angular momentum \( S \) and the orbital angular momentum \( L \) of the electrons, which is expressed in terms of a new quantum number \( J \) that redefines the energy levels of the electrons.\(^1\) Spin–orbit coupling perturbs the wavefunctions of the electrons under its influence and the magnetic forces acting on them, affecting their spatial distributions and magnetic moments. We will discuss it in the context of its effect on the magnetic and electronic states of transition metal compounds, which strengthens on moving from 3d to 4d and 5d systems.

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As such, materials featuring 4d or 5d transition metal cations with \( t_{2g} \), \( t_{2g}^2 \) or \( t_{2g}^3 \) electron configurations, such as \( \text{Ir}^{4+} \) and \( \text{Ru}^{4+} \), are sought after as hosts for the exotic electronic states that can result from strong spin–orbit coupling. It is worth noting that many of the materials in which such states have been found have actually been known for decades,\(^2\) but the development of new theoretical frameworks and experimental probes was required.

Perspective

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materials discovery and development is progressing along a strikingly similar path to that in the quantum materials field, for example through the use of soft-chemical and electrochemical methods to access and manipulate lithium-, sodium- and hydrogen-containing materials. In fact, the exact same materials are often under study by the two communities at the same time.

At the outset it is also worth noting that, although materials such as iridates have so far been the most popular for studies of d electrons with strong spin–orbit coupling, it is not only oxides that host such states but also e.g. halides and intermetallics, and in addition to compounds of the platinum group metals that we focus on there are a growing number of materials based on other 5d elements, such as rhenium and tantalum. Developments in theory also continue to motivate efforts in this field.

2 Exotic electronic and magnetic states in materials with strong spin–orbit coupling

The current explosion of interest in the effects of strong spin–orbit coupling on the behaviour of d-electron systems was sparked by the discovery, in 2008, that the iridate Sr₂IrO₄ has an insulating ground state. With its spatially extended 5d valence orbitals this material was expected to be metallic, like its 4d analogues Sr₂RuO₄ and Sr₂RhO₄. Initial tempering of surprise may have been sought by turning to the Mott-Hubbard model, which accounts for many 3d oxides expected to be metallic actually being insulating by considering the magnitude of the electrostatic repulsion U between electrons occupying orbitals of the same atom. A larger U encourages the electrons to remain localised, and above a critical value manifests as the opening of a band gap at the Fermi energy \( E_F \) that is the cause of insulating behaviour in materials without otherwise filled bands. However, that critical value increases with the improvement of d-d orbital overlap from 3d to 4d and 5d systems that broadens the valence d band, and a sufficiently large value cannot normally be obtained for a Mott-insulating ground state to be realised. However, such a state is realised in Sr₂IrO₄ because the \( t_{2g} \) valence band is first split by strong spin–orbit coupling into two distinct ones defined by their different effective values \( J_{eff} \) of the total angular momentum quantum number \( J \) (Fig. 2). The higher-energy \( J_{eff} = \frac{1}{2} \) band straddles \( E_F \), and being now separate from the lower-energy \( J_{eff} = \frac{3}{2} \) band is sufficiently narrow to be susceptible to the Mott-Hubbard splitting that makes Sr₂IrO₄ an insulator.

Mott behaviour and the effect of \( U \) has underpinned the rich physics of 3d transition metal oxides that has been revealed over the last decades. The realisation since 2008 that strong spin–orbit coupling can also provide access to novel states has led to the search for – and discovery of – unprecedented exotic behaviours in iridates and other 4d and 5d systems. A prominent contemporary example is Kitaev magnetism, including the Kitaev quantum spin liquid state. A spin liquid is a magnetic state in which the spins of different atoms are entangled with...
one another through space but don’t establish long-range magnetic order. The Kitaev model is a fully-solvable model of $S = 1/2$ spins on a honeycomb lattice and describes the possibility of a spin liquid arising from ‘bond-dependent’ magnetic interactions (see Fig. 1(a)) that contrasts with the physics of ‘traditional’ Cu$^{2+}$-based spin liquid candidates.4–6,11,20,25–29

Electronic phases with unusual band structure topologies, such as the topological semimetal phases found in some pyrochlore iridates and in SrIrO$_3$, are also a feature of materials with strong spin–orbit coupling. Another recent development has been the exploration of potential excitonic magnetism in d$^3$ compounds. The effects of spin–orbit coupling and $U$ generally cause t$_{2g}^4$ systems to have a non-magnetic $J_{\text{eff}} = 0$ ground state, but an exotic form of magnetism dubbed excitonic magnetism that is mediated by excitations to the $J_{\text{eff}} = 1$ state may be found.2,3,4 This can also compete with the formation of orbital molecules, where a covalent-like sharing of electrons between cations forms $M$–$M$ bonds.11 This bonding does not solely originate from strong spin–orbit coupling – the essential requirement is simply for a structure in which the d orbitals of neighbouring cations can interact directly.5,6,42,43,51 However, as many of the materials being studied for the effects of strong spin–orbit coupling have such a structure, orbital molecule states are amongst those being found.52–57

There are many other exotic electronic and magnetic states that have been found in materials with strong spin–orbit coupling which we will not discuss here (see e.g. ref. 58–60). We also confine ourselves to discussion of bulk materials, though thin films and heterostructures provide unique and exciting access to further tuning parameters and exotic phases.21 Many excellent reviews exist for the interested reader.4–6,11,20,25–29

3 Crystal chemistry

In the majority of the materials with strong spin–orbit coupling that have been studied to date, the active transition metal cation $M$ is octahedrally coordinated by oxide ligands. MO$_6$ octahedra therefore constitute the basic building block of their crystal structures, which are constructed by connecting these octahedra through different patterns of corner-, edge- and face-sharing. These connectivities define the $M$–$M$ and $M$–O–$M$ orbital interactions, which in turn determine the electronic band structure and magnetic exchange interactions for a given material. So that the relationship of the exciting properties arising from these interactions to the underlying structure of the host materials can be better understood, we now outline some common structure types and their relationships in terms of the connectivity of the MO$_6$ octahedra (Fig. 3). In general formulae we use M to indicate spin–orbit-coupling-active transition metal cations (commonly Ir or Os in the materials we discuss) and A to indicate non-active cations (typically Group 1, Group 2 and rare-earth elements). Furthermore, whilst we refer to these in general as oxides, many of these structure types can be realised with halide and chalcogenide anions too.41,61,62

This section is not intended to replicate comprehensive reviews of all known structure types and their possible variations.63,64

In particular, the range of known polymorphs and superstructures, arising for example from perovskite rotation/tilting distortions, different stacking arrangements in layered rocksalt-type materials and hexagonal perovskites, or the (dis)order of vacancies in topochemically-tunable materials, is too great to be covered here. Instead, this section is intended to be an introduction to relevant crystal chemistry, upon which the nuances of the detailed
Fig. 3 When a transition metal cation is octahedrally coordinated, its five valence d orbitals are split into $t_{2g}$ and $e_g$ sets by the octahedral crystal field (CF). The $t_{2g}$ orbitals retain orbital angular momentum, so for species with $t_{2g}^1$, $t_{2g}^2$, $t_{2g}^4$ or $t_{2g}^5$ electron configurations, these can be further split by spin–orbit coupling (SOC) into states defined by their effective total angular momentum $J_{\text{eff}}$. Many of the materials currently of interest for the effects that spin–orbit coupling has on their magnetic and electronic properties have structures built from such octahedra and a selection of more prominent examples are depicted here, categorised according to the different connectivities of the octahedra. The compositions of common structure types are given with a general formula, where $M$ is the spin–orbit-coupling-active transition metal (blue atoms) and $A$ is a different, non-active species (green atoms). For simplicity all such compositions are shown as oxides (red atoms), though many of these structure types are also adopted by chalcogenide and halide materials. Crystal structures visualised using VESTA.\textsuperscript{65}
structural characterisation that understanding quantum states requires will be developed in later sections.

### 3.1 Rocksalt-related structures

We start with a discussion of materials with structures derived from the rocksalt type, as a number of the most well-studied materials with interesting spin–orbit coupling effects fall into this category. The rocksalt structure comprises a face-centred cubic anion lattice within which cations occupy all the octahedral interstitial sites, giving a 1 : 1 ratio of cations to anions. Structurally, each edge of each MO₆ octahedron is shared with a neighbouring one. Derivative structures are realised when there are multiple cationic species that have an ordered distribution over these octahedral sites. In z-NaFeO₂-type materials the two cationic species, the transition metal M and a Group 1 cation A, order into alternating layers perpendicular to the 111 axis of the cubic rocksalt unit cell, generating a rhombohedral supercell. Within each layer the MO₆ octahedra share edges with six neighbours, forming a hexagonal lattice. However, changing the ratio of the cationic species from 1 : 1 forces one species to occupy sites in the layer of the other. In Na₂IrO₃ and z-Li₃IrO₃ (better described as e.g. Na₁[NaIr₂]O₆) that extra Na/Li cation occupies ordered sites such that they sit in the centre of the ‘cells’ of a honeycomb lattice formed by the remaining Ir. Thus, the 2D honeycomb structure that is the basis for the Kitaev spin liquid state is realised. x-RuCl₃ (and IrCl₃, though being d⁶ it does not exhibit a magnetic ground state) is also structurally – they are effectively the same materials with all Na/Li removed.⁶⁶,⁶⁷ Analogous z-OsCl₃ is not known, though substoichiometric Os₃Cl₃ (in which honeycomb nanodomains may be realised through local ordering of the Os-site vacancies) and partially-lithiated Li₁₅Os₀₈₅O₃ have the same structural basis.⁶⁸,⁶⁹ The layered motif of these materials facilitates topochemical removal/exchange of the interlayer species. In this way, A’IrₓOs₃O₈ (A’ = H, Cu, Ag) have been prepared.⁷⁰–⁷² As these A’ cations favour the linear coordination of an O–A’=O dumbbell over the octahedral AO₆ coordination of Li/Na in the parent materials, these materials have a different stacking pattern of the honeycomb layers than in z-AIrO₃, though that structural motif is retained with the remaining Li cation in the cells of the honeycomb lattice.⁷⁰ The exchange of these in addition to those located between the honeycomb layers has been demonstrated with Cu₁IrO₃.⁷₃ Li₁IrO₃ also has β and γ polymorphs in which the zigzag chains of edge-sharing IrO₆ that define the honeycomb lattice in the α phase propagate in three dimensions instead of two, generating 3D networks described as hyperhoneycomb and stripy honeycomb, respectively.⁷₄ Despite these different long-range connectivities, the local environment in which each IrO₆ octahedron shares its three orthogonal edges with three neighbours is the same in all three polymorphs, so where z-Li₃IrO₃ provides a structural template for 2D Kitaev physics its β and γ polymorphs allow for the study of the same physics in 3D. The sodium analogue, β-Na₃IrO₆, is of interest for the same reason and was actually first synthesised by the battery community.⁵⁵

A face-centred cubic anion lattice is the basis not only of the rocksalt structure but many others, defined by their different fillings of the octahedral and tetrahedral interstices. In spinels, tetrahedrally coordinated cations A and octahedrally coordinated cations M are present as AM₄O₁₂. The MO₆ octahedra share edges with six neighbours, forming infinite chains that run along the six 110-type directions of the cubic structure. Only a few AM₄O₁₂ spinels with 4d/5d M-site cations are known, though the edge-sharing chains allow for interesting orbital effects.⁵⁵,⁷₅ However, spinel-type phases can be accessed by increasing the ratio of A to M cations, such that the former partially occupy the sublattice of the latter. If this occupation is spatially ordered the A cations can ‘block’ particular M–M interactions, defining for example a hyperkagome Ir sublattice in Na₃IrO₆ and Na₃IrO₅.⁷₆ The kagome lattice is a 2D network in which each cation interacts with four nearest neighbours, and has become an important structural template for studying the effect of geometric frustration on magnetic ordering – due to competing interactions in vertex-sharing triangles of cations, more exotic ground states may realised.⁷⁷ The hyperkagome lattice is its 3D analogue, and as with the honeycomb and hyperhoneycomb lattices described previously, despite their different long-range connectivities the local environments of the octahedrally-coordinated M species are the same in both the kagome and hyperkagome networks, providing 2D and 3D templates upon which exotic ground states can be realised.⁷⁷

### 3.2 Fluorite-related structures

The MO₃ fluorite structure is also derived from a face-centred cubic lattice, though here it is formed by the cations, whilst the anions occupy the tetrahedral interstices. The cations have eight-fold cubic coordination, though this coordination number is reduced in derivative structures when the anion-to-cation ratio is reduced. One such derivative is the A₂MO₃ family of osmates and iridates (A is a rare-earth cation), in which one-dimensional chains of corner-sharing MO₆ octahedra run in parallel to the c axis of the orthorhombic unit cell.⁷⁸–⁸⁰ The cations M are also octahedrally coordinated in pyrochlores, A₂M₂O₇, though here the corner-sharing chains have 3D connectivity. It is notable that, although one is based on octahedra sharing corners and the other on them sharing edges, the spatial arrangement of the M cations in pyrochlores and spinels is the same – they themselves form a network of corner-sharing tetrahedra known as a pyrochlore lattice that is also a template for geometric frustration. Iridium and osmium both form pyrochlores with a range of A-site species such as Cd, Hg, main-group and rare-earth elements.⁸¹–⁸⁴ However, as one of the seven oxygens in the pyrochlore formula unit is crystallographically distinct from the other six and is not involved in the MO₆ coordination octahedra, pyrochlore materials are susceptible to oxygen non-stoichiometry, as in for example Pb₂Ir₂O₆.⁵⁵

### 3.3 Perovskite-related structures and isolated octahedra

The structure most commonly described in terms of its construction from octahedral MO₆ units is perhaps the perovskite type, adopted by materials with AMO₃ ideal stoichiometry, in which corner-sharing MO₆ octahedra surround a central cavity occupied by the A cation. The AMO₃ lattice with a vacant A site
can by itself be stable, as demonstrated by the binary oxides ReO$_3$ and WO$_3$; however, the attractiveness of the perovskite structure to materials chemists derives from its flexibility to accommodate $A$-site cations (either single species or solid solutions) with a wide range of ionic radii relative to that of the chosen $M$. This allows careful control not only of the number of valence d electrons available to the $M$ species, through variation of its oxidation state, but also of how these interact with one another, because this flexibility is achieved through cooperative rotations and tiltings of the MO$_6$ octahedra in their corner-sharing network. Judicious choice of the $A$-site structure therefore allows sensitive manipulation of the O-M-O bond angles. The ideal perovskite structure, in which these angles are all 180 degrees, is in fact relatively rare – more commonly, the $A$-site structure is smaller and less ideal, and this manifests in a rich variation of distorted variants, such as the orthorhombic GdFeO$_3$-type adopted by SrIrO$_3$, NaOsO$_3$ and others.\footnote{86,87}

The corner-sharing perovskite network is the basis for numerous derivative structures. Layered variants such as the Ruddlesden–Popper ($A_nB_2O_{3n+1}$) and Dion–Jacobson ($AA_nB_{2}O_{3n+1}$) types separate the framework of MO$_6$ octahedra into 2D layers, though as in the 3D analogues the MO$_6$ octahedra are susceptible to tilting distortions that can dramatically affect quantum properties determined by orbital interactions. These can be further impacted by the reduced dimensionality, which reduces the width of the valence d band, leading to stronger Mott effects. A similar effect can also be achieved by introducing a second octahedrally-coordinated species $M'$ which, when $M$ and $M'$ have very different ionic radii or charge states, is stabilised in a cation-ordered structure.\footnote{88} In these $A_2M'M''O_6$ double perovskites, such as BaMgReO$_6$ and Ba$_2$NaOsO$_6$, the $M$ and $M'$ species order in a rocksalt-like pattern, and this provides a template upon which the magnetic $M$ species are spatially separated by the non-magnetic $M'$ ones, reducing the width of the valence d band and allowing the electrons to experience both strong spin–orbit coupling and a large $U$.\footnote{89} This separation of the MO$_6$ octahedra is taken to its extreme in the halides Cs$_6$TaCl$_6$ and Rb$_2$TaCl$_6$, which effectively have a rocksalt-ordered double perovskite structure in which the $M'$ site is vacant.\footnote{91} In other words, the TaCl$_6$ octahedra have no structural connections to one another. Isolated IrO$_6$ octahedra are also found in tetragonal Sr$_3$La$_{11-\delta}$Ir$_{2}$O$_{24}$.\footnote{92}

### 3.4 Multimer-based structures

AMO$_3$ perovskites in which the $A$ cation is too large adopt a different type of structure, forming a family of materials known as hexagonal perovskites. Here, instead of sharing corners the MO$_6$ octahedra share faces, forming dimer, trimer and larger multimer units oriented parallel to the axis of three/six-fold rotational symmetry of the unit cell. This symmetry element has made hexagonal perovskites of emerging interest because it again provides a structural basis for geometric frustration, whilst the face sharing allows for unusually short $M-M$ distances and small $M-O-M$ bond angles that can dramatically influence their quantum properties.\footnote{90} Hexagonal perovskites effectively have a stacked structure, formed by the face-sharing multimers sharing corners with neighbouring multimers (which can be of the same or different types), and this gives rise to extensive polymorphism derived from different stacking arrangements. AMO$_3$ materials can also show polymorphism between hexagonal and conventional perovskite structures – for example, the ambient pressure polymorph of SrIrO$_3$ is actually of the hexagonal 6H type, and the conventional GdFeO$_3$-type polymorph mentioned previously is a metastable phase accessed through high-pressure synthesis.\footnote{86,94}

Dimers of MO$_6$ octahedra are also found in materials adopting the KSO$_3$-type structure, though here the octahedra in each dimer are connected by a shared edge. The dimers themselves are linked by shared vertices and form a three-dimensional tunnel-like structure. As well as materials with AMO$_3$ stoichiometry such as KIrO$_3$,\footnote{93} variants the same tunnel-like structural framework but different stoichiometries, such as $A_3M_2O_9$, $A_3M_3O_{10}$ and $A_4M_6O_{19}$, can be prepared through control of the synthesis method and the choice of $A$-site cation.\footnote{93}

In both edge- and face-sharing multimers the $M-M$ distances are relatively short, and this may allow for sufficient direct interaction between the valence orbitals of the cations of neighbouring octahedra that they form multi-site molecular orbitals over which the valence d electrons delocalise. In other words, the magnetic ‘units’ of the structure may no longer be the individual MO$_6$ octahedra but multimer ‘molecules’.\footnote{90,94,95}

#### 3.5 1D chain structures

The edge- and face-sharing connectivity of MO$_6$ octahedra is not limited to forming just small multimer units. We have already introduced a few structure types in which edge-sharing octahedra form infinite chains, such as $\pi$-NaFeO$_2$ layered rocksalt and AM$_2$O$_4$ spinel, though in these the octahedra share multiple orthogonal edges such that the chains intersect and form two- and three-dimensional networks respectively. One dimensional systems, in which infinite chains run parallel to one another without intersecting, are also known. In the rutile structure adopted by metal dioxides such as OsO$_2$ and IrO$_2$, chains of edge-sharing octahedra chains run along the short axis of the tetragonal unit cell. Corner-sharing of all six vertices of each octahedron connects each chain to four neighbours. Comparatively, in the structure adopted by CaIrO$_3$ and NaIrO$_3$ only the axial vertices of the octahedra are corner-sharing, whilst the equatorial ones are only involved in edge-sharing, giving an arrangement of 1D chains connected in 2D planes.\footnote{96}

The CaIrO$_4$ structure also features edge-sharing chains but with no corner-sharing, so the 1D chains are isolated from one another.\footnote{64} Isolated 1D chains are also found in structures with corner- and face-sharing connectivities of the MO$_6$ octahedra. The former are found in, for example, the fluorite-type $A_3M_2O_9$ materials mentioned previously, whilst the latter are found in recently-reported K$_3$Ir$_2$O$_6$.\footnote{97}

### 4 Structure–property relationships

The determination of structure–property relationships is a key principle of materials chemistry, and in materials with strong
spin–orbit coupling this can be exceptionally challenging. This challenge is the result of their often extreme sensitivity to subtle structural distortions, as the different forces acting on the electrons are finely balanced and exquisitely responsive to changes of $M$–$M$ distances and $M$–$O$–$M$ angles that perturb the band structure and magnetic exchange. In honeycomb iridates, for example, minute distortions of the $\text{IrO}_6$ octahedral network or spacer ion configuration are sufficient to drive the material into a completely different phase (Fig. 4(a)), and many target phases occupy only a tiny volume of parameter space (Fig. 4(b)). Furthermore, the positions and concentrations of light elements, which are particularly difficult to characterise, are often crucial to the exotic behaviour displayed. It has been estimated that the presence or absence of hydrogen at a particular site can change the magnitude of the Kitaev coupling in $\text{H}_2\text{LiIr}_3\text{O}_6$ by a factor of 3.100

It is obvious in such cases that, if the structures and compositions of materials with strong spin–orbit coupling are not accurately known, the virtuous cycle between experiment and theory that is so essential for understanding these materials cannot be optimal. In addition, highly detailed (and often painstakingly collected) physical characterisation measurements cannot be correctly assigned to a stated material if the nuances that distinguish real-world materials from theoretical models have not been considered. In the following sections we will discuss some of the challenges that the study of quantum materials faces in this regard, and outline potential solutions that are coming to light – and which may even seem intuitive to materials chemists from other fields. We will also present some notable examples from the recent literature that illustrate the confusion that can result when these challenges have not been addressed, with the aim of motivating more detailed characterisation of new materials immediately upon their discovery and thorough re-investigations of existing systems where necessary; such efforts have proven extremely fruitful to date.66

5 Materials chemistry challenges

5.1 Structure determination and refinement

As introduced in Section 3, materials with strong spin–orbit coupling almost always contain both very heavy metallic elements and very light elements such as oxygen, and perhaps even lithium or hydrogen. This poses a challenge for structural characterisation as the most common structural probe available in home laboratories is X-ray diffraction. In the presence of heavy 5$d$ elements which scatter X-rays strongly the extremely weak signal from the oxygen positions, for example, is simply swamped. Absorption, particularly for diffractometers operating with a capillary/transmission geometry, is also a significant problem when 4$d$ and 5$d$ elements are present. Furthermore, techniques such as neutron diffraction that are often better suited to the characterisation of materials containing light elements such as oxygen alongside heavy transition metals (Fig. 5) have their own difficulties, such as the large absorption cross-section of natural Ir for neutron studies of iridates. The neutron scattering lengths and scattering/absorption cross-sections of different elements and isotopes are available in standard references,101,102 and the values in the first of these are also available in an online database maintained by NIST.103

This particular problem has been previously tackled for absorbing compounds by e.g. using high-flux instruments and optimising experimental geometry.104 More recently, we have achieved a significant breakthrough by developing the use of $^{191}\text{Ir}$, the lower-absorption isotope of the two naturally-occurring ones (37.3% $^{191}\text{Ir}$, $\sigma_{\text{abs}} = 954$ barns; and 62.7% $^{193}\text{Ir}$, $\sigma_{\text{abs}} = 111$ barns). Whilst $^{193}\text{Ir}$ is commercially available its use in neutron scattering had not been reported in the literature and no scattering length, which parameterises the scattering strength and is necessary for data analysis such as Rietveld refinement,
was available. Using a novel total scattering-based technique we determined the scattering length of $^{193}$Ir, allowing samples enriched in this isotope to be used in neutron scattering studies effectively.\textsuperscript{105} As such, high-resolution structural data (including detailed studies at high pressures that were effectively impossible previously\textsuperscript{79}) can now be obtained for iridates as a matter of course, allowing unprecedented access to the structural and spectroscopic information that is crucial for unravelling the origin of the exotic ground states and properties of these materials.\textsuperscript{77,105}

### 5.2 Determining composition

Accurately determining the composition of many of the materials of interest is also a significant challenge, and if sufficient care is not taken with this it is easily possible to assign physical properties to a mis-identified compound. Commonly-available techniques such as energy-dispersive X-ray analysis (EDX), inductively-coupled plasma analysis (ICP) and X-ray fluorescence spectroscopy (XRF) may not be sufficiently sensitive to light elements such as oxygen, lithium and other problematic light elements.\textsuperscript{110}

Oxygen content can be determined by structural refinements using neutron diffraction data, but this is only reliable when the data are not affected by other complications such as absorption, oxidation state variability and crystal structure uncertainties.

The difficulty in determining composition is exemplified by the cases of Na$_4$Ir$_3$O$_8$ and Na$_3$Ir$_3$O$_8$. These two compounds have closely related crystal structures that are almost indistinguishable in laboratory powder X-ray diffraction patterns. Although a continuous transition from one structure to the other is not possible,\textsuperscript{76} Na$_4$Ir$_3$O$_8$ was discovered during experiments aimed at growing single crystals of Na$_3$Ir$_3$O$_8$ as they occupy similar synthetic parameter space.\textsuperscript{106} Furthermore, Na$_3$Ir$_3$O$_8$ is amenable to Na-deficiency, allowing Na$_4-x$Ir$_3$O$_8$ to be synthesised – deliberately or otherwise – for a range of $x$.\textsuperscript{107} Therefore, a synthesis can easily produce a sample that is not of the expected composition but without that being realised.

Related to this is the characterisation of chemical (in)homogeneity, which is often not considered until late in the investigation cycle. Certain synthesis methods also present particular characterisation challenges. For example, it is difficult to determine if an ion-exchange reaction has progressed to completion if the starting content of the species to be exchanged has not been determined. Another challenge lies in ensuring that the target transition metal oxidation state is actually realised, or whether mixed valences and/or non-stoichiometry are present.

Examples of this include the double perovskites Ba$_2$CaIrO$_6$ (for which the stoichiometric Ir$^{6+}$ phase is only stabilised by high-pressure synthesis)\textsuperscript{108} and Sr$_2$CoIrO$_6$ (in which only 10\% of the iridium content is Ir$^{6+}$)\textsuperscript{109} and pyrochlores such as Bi$_2$Ir$_2$O$_7$.\textsuperscript{85} One recent development that may be particularly helpful in many cases is muon elemental analysis, which is sensitive to oxygen, lithium and other problematic light elements.\textsuperscript{110}

### 5.3 Oxidation states and mixed valence

When the spin-orbit-coupling-active species is of mixed valence, correct assignment of oxidation states to its different crystallographic sites is imperative if the interactions between sites are to be correctly understood. Osmates are a prime case as osmium has several stable oxidation states, which compounds the difficulties derived from the determination of composition. For example, Li$_{2.15}$Os$_{0.85}$O$_3$ is the first reported honeycomb osmate but, in contrast to its single-valence iridate analogue, samples of this material were found to be approximately 70\% Os$^{5+}$ and 30\% Os$^{4+}$.\textsuperscript{68} This has a significant effect on the magnetism as the mixture of nominally $J_{\text{eff}} = 3/2$ and $J_{\text{eff}} = 0$ ions creates disorder on

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**Fig. 5** (a) The relative X-ray scattering strengths of different elements are dependent on their atomic numbers, represented graphically here for Sr$_2$IrO$_4$ by the size of the atoms (Sr in green, Ir in blue, O in red). (b) An analogous representation for neutron scattering. Neutron scattering lengths are not dependent on atomic number, facilitating the location of light elements such as oxygen. (c) Simulated X-ray (upper) and neutron (lower) powder diffraction patterns of Sr$_2$IrO$_4$, with the intensities of the reflections normalised to the most intense in each. For both the X-ray and neutron simulations, the black pattern is for stoichiometric Sr$_2$IrO$_4$ whilst the red pattern is for a hypothetical analogue, Sr$_2$Ir$_{0.8}$O$_4$, that has an identical unit cell but all oxygen sites vacant (indicated by □). The asterisks indicate two superstructure reflections that are the result of a rotation of the IrO$_6$ octahedra. This rotation is crucial in determining the physical properties of the material but is almost invisible to X-rays.
the honeycomb spin network. Another intriguing example is the Os\(^{5+}/ Os^{6+}\) mixed oxide Sr\(_5\)Os\(_{3}\)O\(_{13}\), which shows complex temperature-dependent behaviour.\(^{111}\) In this material, one of the three osmium sites remains Os\(^{5+}\) across a wide temperature range, but the other two sites seem to show charge-ordering that varies continuously down to the Neél temperature, at which point full charge-order locks in alongside antiferromagnetic ordering. Precisely determining the oxidation states present and whether they are ordered, clustered or disordered is key to understanding the structures and magnetism of these materials but is non-trivial, often requiring facility-based techniques such as X-ray absorption spectroscopy (XAS).\(^{66}\) However, recent progress in oxidation state control by a variety of synthetic routes is showing that the effort can certainly be worthwhile.\(^{112,113}\)

5.4 Disorder

The effects of strong spin–orbit coupling are incredibly sensitive to structural perturbations, so if the virtuous cycle of experimental observations feeding into theoretical predictions and vice versa is to be maintained, all the real-world nuances of the materials under study must be accounted for. This includes the identification and characterisation of disorder. Disorder is commonly seen as a negative complication that drives materials away from desirable exotic ground states but it is ubiquitous, and in fact can sometimes stabilise states that would not otherwise be accessible – for example, the formation of the spin liquid state in H\(_4\)LiIr\(_2\)O\(_6\) is thought to be closely linked to hydrogen disorder.\(^{7,114}\) However, this can be compared to honeycomb Li\(_2\)RhO\(_2\), in which the effect of structural disorder on the magnetic ground state is to drive the formation not of a dynamic spin liquid but of a static glassy phase.\(^{115,116}\) Disorder is difficult to characterise structurally, particularly if diffraction alone is used as it only reveals the periodic structure of crystalline materials, meaning that disorder can be overlooked. Complementary local-structure probes such as muon spin relaxation (\(\mu\)SR)\(^{7,117}\) solid-state NMR\(^{118}\) and total scattering\(^{20,119,120}\) therefore have the ability to make significant contributions towards unravelling the complexities of real quantum materials.

5.5 Stacking faults

Stacking faults are a particular type of disorder that merit further elaboration. These microstructural defects are discontinuities in the stacking pattern of layered materials, and as such are relevant in the honeycomb and Ruddlesden–Popper materials that are of principal interest for strong spin–orbit coupling at this time. In fact, the presence of stacking faults in these materials provides a huge challenge for the determination of their structure–property relationships and the connecting of experiment and theory.

Firstly, even a low concentration of stacking faults complicates structure determination. As they affect both the shapes and intensities of the peaks in a diffraction pattern in a complex manner it can be difficult to disentangle the contribution of stacking faults from those of factors such as occupational disorder, hence data are often misinterpreted. Until recently simulation was the only way to easily quantify stacking faults in powder diffraction patterns (Fig. 6); however, recent developments in Rietveld refinement software are making the inclusion of stacking faults into structural models more feasible, thus allowing for improved refinements and the more accurate structure determination for significantly faulted materials.\(^{121,122}\) A rare trio of examples in the quantum materials field are the TOPAS refinements of H\(_4\)LiIr\(_2\)O\(_6\), Ag\(_3\)LiRu\(_2\)O\(_6\) and Ag\(_3\)LiIr\(_2\)O\(_6\).\(^{70,120}\) Single crystal diffraction can also be used to identify and study stacking faults, through the rods and streaks of diffuse scattering they generate in single crystal diffraction patterns (see e.g. ref. 66 and 123–125 for a range of examples).

Stacking faults can also influence the behaviour of the electrons in a material. Early studies of \(\alpha\)-RuCl\(_3\) produced conflicting reports of physical properties that were shown to be largely due to high concentrations of stacking faults in certain samples. A comprehensive diffraction study was required before awareness of this issue became widespread.\(^{66,126}\) Another clear example of the effect of stacking faults on physical properties may be found in the layered honeycomb iridates and ruthenates, a case in point being Ag\(_3\)LiIr\(_2\)O\(_6\), for which samples show ordering or an absence of ordering depending on the level of faulting.\(^{127,128}\)

Complementary techniques such as electron microscopy have the potential to accelerate our understanding of, and ability to control, stacking fault disorder.\(^{70}\) Furthermore, efforts to study and control stacking faults in \(\alpha\)-Li\(_2\)IrO\(_3\) and relatives are paralleled by current efforts in the energy materials field with regard to e.g. Li\(_2\)MnO\(_3\).\(^{129}\) Another example of the generality of the materials science issues faced by quantum materials researchers.

5.6 Challenges related to synthesis

Not only is the progress of materials discovery hampered by the characterisation challenges outlined above, but the synthesis of
materials with strong spin–orbit coupling is often non-trivial. Particular synthesis methods bring their own specific issues; for example, soft-chemistry techniques can be difficult to carry out reproducibly, and can also give products with very small particle sizes that complicate diffraction analysis through greater peak broadening. Single crystal growth is often very difficult and obtaining high quality crystals of large size is a particular issue, as techniques such as floating zone growth are often not suitable. Furthermore, single crystals often present different physical properties to polycrystalline samples that are nominally of the same material, and determining which form is displaying the intrinsic behaviour is not straightforward. The oft-made assumption that single crystals are ‘better’ than powders is not a reliable one.131,132

One other major challenge is polymorphism. Syntheses of Li2IrO3, for example, often do not reliably produce the target polymorph, even when a previously-successful method is followed.4,10,74 This can be related to reagent quality, as obtaining platinum group metal oxides of reliable composition may not be trivial – it is not unusual to find that batches of oxides contain fractions of the native element. Furthermore, when using isotopically enriched precursors for e.g. neutron scattering or NMR studies the range of available reagents may be reduced, forcing synthetic procedures to be re-designed. Control of stoichiometry is particularly difficult in soft-chemistry cases where the elemental composition makes chemical characterisation very difficult, and comparisons of the properties of samples prepared by different groups are limited if sufficient synthesis details and/or characterisation data are not included in publications. A further point of difficulty, particularly in the case of osmates but also not uncommon in iridates,133 is the stabilisation of the nominal oxidation state. High-pressure (multi-GPa) or high-oxygen-pressure methods are two approaches that have been used to achieve this and access the higher oxidation states of osmium, such as Os8+ without significant production of volatile and highly toxic OsO4 (which most osmates are unstable in ambient air).81,134,135 The stabilisation of single (or multiple controlled) oxidation states and further elucidating the chemistry of osmium in the solid state is a focus of current developments.81

6 Literature examples

The literature on iridates and other materials with strong spin–orbit coupling has focused on detailed explorations of the physics of their exotic electronic and magnetic states by property measurements, computation and the development of theory. In the understandable rush to discover and understand these exciting materials, detailed structural and chemical characterisation has often been neglected. This has led to conflicting reports of properties and ground states for particular materials, and theoretical models based on structures determined only by preliminary lab X-ray data. A prominent example is given by Na4Ir3O8. Four years after the properties of this material were first reported a detailed study demonstrated its air-sensitivity,138 yet in reports to that point samples were routinely handled in air. Inspection of published diffraction patterns now allows the likely partial-decomposition to be identified. Studies of the ruthenium analogue of this material, Na2RuO3, have also been beset by issues of air sensitivity, with recent reports suggesting an entirely different ground state in samples that were handled in an air-free environment (Fig. 7).132,137 Na4Ir3O8 also degrades on storage under ambient conditions, with sodium gradually being lost and the physical properties changing substantially.117,139,140 Such problems are not limited to materials containing volatile elements such as Na - in 2019 two groups reported property measurements of the K2O-type iridate La2IrO4,11 even though one found an anomaly at 3.5 K whilst the other did not, they cannot truly have been measuring the same material.141,142 The tendency of pyrochlore iridates to oxygen non-stoichiometry underlies another example. A large literature generated on ‘pure’ A2Ir2O7 materials claimed that these undergo a metal–insulator transition, but detailed investigations eventually showed that the stoichiometric materials do not undergo such a transition at all (Fig. 8).133 Even more extremely, the compound SrIrO3 was reported to exist in the Sr–Ir–O phase diagram but was later found to be simply the result of intergrowths of the n = 1 Ruddlesden–Popper phase Sr3Ir2O7 in the n = 2 phase Sr2Ir2O7.143 In other words, Sr2Ir2O7 as a distinct compound does not in fact exist, although at the time of writing a search of the Inorganic Crystal Structure Database144 suggests that it does, and the unwary user may inadvertently misidentify their material.

A final key difficulty in following the literature on these materials arises from the tendency to use physical property

![Fig. 7](https://example.com/figure7.png)
measurements for the characterisation of samples when chemical and structural methods are difficult. Whilst such measurements are essential for thorough characterisation, and in certain cases provide significantly more insight into sample quality than e.g. lab X-ray diffraction can,145 it should not be used as a primary method unless the applicability of the technique has been rigorously determined. The reported physical properties of many compounds have often been shown later to be non-intrinsic, instead being due to decomposed samples, impurity phases or incorrect compositional assignment. Such problems have led to significant issues of reproducibility and held back our understanding of these important materials.

7 Conclusions

In this Perspective we have provided a brief introduction to the many challenges and opportunities that quantum materials with strong spin–orbit coupling afford to materials chemists. The often extreme sensitivity of their physical properties to disorder, distortions and other chemical factors makes the input of chemists all the more essential, with potential payoffs through more rigorous investigations and a significant strengthening of the virtuous cycle between theory and experiment that could pave the way for critical developments in a range of fundamental and applied fields. These challenges may well be familiar, and our hope is that by recognising the complementary efforts currently being pursued in a wide variety of materials chemistry communities interdisciplinary exchange will be promoted and our understanding of these fascinating materials advanced.

Author contributions

AJB: conceptualisation, investigation, visualisation, writing—original draft, writing—review & editing. AK: investigation, writing—original draft, writing—review & editing. ASG: conceptualisation, supervision, funding acquisition, project administration, investigation, visualisation, writing—original draft, writing—review & editing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

ASG acknowledges funding through an EPSRC Early Career Fellowship EP/T011130/1. ASG and AK would like to acknowledge Dr T. Takayama and Prof. H. Takagi for their invaluable support, discussions and insights into materials with strong spin–orbit coupling (and many others) over the years. ASG acknowledges Dr K. S. Knight, Dr A. D. Fortes and Prof. W. I. F. David for insightful discussions.

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