REVIEW

Recent studies of cements and concretes by synchrotron radiation crystallographic and cognate methods

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The portfolio of available synchrotron radiation techniques is increasing notably for cements and pastes. Furthermore, sometimes the terminology is confusing and an overall picture highlighting similarities and differences of related techniques was lacking. Therefore, the main objective of this work is to review recent advances in synchrotron techniques providing a comprehensive overview. This work is not intended to gather all publications in cement chemistry but to give a unified picture through selected examples. Crystallographic techniques are used for structure determination, quantitative phase analyses and microstructure characterization. These studies are not only carried out in standard conditions but synchrotron techniques are especially suited to non-ambient conditions: high temperatures and pressures, hydration, etc., and combinations. Related crystallographic techniques, like Pair Distribution Function, are being used for the analysis of ill-crystalline phase(s). Furthermore, crystallographic tools are also employed in imaging techniques including scanning diffraction microscopy and tomography and coherent diffraction imaging. Other synchrotron techniques are also reviewed including X-rays absorption spectroscopy for local structure and speciation characterizations; small angle X-ray scattering for microstructure analysis and several imaging techniques for microstructure quantification: full-field soft and hard X-ray nano-tomographies; scanning infrared spectro-microscopy; scanning transmission and fluorescence X-ray tomographies. Finally, a personal outlook is provided.

Keywords: synchrotron techniques; quantitative phase analysis; imaging and mapping; binders; cement hydration; cement paste microstructure; C–S–H and C–A–S–H gels

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Nomenclature, Acronyms

Cement nomenclature: C=CaO, C=CO₂, S=SiO₂, S=SO₂, A=Al₂O₃, F=FeO₃, M=MgO, K=K₂O, N=Na₂O and H=H₂O. Therefore, for instance, Ca₃SiO₅ is C₃S, CaCO₃ is C(CO₂), and CaSO₄·2H₂O is CSH₂.

AFm ‘Al₂O₃–FeO₃–mono sulphate’ (set of phases)

Aft ‘Al₂O₃–FeO₃–tri sulphate’ (aka, ettringite)

ASR alkali silica reaction

CSA calcium sulfoaluminate cement

C–S–H calcium–silicate–hydrate amorphous gel

DTA/TGA differential thermal analysis/thermo gravimetric analysis

MIP mercury intrusion porosimetry

OPC ordinary Portland cement

PCE polycarboxylate ether

SCM supplementary cementitious materials

w/c water-to-cement mass ratio

AChX-nCT absorption-contrast hard X-ray nano-computed tomography

AChX-µCT absorption-contrast hard X-ray micro-computed tomography

ACsX-nCT absorption-contrast soft X-ray nano-computed tomography

ACsX-nM absorption-contrast soft X-ray nano-microscopy

BCDI Bragg coherent diffraction imaging

BL beam line

CAT computed axial tomography (aka, computed tomography)
1. A very brief introduction to Portland cement and concretes

The term cement (building material) may be used for almost any type of binder, with chemistry ranging from fully organic to totally inorganic. The oldest organic ‘cements’ were made from plant extracts like starch. The oldest pure inorganic cements were based on muds and subsequently on gypsum plasters. However, since the development of Portland cement about 180 years ago (http://en.wikipedia.org/wiki/Portland_cement), it has become the dominant binder used in concrete (http://en.wikipedia.org/wiki/Concrete) for construction. Concrete is a hierarchically complex hardened composite material formed from the mixing of water and aggregates (both fine and coarse) with Portland cement. Some other additions and admixtures may be added depending upon the type of concrete.[1] Portland cements are used as poorly soluble but highly reactive powders in water. The initial suspension, named as fresh cement paste, firstly undergoes setting and on hardening yields the cementitious matrix. The use of water for achieving the hardening results in the term ‘hydraulic binders’, to highlight that the reaction of cement with water yields the concrete which can harden even underwater.
Praised for its versatility, durability, fire-resistance, on-site casting and economic value, Portland concrete is receiving recognition also for its relatively low embodied energy compared to other building materials (see Figure 1). Furthermore, because of its use of local materials, by which there is reduction in energy and pollution costs associated with material transport, it is also an important asset. Annual worldwide Portland cement production is approaching 3 Gt.[2] World yearly consumption of concrete is well over 6 km$^3$; its magnitude may be, perhaps, better perceived by referring this value to every person in the planet: 2.5 tonnes per person per year. This is just second only to our consumption of fresh water. However and in spite of its universal use, Portland cements are one of the most environmentally contentious materials. Worldwide production of cement accounts for approximately 6% of the total anthropogenic CO$_2$ production.[3]

Research in cements and concretes has many facets ranging from the understanding of phase assemblage and microstructure to ensure extended durability, to reduce the CO$_2$ footprint of cements to mitigate the current man-increased green-house effect. An overview of the research carried out in cements can be gained from the scientific programme of the International Congress on the Chemistry of Cements which takes place every four years with the last edition taking place in October-2015 at Beijing (www.iccc2015beijing.org/).

Portland cement is manufactured by grinding the Portland clinker with the setting regulator that may be close to 4 wt% of gypsum. A typical Portland clinker chemical composition is close to 67 wt% CaO, 22 wt% SiO$_2$, 5 wt% Al$_2$O$_3$, 3 wt% Fe$_2$O$_3$ and 3 wt% of minor components. With this elemental chemistry, OPCs usually contains four major phases: 50–70 wt% alite, Ca$_3$SiO$_5$ or C$_3$S, 15–30 wt% belite, Ca$_2$SiO$_4$ or C$_2$S, 5–10 wt% tricalcium aluminate, Ca$_3$Al$_2$O$_6$ or C$_3$A and 5–15 wt% tetracalcium aluminoferrite, Ca$_4$Al$_2$Fe$_2$O$_{10}$ or C$_4$AF. Most cement compounds are not pure stoichiometric phases but they (may) incorporate many ions as extensively discussed in classical papers and books.[4]
Finally, some crystalline and amorphous hydrated phases should also be mentioned. AFm are a set of phases with crystalline layered structures. The AFm phase refers to a family of hydrated calcium aluminates based on the hydrocalumite structure, \( \text{Ca}_4\text{Al}_2(\text{OH})_{12}[\text{Cl(OH)}]6\text{H}_2\text{O} \). The archetype AFm phase is \( \text{C}_3\text{A} \cdot \text{CaSO}_4 \cdot 12\text{H}_2\text{O} \) or \( \text{Ca}_4\text{Al}_2(\text{OH})_{12}[\text{SO}_4]6\text{H}_2\text{O} \), known as Kuzelite, but Al can be partly replaced by Fe and \( \text{SO}_4^{2−} \) can be partly or fully replaced by \( \text{OH}^{−} \), \( \text{Cl}^{−} \), \( \text{CO}_3^{2−} \) and several other anions.\(^{[5,6]}\) In a similar way, AFt are a set of crystalline tridimensional framework compounds. By far the most common AFt phase is ettringite which has the stoichiometry \( \text{C}_3\text{A} \cdot 3\text{CaSO}_4 \cdot 32\text{H}_2\text{O} \) that can also be written as \( \text{Ca}_6\text{Al}_2(\text{SO}_4)_3(\text{OH})_{12} \cdot 26\text{H}_2\text{O} \). On the other hand, the principal binding reaction product of cement hydration is amorphous calcium–silicate–hydrate (C–S–H) gel, one of the most complex of all gels.\(^{[7]}\) Saturated C–S–H gel has the approximate formula \((\text{CaO})_{1.7}\text{SiO}_2(\text{H}_2\text{O})_4\), including liquid water between the particles, but the \( \text{Ca}/\text{Si} \) ratio and the water content evolves with time and it depends upon the composition of the starting binder.

2. A very brief introduction to synchrotron radiation and properties

Current synchrotron light sources are a particular type of circular particle accelerator designed and built to produce electromagnetic radiation of outstanding properties. These laboratories are user-oriented (both academic and industrial) Large Facilities where staff of very different backgrounds (photon scientists: physicists, chemists, geologists, biologists, physicists, etc.; and support scientists and technologists: accelerator physicists, mechanical/vacuum/computing/control engineers, technicians, etc.) work together with the final goal of offering the best possible service to the users.

A synchrotron light source usually contains three types of electron accelerators: (i) the linear accelerator (linac), (ii) the booster and (iii) the storage ring. The electrons are produced and initially accelerated in the linac. The electrons from the linac are injected into the booster where they are further accelerated before being kicked into the storage ring. However, some synchrotron light sources inject the electrons directly from the linac into the storage ring which is a circularly shaped accelerator where the kinetic energy of the electrons is kept constant and the desired electromagnetic radiation is generated. The readers are directed to two recent books in order to learn more about synchrotron light sources.\(^{[8–10]}\)

Figure 2 shows some of the most important components in a synchrotron facility from the storage ring of the sample to be studied. The design of beamlines varies tremendously depending on the methods (and wavelength range) to be used. For this reason only a very general overview of a beamline set-up is given in Figure 1. From the user’s point of view, everything starts at the photon source where the synchrotron radiation is produced by moving the electrons in the appropriate way within the magnetic device (see Figure 1). The SR produced by the source is pre-conditioned in the front-end before leaving the tunnel to enter into the optic hutch. It must be highlighted that both the energy range and the brightness of the SR strongly depend on the photon source but their discussion is out of the scope of this paper.

The unmonochromatized beam is called a white beam and must be ‘conditioned’ before it can be used to analyse the sample in the experimental hutch. It is not possible to review all possible optics devices but we can highlight just a few: (i) Mirrors to transport the beam (and sometime to focus it as well as to collimate and reject unwanted radiation); (ii) Monochromators to select a particular wavelength (or set of wavelengths) from the incoming beam; (iii) Focusing optic elements (if needed) to match the size of the beam to the requirements of the employed technique and sample; (iv) Slits, filters, diagnosis elements, etc. Monochromators can select a very narrow energy bandwidth (usually \( \Delta E/E = 10^{-4} \) or smaller) for instance using a double-crystal-monochromator and this radiation is called monochromatic, but it can also select a
much wider set of wavelengths (f.i. $\Delta E/E = 1–2\%$ with a multilayer monochromator) for having higher flux and this type of radiation is commonly named pink beam.

The conditioned SR is then used in the experimental hutch to analyse the sample. The appropriate sample answer is recorded in the detector that for combined studies can be more than one. Most of the synchrotron BLs are in the hard X-ray domain (roughly between 4 keV and 50 keV) or soft X-ray domain (roughly between 200 eV and 4000 eV). The optics (and sample set-ups) of soft and hard X-ray BLs are quite different as low-energy implies vacuum which places a lot of restraints in many experiments. It must also be mentioned that IR spectro-microscopy BLs are also starting to play a role in cement studies.

The sample (as well as some optics elements) must be positioned with high mechanical accuracy. Hence, high resolution mechanical and piezoelectric stages are commonly used. Custom-made sample environments also allow in-situ experiments. This term should be understood to include, but not be limited to, time-resolved works (under the influence of external parameters like chemical gradients (water, CO$_2$, etc.), temperature, pressure, etc.; or combinations!) usually in the subsecond to hour timescale which is the relevant scale for building material experiments.

The main properties of SR that benefit building material studies are summarized in Table 1. The outstanding properties of SR have encouraged their uses in many researches related to cements. Comprehensive information regarding the uses of synchrotron can be found in recent books.[7–10] Very useful properties of SR like its intrinsic time structure (that allows following
Table 1. Main properties of synchrotron radiation of interest for building material studies.

<table>
<thead>
<tr>
<th>Property of the SR</th>
<th>Features / uses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extreme brightness and flux (combined with high collimation)</td>
<td>• Very high signal-to-noise ratio, allowing high quality data&lt;br&gt;• Fast data collection, including kinetics studies (typ. &gt; 1 ms)&lt;br&gt;• Complex sample environments that absorb radiation, for instance in high temperature, high pressure studies, etc.</td>
</tr>
<tr>
<td>Energy tunability (from infrared to very high-energy X-rays)</td>
<td>• Chemical sensitivity through the absorption edges&lt;br&gt;• High-energy minimizes absorption in complex sample environment experiments&lt;br&gt;• High energy allows to access to large momentum transfer values</td>
</tr>
<tr>
<td>Beam size tunability (from tens of mm to tens of nm)</td>
<td>• Enhance contrast in soft X-ray imaging, f.i. ‘water-window’&lt;br&gt;• Enable a very large range of imaging techniques from cm in paleontology to nm in nanotechnology&lt;br&gt;• Matches the size of the beam to the experiment requirements&lt;br&gt;• Allows to design sample environments with large degree of freedom</td>
</tr>
<tr>
<td>Parallel geometry</td>
<td>• Improved full-field reconstruction for imaging&lt;br&gt;• Minimize the errors in powder diffraction&lt;br&gt;<strong>Limited magnification of X-ray beams</strong></td>
</tr>
<tr>
<td>Partial transverse coherence</td>
<td>• Used in phase contrast imaging techniques&lt;br&gt;• Allows coherent diffraction imaging</td>
</tr>
<tr>
<td>Stability</td>
<td>• High resolution and high reproducibility</td>
</tr>
</tbody>
</table>

Notes: Characteristics in italics note possible drawbacks. Polarization properties and temporal resolution due to the electron bunches are not included as they have not been used in cement studies.

Also to the best of my knowledge there is no general review about the uses of synchrotron radiation for characterizing cement binders. I am aware of three focused reviews and they are discussed at the beginning of the next section. However, there are general reviews about the uses of synchrotron radiation applied to other fields like cultural heritage.[11,12] In particular, I found the second, very general, work [12] extremely enlightening. Furthermore, some techniques not treated here (because they have not been applied to building materials) are gathered and discussed there. It must also be noted that the evolution of synchrotron radiation, including its growing importance in crystallography, has also been reviewed [13] and also the starting uses of XFEL.[14]

In this review article, I highlight recent uses of SR applied to a better/deeper characterization of building materials through selected examples. I restrict the discussion to the last decade and references to previous works are only carried out for a few selected cases. I**convey my apologies** Q5 to these authors whose work(s) are not referenced here but it would not be possible to account for all works dealing with SR and building materials. I have selected examples to show the wide range of applicabilities, and this work is not intended to be a thorough work compiling all references but the updated summary of some key developments.

3. Uses of synchrotron radiation for cements, pastes and concretes

There are three recent reviews devoted to the characterization of building materials by synchrotron characterization techniques. Prof. Monteiro’s group has reported two reviews [15,16]...
Figure 3. Summary of the different synchrotron techniques employed for characterizing cements and pastes. The interaction of the SR can be elastic where momentum is transferred (yielding the scattering techniques) or inelastic where energy is exchanged (resulting in the spectroscopic techniques). Furthermore, there are many imaging tools profiting from these interactions as well as others like absorption. Moreover, time-resolved studies allow to study kinetics of chemical reactions which are key to follow the evolution in pastes in different time domains. (Acronyms have been defined in the beginning of this paper.)

mainly focused on their own activities covering some techniques: (i) full-field soft X-ray nano-tomography; (ii) scanning transmission X-ray microscopy; (iii) scanning X-ray microdiffraction microscopy; and (iv) high pressure X-ray powder diffraction. On the other hand, Prof. Provis’s group has also reported one review,[17] mainly focused on their own activities arising from BL-based studies (synchrotron and neutron), dealing with the multiscale understanding of the microstructure and chemistry of geopolymer binders. In this case, the reviewed synchrotron techniques were: (i) scanning infrared spectro-microscopy; (ii) scanning fluorescence X-ray microscopy; and (iii) X-ray nano- and micro-tomographies.

With the information that I have gathered, Figure 3 displays a scheme containing the different SR techniques that have been employed for the characterization of cements and pastes. Figure 4 schematically represents the set-ups used for the main characterization techniques discussed here. As it can be seen from both figures, many synchrotron techniques have been used for the characterization of building materials.

These techniques range from scattering for the analysis of phases as well as phase evolutions to spectroscopic tools for a better characterization of chemical species and elemental compositions. Spatially resolved information can be obtained from a number of approaches. In this context, it is appropriate to distinguish between mapping (or raster-scanning) which consists in the sequential measurement of data from adjacent regions of a sample achieved by moving each region into the photon beam, and (full-field) imaging where an image of the sample is recorded onto an array detector where the signal coming from each region of the sample is measured at each pixel. In addition to the mapping and imaging approaches, hybrid approaches are also being developed. There is no doubt that spatially resolved data are very important in building material characterization as they are being used to reveal the complex hierarchical microstructure of the hydrated pastes without alteration, which is not the case for other characterization techniques like electron microscopies.
For mapping and combined approaches, the synchrotron beam must be focused on a tiny spot of the appropriate size (see Figure 4). However, focusing the synchrotron X-rays is challenging since the optical index $n$ of most materials is close to unity for these energies. This is being circumvented by several approaches and Figure 5 gives the four main types of focusing optics. For a deeper insight, the reader is directed to a recent review [18] where the progress in the development of hard X-ray microscopy techniques for material characterization at the nanoscale was discussed.

Reflective optics (KB mirrors and capillaries) are achromatic, allowing stable beam focusing even when the energy of the incoming X-ray beam varies. Therefore, they are suited to X-ray
absorption experiments. Some mirrors have a fixed bending, while others use mechanical benders to change the focal spot size (or position). For hard X-rays, the most common configuration used for focusing is the KB system, in which one mirror focuses the beam in the horizontal and a second mirror focuses in the vertical direction. Both mirrors can be moved (and bent) independently of each other, giving additional flexibility to the system.

Diffractive optics use nanostructures to focus the beam. They are called Fresnel lenses and are very effective for soft X-ray energies. At hard X-ray energies, it becomes increasingly difficult to manufacture nanostructures that provide the necessary phase shift although big advances are taking place.[18] FZP optics is chromatic which means that the focal points move with the energy. Therefore, they require adjustments when varying the energy (e.g. µ-XAS in STXM experiments).

Refractive optics work profits from the index of refraction being smaller than one, and so they look like inverted lenses. Since the index of refraction is quite close to 1.0, many lenses are needed (often more than 10) to provide suitable refraction power. Most lenses are made out of beryllium, although other materials are also used. The main disadvantages of refractive lenses are their chromaticity and the low flux that they provide at soft X-ray energies. For hard X-rays and limited energy ranges, their performances are competitive.

3.1. Diffraction for characterizing crystalline phases (long-range order)

SR X-ray diffraction is used to determine the crystal structures of phases present in building materials which develop long range periodic order. These phases can be fine powder, and then the suitable technique is XRPD, or single crystal, and then the right technique would be SCXRD, (see Figure 3). Now, microdiffraction is becoming important as crystals considered powder a decade ago (sizes ranging 1 to 10 µm) can now be studied as single crystals using microdiffraction techniques. To the best of my knowledge, microcrystal synchrotron X-ray diffraction has still not been used to solve crystal structures in the cement field but has been used for imaging samples. This application will be treated in sections 3.7.3.4 and 3.7.3.5.

Examples of crystal structures studied by SXRPD are numerous. I can highlight the M3-superstructure of C₃S from synchrotron and neutron powder diffraction,[19] the T₁ and T₂ polymorphs of C₃S,[20] the T₃ superstructure of C₅S,[21] an improved structural description for gypsum,[22] cell dimensions and compositional details of mechanochemically prepared C–S–H gels [23] and approximate structural descriptions of 1.0 C–S–H and 1.5 C–S–H gels,[24] wet and dried ettringites,[25] Fe–Si-hydrogarnets (hydroandradite),[26] the pseudo-cubic structure of doped ye’elimite [27] and several AFm-type phases including Fe-hemicate, [28] hemicarboaluminate and carbonated hemicarboaluminate [29] and the double-anion Kuzel salt, Ca₂Al(OH)₆(Clₒ.₅[SO₄]ₒ.₂₅·2.₅H₂O).[30]

3.2. Techniques for characterizing amorphous phases

If the samples are crystalline (periodically ordered in the long range), the suitable technique to characterize the (crystal) structures is diffraction. However, very interesting samples (for instance C–S–H gel) are not crystalline and so alternative methodologies have to be employed to study their local (short-range) structures (see Figure 3). From the set of suitable techniques for characterizing local structures with SR, total scattering techniques and XAS are the most widely used approaches.

3.2.1. Total scattering techniques for studying short-range order

Total scattering methods (often referred to as PDF or radial distribution function) have been used since the thirties of the past century to get insight into the disordered structure of liquids and
amorphous materials. The experimental set-up essentially comprises a collimated, monochromatic beam impinging on a sample and the scattering data are nowadays recorded in a suitable 2D detector. Very short wavelengths (and so highly energetic X-ray photons) allow access to large momentum transfer values as well as to reduce experimental artefacts. The PDF method involves the sine Fourier transform of the measured structure factor over the widest possible momentum transfer range, providing a direct measure of the probability, G(r), of finding an atom surrounding a central atom at a radial distance. The weighting factor of each atomic species is scaled by their concentration and by the number of electrons (for X-rays). The advantage is that average structural information may be obtained when no (sharp) Bragg peaks are present in the measured diffraction pattern and all the scattering data are taken into account (including diffuse scattering), hence the name total scattering. A work that was recently focused on has reviewed the use of total scattering methods in building materials to characterize amorphous and poorly crystalline phases.[31]

The PDF technique was employed to study the local structure of synthetic C–S–H(I) showing nanocrystalline ordering with particle diameter close to 3.5 nm which are similar to a size-broadened 1.1 nm tobermorite crystal structure.[32] The C–S–H component in hydrated tricalcium silicate was found to be similar to C–S–H(I); only a slight bend and additional disorder within the CaO sheets were required to explain its nanocrystalline structure. In a subsequent work,[33] these researchers studied different C–S–H samples with varying Ca/Si ratios (between 0.6 and 1.8). The PDF analysis results suggested that the C–S–H structure evolves from tobermorite-like to jennite-like as a function of the increasing Ca/Si ratio. Evolution of these short- and medium-range order structural characteristics was associated with the alteration of the Ca–O layers and silicate depolymerization.

The PDF approach using SR has also been employed to study several other amorphous (or ill-crystalline) phases in cement research including ASR gel,[34] silica fume,[35] metakaolin[36,37] and geopolymers.[38–40]

### 3.2.2. X-ray absorption techniques for studying short-range order and chemical states

Synchrotron-based XAS techniques provide element-selective complementary information about the local structure and chemical speciation of the selected element by fine-tuning the X-ray photon energy to the absorption edge of the absorber. Most frequently used XAS techniques are XANES and EXAFS. The XANES region covers up to \( \approx 50 \text{ eV} \) above the absorption edge and it is mainly used to study the oxidation state of the absorber atom and for fingerprinting on the basis of a comparison of reference spectra with the unknown spectrum of the element under study. The first coordination shell (e.g., tetrahedral vs. octahedral coordination) can also be discriminated from the XANES data under some conditions. The EXAFS region can span up to 1 keV above the absorption edge and it can provide information about the local structure (i.e., type of neighbouring atoms, bond length and coordination numbers, second neighbours) of the studied atom. In this case, a deeper mathematical treatment of the recorded signal is needed. Chiefly, in addition to crystalline materials, XAS can be used to study amorphous solids with elements at very low concentrations (detection limit variable but usually lower than a few tens of ppm). In addition, XAS techniques combined with a micro X-ray beam can provide spatially resolved information on the micrometer scale about the speciation of the studied element(s); examples of this mapping approach are given in section 3.7.3.3.

Selected examples of XAS techniques applied to building materials are discussed next. The iron incorporation in hydrated cement phases is always an issue as it can replace a number of elements including aluminium. Hence, iron in carbonate containing AFm phases was studied by SXRPD and EXAFS around the Fe K-edge (\( \sim 7120 \text{ eV} \)).[28] In a latter study,[41] Fe K-edge EXAFS data were used for studying iron along OPC hydration from very early ages (hours)
up to one year, with hydration having taken place at 20°C and 50°C. Disregarding anhydrous phases, ferrihydrite was detected during the first hours of the hydration process. After 1 day, the formation of Al- and Fe-siliceous hydrogarnet was observed, while the amount of ferrihydrite decreased. In a subsequent work,[42] it was shown, also from Fe-edge EXAFS data, that at long hydration ages (many years), and upon selective dissolution of the pastes, independent of the chemical compositions of cements, formation of the mixed Fe–Al siliceous hydrogarnet is thermodynamically favoured. Other work used Si K-edge (∼1850 eV) XANES data to study the silicon environments in two hydrated samples: an OPC paste and a related paste with 35 wt% of slag as SCM, which showed no major differences.[43] Ca K-edge (∼4400 eV) EXAFS spectra were obtained, in addition to other complementary techniques, to investigate and characterize the structural disorder in C–S–H samples with Ca/Si ratios similar to tobermorite.[44] It was shown that C–S–H gels with Ca/Si ratio of ∼0.8 are structurally similar to nano-crystalline turbostratic tobermorite. Finally, XANES spectra of Fe K-edge, Ca K-edge and S K-edge (∼2480 eV) have been very recently used for investigating the nature of damaged interior walls.[45]

3.3. Diffraction for quantifying crystalline phases and in many cases phase evolution(s)

XRPD in general, and SXRPD in particular, is being widely used for quantifying the crystalline (and overall amorphous) phase contents. There are several approaches to analyse the powder diffraction patterns, and thus to derive the phase assemblages. However, the Rietveld method is by far the most commonly employed methodology. Recently, three review articles have been devoted to RQPA of anhydrous Portland cements,[46] Portland cements, blended Portland cements and their hydration products,[47] and building materials in general.[48] In these review articles, results derived from SXRPD analyses were mentioned but the uses of SR were not thoroughly discussed. The main uses of RQPA for giving information about building materials are compiled in Figure 6 and as it can be seen, there are many applications. Most of these uses are attained from laboratory XRPD, but SXRPD provides higher resolution data which are very important for studying very complex samples. SXRPD has also been used to validate analysis protocols later implemented with laboratory data. Furthermore, SXRPD allows very demanding experimental conditions that can hardly be attained with laboratory data (f.i., high pressure studies, very high temperature analyses, etc.). In the next subsections I will focus on the uses of SXRPD for phase quantification (including phase evolution) under a large variety of conditions.

3.3.1. Quantitative phase analysis of anhydrous building materials

The initial work on RQPA of a Portland clinker was published in 1993[49] but it was not until 2001 that SXRPD was applied to the RQPA of OPCs.[50] In this work, the main benefits of using synchrotron radiation were discussed: (i) very high resolution (by using a crystal analyser system in the diffracted beam) for polymorph identification/quantification; and (ii) high-energy radiation (E = 28.2 keV, or 0.44 Å) for sampling a large amount of clinker and thus to have accurate intensities that yield accurate phase quantification. In a subsequent work, these authors focused on the use of SXRPD for establishing the accuracy in RQPA of OPC cements.[51,52] Other authors also compared RQPA results from SXRPD, from neutron powder diffraction data and from laboratory data.[53] The very high resolution data obtained at synchrotron powder diffractometers allowed to establish alite phase coexistence in many clinkers.[54,55] Finally, SR was also employed for the quantification of other cements like calcium aluminate cements[56] and belite cements.[57]

Figure 7 shows a typical Rietveld plot as a visual outcome of a quantitative phase analysis for an OPC clinker.[47] The main peaks due to a given phase are labelled. Furthermore, this figure compares the Rietveld plots for data taken with three powder diffractometers for the same clinker.
3.3.2. Quantitative phase analysis of hydrating binders

SXRPD has been used more often for analysing hydrating binders than for characterizing anhydrous clinkers/cements. This is a consequence of the more complex nature of the hydrating building materials where many crystalline phases (and some amorphous ones) can coexist/develop. Therefore, the high-resolution data of SXRPD coupled with the larger penetration of high energy radiation (which leads to better particle averaging and so more accurate powder diffraction intensities) are two key assets. There is no need to say that fast powder diffraction data can be recorded (which depends upon the used detector system). Fast kinetics can be recorded from sub-second [58] to hours – time resolution. Firstly, I will focus on some selected studies of single phase hydration (or artificial mixtures of crystalline phases) and, secondly, I will review key works dealing with hydration of cements.

The hydration of C3A and C4AF at variable temperatures, between 25°C and 170°C, was studied by SXRPD. The diffraction pattern evolutions were reported including the identification of intermediate phases but phase quantifications were not carried out.[59] In a related work, energy-dispersive SXRPD was used to study the hydration of C4AF and CSH2 at temperatures ranging between 30°C and 150°C. The diffraction peak evolution was again followed, including intermediate phase identification, but phase quantification was not carried out.[60] In a step forward, Rietveld methodology was employed for quantifying the phase development of C3A+CSH0.5 samples with and without superplasticizer. Ettringite was quantified and also its texture evolution was also reported.[61] Rietveld methodology has also been employed to study the role of ye'elimite polymorphism in hydration reactions (and kinetics) by addressing the standard
Figure 7. Selected region of the Rietveld plots (three different experimental set-ups) used for quantitative phase analysis of the same commercial Portland clinker. Data were collected at: ID31 diffractometer of ESRF synchrotron (top), a laboratory diffractometer with a Ge(111) primary monochromator, CuK\(_{\alpha_1}\), (middle), and a laboratory diffractometer with a graphite secondary monochromator, CuK\(_{\alpha_{1,2}}\), (bottom). Please note the different resolution in the recorded data. Peaks arising from different phases are labelled. Reprinted from reference 45 with permission from the Mineralogical Society of America.
variables: water-to-ye’elimite ratio and sulphate sources.[62] Very recently, the hydration of C₄AF has been followed by RQPA of SXRPD at room temperature in different conditions: (i) without gypsum; (ii) with gypsum; (iii) with gypsum and stoichiometric ye’elimite (orthorhombic); and iv) with gypsum and doped ye’elimite (pseudo-cubic). Different behaviours have been observed and discussed including a delayed C₄AF hydration by the presence of ye’elimite, that it is even dependent of the doping content of the ye’elimite phase.[63] To end this section, I would like to highlight that SXRPD can also be used to develop/validate methods for analysis of key phases. This is the case of two reports [64,65] where a method for quantifying portlandite was developed.

Concerning the hydration of cements, a first key paper was devoted to monitoring the influence of cellulose ethers on hydration kinetics of cements with a time resolution of one minute with the first dataset taken only 90 seconds after water mixing.[66] In this work, the ettringite crystallization was followed but quantitative phase contents were not derived. Later, Rietveld methodology was used to extract the phase contents in hydrating OPC pastes with time resolution of minutes.[67] This hydration study was carried out in the presence of additives such as superplasticizers and setting accelerating agents. A similar experimental set-up and analysis methodology were followed to study the hydration of belite and active belite cements.[68]

The hydration of eco-cements, including phase content evolution by Rietveld methodology, has also been analysed for aluminium-rich belite sulfoaluminate cements,[69] active sulphobelite cements [70] and active sulphobelite cements at very early ages.[71] SXRPD coupled with the Rietveld method can also be used to characterize the long-term leaching behaviour of concretes by quantifying portlandite dissolution at different hydration times.[72] These results were compared to those obtained by DTA/TGA.

It is also important to discuss some hydration studies where SCM were blended with OPCs. Time-resolved SXRPD coupled with the Rietveld method was used to quantify the early-age hydration and pozzolanic reaction in OPC blended with natural zeolites. One key conclusion of this study was that the addition of natural zeolites accelerates the onset of C₃S hydration and precipitation of CH and Aft.[73] In a subsequent work, these authors enlarged the study by analysing the early-age hydration of OPC blended with micronized zeolitite and quartzite powders.[74]

To conclude this section, not only has OPC been studied but other cementitious systems have been analysed as well. For instance, SXRPD, in combination with other characterization techniques, was used to determine the reaction products of alkali-activated Class C fly ash-based aluminosilicate materials.[75] Finally, SXRPD and other characterization techniques were used to follow the phase developments and microstructural characteristics of geopolymers obtained by alkaline activation of natural pozzolans.[76]

### 3.4. Variable-temperature diffraction to characterize phase evolution(s) and reactions

SXRPD have been also used to characterize cements at quite high temperatures. The main objectives are twofold: on the one hand, to characterize the formation and decomposition products in cement systems and, on the other hand, and at higher temperatures, to study in-situ clinkering processes. These two uses are discussed next.

#### 3.4.1. In-situ thermal formation and decomposition studies at moderate temperatures

I will follow the approach already exercised in previous sections. First, I will discuss some examples of the decomposition of single phases of interest in cement chemistry and, second, I will give examples of thermal decomposition of cement pastes.

Calcium sulphates are key phases for regulating the setting behaviour of cement through the control of the soluble sulphate content at very early ages which is of paramount importance for
aluminate reactivity. Therefore, I highlight the work studying the formation and reactivity of five calcium sulphates studied by SXRPD and the Rietveld method.[77] More complex phases can be studied in-situ, both formation and decomposition. As an example: a slurry of synthetic ettringite was heated to form AFm-14 (and bassanite), at about 115°C, on saturated water pressure. On further heating, AFm-14 decomposes to yield hydrogarnet above 210°C. A full transformation pathway was reported by time-resolved SXRPD using a capillary cell.[78] The thermal decomposition of other cement phases has also been studied by SXRPD, like 3Mg(OH)2⋅MgCl2⋅8H2O, the main phase of Sorel cement.[79]

High-temperature SXRPD has been used to follow the thermal transformations of special cements or mixtures. This is the case of cement-asbestos decomposition studies.[80,81] The highest reported temperature was 900°C as the authors used an air blow heating system with the sample within quartz capillaries. This type of preliminary studies is important for safe treatments of asbestos in novel industrial reactors.

3.4.2. In-situ clinkering studies at very high temperatures

High-temperature SXRPD can also be used for the in-situ study of the reactions taking place in the clinkering process. These are complex experiments as the temperature can be as high as 1400–1500°C with important fraction of the sample melt, and so the capillary must be inert towards this very reactive environment. Using a halogen furnace that can heat up to 1600°C, the clinkering reactions of active belite clinkers were reported using platinum tubes.[82] The selected wavelength was very short, \( \lambda = 0.30\text{Å} \), to go through the highly absorbing Pt capillary. Rietveld methodology was successfully applied to the recorded data to obtain quantitative phase analyses. In a subsequent work, the same experimental conditions were used to study the clinker reactions of iron-rich belite sulphoaluminate cements, aka sulphobelite.[83] In this study, the high-temperature reactions were established and quantified and the role of borax for activating these cements was also characterized.

Figure 8 shows a sequence of steps to carry out this type of in-situ experiments. First, the precalcined raw material mixtures(s) are loaded into Platinum tubes.[82] A precalcining step is needed to release the CO2 and avoid overpressures in this type of experiments, where the sample is sealed within the capillary. Furthermore, expensive Pt capillaries/tubes are needed because the heated mixtures contain a relatively large fraction of highly reacting liquid (above 1200–1400°C), and so inexpensive ceramic tubes/capillaries cannot be used. The Pt tube is isolated from the diffractometer by an MgO ceramic bar which is bonded by a refractory glue (dark powder between Pt and MgO in Figure 8, top left). The cell is assembled, mounted within the diffractometer axis and the temperature is coarsely controlled by the calibrated voltage applied to the halogen lamps. Then, the SXRPD patterns are collected at (approximate) temperatures. The temperature can be accurately known from the values of the refined unit cell parameters of platinum, as its thermal expansion is well known. Finally, RQPA is carried out based on the pattern regions which contain diffraction peaks from the sample and are not severely overlapped with the Pt diffraction peaks. Under these very tough conditions (high temperatures with samples containing about 30 wt% of a highly reactive ceramic melt), the SXRPD patterns were of sufficient quality to allow appropriate quantification of the phase evolution.[82,83]

3.5. Diffraction under pressure to characterize phases and chemical reactions

In this section, I will review some SXRPD studies of single phases or cements under pressure. I have subdivided this part into three sections depending upon the level of pressure reached in the different approaches. Hydrothermal reactions take place at relatively low pressure, usually
lower than 20 bars (2 MPa). There are capillary cells that can withstand up to 200 bars (20 MPa) and these works are reviewed in the second section. Finally, a DAC can withstand more than 1 Mbar (100 GPa) and is normally used to study the high pressure behaviour of hydrated cement phases to derive mechanical properties like the bulk modulus. These works are reviewed in the last subsection.

3.5.1. Chemical reactions at relatively low pressures in capillary cells (hydrothermal conditions)

The formation reactions of AFt and two types of AFm phases were studied by SXRPD and hydrating different calcium aluminate phases in the presence of gypsum.[84] The reactions were investigated with the samples loaded in quartz capillaries, and the maximum internal pressure was 17 bars, which was obtained by supplying N₂ gas. The studied temperature range was 25–170°C, which was obtained by heating the capillary with a hot air flow system. In a second work,[85] these authors studied the hydrothermal transformations of the calcium aluminium oxide hydrates, CaAl₂O₄·10H₂O and Ca₂Al₂O₅·8H₂O, to yield crystalline Ca₃Al₂(OH)₁₂, and
where intermediate phases were identified. On the other hand, instead, a capillary cell, an autoclave cell with Be windows was designed and built, allowing the study of cements by SXRPD up to 12 bars and 200°C.[86] This autoclave cell was initially used for studying the hydrothermal formation of tobermorite under different conditions [86,87] and later it was used for investigating the phase evolutions during the production of autoclaved aerated concretes.[88–90]

3.5.2. Hydrating reactions at medium pressures and temperatures

Several hydrothermal reactions take place at pressures up to 20 bars, but there are other conditions where (much) higher pressures apply. For instance, deep oil-well cement slurries can be subject to pressures of up to 1 kbar. Therefore, to study cement hydration in the range of 100–200 bars it is very interesting for several applications including oil-well cements which surround metal oil-well liners to form a gas tight seal between the bore wall and the liner. For this type of studies, a cell was designed and built for SXRPD data collection, in optimum conditions, up to 500 bars and 200°C.[91] Later, this cell, which performs best with sapphire capillaries, has been used for studying a number of hydrating cement slurries including: (i) Class A and H oil well cements mixed with variable amounts of CaCl2 for accelerating the hydration [92]; (ii) Class H oil well cements mixed with silica flour, silica fume and a natural zeolite which are used to improve the mechanical properties of the binder [93]; (iii) White cement, class G and class H oil well cements, without additives, for analysing the reaction kinetics of the main phases [94]; and (iv) C3S in the presence of several retarders to counterbalance the accelerating behaviour of pressure.[95] These authors have also developed a cell to allow simultaneous measurement of SXRPD data and ultrasound shear-wave reflection data from cement slurries at variable temperatures and pressures.[96]

3.5.3. Phase characterization at very high pressures in diamond-anvil-cells

The pressures employed in this type of studies (dozens of GPa equivalent to hundreds of kbars) are much higher than what could exist in cement pastes (mortars or concretes); however these works are important to derive mechanical properties of the hydrated phases as well as to study the stability of cement phases which may help to understand reaction mechanisms. The pressure transmitting medium is important as it should not induce any structural modification to the studied phase and it must be easy to work with. Silicone oil as well as methanol/ethanol mixtures have been used for studying hydrated cement phases at high pressures in DAC systems.

Figure 9 shows a scheme with components of a DAC. There are different types of DACs, but its discussion is out of the scope of the present work. When the sample is loaded within the DAC, the set-up is placed in the diffractometer and the micro-focused beam interacts with the sample. In order to reach higher pressures, smaller samples are employed. The diffracted beam is recorded in an appropriate 2D detector (commonly a CCD detector). Forces are varied with the pressures being measured, commonly, from the fluorescence signal of a ruby chip. Finally (from the data collection point of view), the 2D patterns are radially integrated to transform them to 1D patterns (see Figure 9) that can be analysed with any Rietveld package.

The DAC approach has been used to study the behaviour of ettringite up to 6 GPa. The two main outputs of this study were the isothermal bulk modulus of ettringite, 27 GPa, and its transformation to an amorphous phase at pressures above 3 GPa.[97] The bulk modulus of gel C-S-H is of utmost importance as it is the main binding component in OPC mortars and concretes. However, as its chemical composition can vary, to report a single number is complicate. In an initial work, two samples were studied, synthetic C–S–H(I) and another gel obtained from the hydration of alkali-activated slag. The powder diffraction patterns were collected up to 4 GPa and the bulk modulus for the two gels were very similar, ≈35 GPa, showing that Al substitution in
Figure 9. Schematic representation showing the experimental work flow in a high-pressure experiment using DAC. (Top right) Scheme of a diamond-anvil-cell with its different components including the ruby chips commonly used to measure the applied pressure. The (micro)focused SR X-ray beam is diffracted by the sample and the high-pressure pattern is collected in a suitable 2D detector. (Top left) A CCD 2-D pattern is collected at a given pressure. Then, the 2D pattern is integrated to yield the typical intensity versus angle 1D pattern. (Bottom) Finally, several data sets are collected at different pressures and the appropriate information is derived (f.i., unit cell variation, crystal structures, amorphization processes, bulk modulus, etc.).

C–S–H(I) does not significantly affect the bulk modulus of C–S–H(I).[98] On the other hand, the two crystalline phases more similar to the ill-crystalline C–S–H gel are 14Å-tobermorite and jennite. Hence, SXRPD data for 14Å-tobermorite were measured up to 5 GPa and the obtained bulk modulus was 47 GPa.[99] A similar study for jennite, measured up to 6 GPa, yielded a bulk modulus of 64 GPa.[100] The isothermal bulk modulus of aluminium-substituted 11Å-tobermorite from relict lime clasts of 2000-year-old Roman seawater harbour concrete was also measured, yielding $K_o = 55 \text{ GPa}[101]$. Other works reported the bulk modulus for selected samples of interest in cement chemistry like stratalitie: $K_o = 23 \text{ GPa}$, hemicarboaluminate: $K_o = 15 \text{ GPa}$, monocarboaluminate: $K_o = 54 \text{ GPa}$ and hydrogarnet: $K_o = 70 \text{ GPa}$.\[102,103\]

X-ray diffraction is the standard method for measuring the unit cell volume variation with the pressure of crystalline materials. For a pressure range where the composition of the phase is constant, this directly translates into the density variation with pressure. Using diffraction to determine the density variation of non-crystalline materials is not straightforward, so alternative methods are being investigated. Recent advances have allowed XAS data, taken for an amorphous material loaded in a DAC, to be used to characterize the density variation with pressure. This approach has been successfully used in cement chemistry.[104] The elastic properties of an ASR gel were studied by XAS and Brillouin spectroscopy measurements. XAS was used to determine the density of the gel as a function of pressure, yielding an isothermal bulk modulus
of 33 GPa. Brillouin spectroscopy was applied to measure isentropic bulk (24.9–34.0 GPa) and shear moduli (8.7–10.1 GPa) of the gel.

Finally, SXRPD data collected in DAC as a function of pressure have also been recorded for anhydrous phases of interest in cements. Experimental data were recorded for C$_3$A which yielded an isothermal bulk modulus of 110 GPa which was compared to the results from theoretical calculations.[105] Other elastic properties were also measured and compared to the theoretical results, a second example being ye’elimite. The reported isothermal bulk modulus was 69 GPa.[106]

### 3.6. Small angle X-ray scattering for microstructural characterization of building materials

When an X-ray (or a neutron) beam passes through a material, under the appropriate experimental conditions, a component of the direct beam is scattered out at a small angle due to heterogeneities (microstructure) of the sample. The angular profile of the SAXS (or SANS) intensity is effectively a Fourier transform of this microstructure. The recorded data are commonly analysed using appropriate microstructure models, and the microstructures are described/quantified through the resulting parameters. Although SAXS data can be obtained with laboratory sources, the quality of the data from SR instruments is much higher. Therefore, most of the SAS studies are carried out at synchrotron (or neutron) facilities. A focused review was devoted to the applications of SAS and SANS characterization techniques to C–S–H gel and cement pastes.[107] In this review, the similarities and differences between SAXS and SANS were also highlighted.

SAS (SAXS and SANS) techniques are nondestructive tools for characterizing density fluctuations over a wide range of length scales without altering the sample (sample preparation that may modify the microstructure is not commonly needed). Furthermore, it can probe interfaces of closed as well as open pores, unlike mercury porosimetry where information about the closed pores is not available. SAS provides statistically averaged information over the bulk of a material, unlike microscopy techniques where the information is limited to the local studied part of the sample. However, data treatment is important to extract the appropriate information and data analysis is far from obvious. There are several microstructure models ranging from Guinier approximation to fractal morphologies and full analysis of coherently ordered microstructures.[107,108]

Some examples of uses can be highlighted. By combining SAXS and SANS data with several other techniques, the mean formula and mass density of the nanoscale C–S–H gel particles in untreated hydrating cement were determined to be (CaO)$_{1.7}$SiO$_2$.1.80H$_2$O and 2.604g/cc, respectively.[109] These values differed from previous reported ones for the C–S–H gel, likely because of the specific drying conditions used in different approaches. Other study focused on determining the pore volume fraction of hydrated cement compacts from SAXS data and their evolution with hydration time.[110] The changes in the microstructures due to the use of cement additives have also been investigated by SAXS and SANS (see Figure 10).[111] The reported results showed that the used PCE tends to increase the size of the disk-like C-S-H globules but has little influence on the thickness of the water and calcium silicate layers within the globules. It must also be noted in importance of SANS studies. As a key example, this technique has been used for monitoring the degree of homogeneity of a cement paste matrix with the final goal of immobilizing low- and intermediate-level radioactive waste.[112]

Finally, SAXS data have also very recently been used to validate 2D high-resolution observations by TEM. Under some approximations, and using the Fourier slice theorem, it has been shown that the spectral density from TEM is a good approximation of the SAXS pattern. This has been used for a deeper characterization of the C–S–H gel where the computed SAXS signal (from TEM) and the measured spectrum agree quite well within the limited overlapping q–range (0.01 to 0.04 Å$^{-1}$).[113]
There are many different types of imaging methods based on different wavelengths (IR, soft X-rays and hard X-rays) as well as on different experimental set-ups. Figure 11 displays a schematic representation of the three more common techniques: full-field microscopy, scanning transmission microscopy and coherent diffraction imaging. All these three set of techniques (with subgroups) have been employed for characterizing the rich hierarchical microstructure details of cement pastes. A recent publication has reviewed state-of-the-art X-ray imaging techniques based on partially coherent synchrotron radiation including: full-field tomography, scanning transmission microscopy, ptychographic forward coherent diffraction imaging and scanning small-angle X-ray scattering [114] but it did not specifically deal with cement research. Another recent review [18] discussed the progress in the development of hard X-ray microscopy techniques for materials’ characterization at the nanoscale. They reviewed state-of-the-art hard X-ray synchrotron nanoscale microscopy techniques which included 3D tomographic visualization, spectroscopic elemental and chemical mapping, microdiffraction-based structural analysis, and coherent methods for nanomaterial imaging.

For full-field imaging, see Figure 11 (top); the synchrotron beam is relatively large and the ability to resolve tiny details of the sample arises from either a magnified projection onto the X-ray detector or from a magnifying lens. In the former approach, image resolution is limited by the pixel size of the detector and by the size of the X-ray source; in the second case, resolution
Figure 11. Scheme of the three main techniques for X-ray imaging highlighting some limitations. (Top) full-field microscopy, where the sample is fully bathed by the X-ray beam. (Intermediate) scanning transmission microscopy, where the sample is raster scanned by a focused beam. (Bottom) Coherent diffraction imaging, where the image is generated by an appropriate reconstruction algorithm. In the three cases, proper sample rotation allows recording tomographic data.

and image quality depend on the X-ray optics. The challenge to increase the resolution resides in producing better X-ray optics elements that combine high numerical aperture, high efficiency and low aberrations. This is assuming that radiation sample damage does not limit the achievable resolution.

For scanning imaging, see Figure 11 (intermediate); the beam is focused on a tiny spot that it is raster scanned to deliver 2D (radiographic) or 3D (tomographic) images. Furthermore, different properties can be measured (absorption, fluorescence, diffraction, etc.) which provide complementary information. For instance, STXM is a real-space imaging technique that utilizes focusing optics (it could be a Fresnel zone plate or a capillary) to deliver a small monochromatic X-ray beam onto a sample and it measures the intensity of the transmitted beam in raster mode, thereby filling an image array. STXM does not require an objective lens, and hence there is no attenuation of the beam between the sample and the detector, which means that the dose delivered to the sample is reduced compared to a full-field, lens-based approach. The procedure allows imaging with classical spectroscopic techniques, see below, at sample-tailored X-ray photon energies.

There is an alternative image-forming approach where the X-rays scattered by the sample are analysed and reconstructed by appropriate mathematical algorithms, see Figure 11 (bottom). Such lensless techniques are often grouped under the term: CDI or diffraction microscopy (not to be confused with X-ray diffraction micro-tomography, see below). It is widely believed that the CDI technique allows image resolution limited only by the tolerance of the samples to radiation damage. Furthermore, single-shot experiments at ultra-fast XFEL sources may even bypass this limitation. In a typical CDI experiment, a small sample is fully bathed with a plane wave (full-field technique), and its far-field scattering pattern is recorded in oversampling conditions. For more information about CDI, the interested reader is directed to a recent review.[115] At this stage, two related techniques should be mentioned that depend on where the detector is placed.

In BCDI, the far-field scattering pattern is collected at the right angle of a diffraction peak for a crystalline microparticle. In FCDI, the far-field scattering pattern is collected at the small angle scattering and the sample does not need to be crystalline. While the scattering process is well known and understood, inverting measurable intensity distributions to obtain an image of the
sample, often known as image retrieval, is a mathematically ill-posed problem where a unique solution has to be ensured. Image retrieval usually comprises many cycles of alternate iterative processes enforcing a priori knowledge on the sample, such as its finite size and consistency with measured data. Such a 'classical' CDI approach requires the samples to be isolated and has very low tolerance to perturbations by signals due to other scatters, i.e. from ice. Very importantly, there is an evolution of CDI where the samples are scanned by a small beam overlapping part of the illuminated samples (ptychographic approach) which favour/ensure the convergence of the reconstructing algorithms.

3.7.1. Full-field soft X-ray nano-tomography

Transmission soft X-ray full field microscopy was originally developed to study biological samples, but later has been applied to the study of the hydration processes of cement-based materials. Working with soft X-rays of an energy of $\approx 520$ (eV), which corresponds to a wavelength of 2.4 nm, it enhances the contrast between the cement particles and developing hydrates which contains more oxygens ‘water window’. Soft X-ray microscopy allows high-resolution imaging of hydrated samples over time and permits complete imaging of samples up to about 5 µm thick. For building materials this limitation effectively means loading dilute suspensions with very high w/c ratios (a severe drawback for many types of studies). In order to prevent excessive dissolution of the cement grains, several works used saturated solutions of CH and/or CSH2. However, saturated solutions cannot overcome the other artefact of dilute solutions, the increased availability of space in which hydration products are growing.

Initially 2D soft X-ray full-field microscopy, ACsX-nM, was used to directly image the hydration of C3S and OPC in a saturated solution of CH and CSH2 with time.\[116\] Figure 12(a) shows a C3S grain which has needle-like hydration products on its surface likely showing the of C-S-H gel formation after the induction period. Figure 12(b) and 12(c) shows the hydration evolution with time of OPC which clearly shows the formation of hexagonal prisms ettringite crystals. ACsX-nM has also been employed to study the differences in real-time hydration between pure (cubic) tricalcium aluminate and Na-doped C3A (orthorhombic) in aqueous solutions saturated in sulphates. The reactions involving cubic C3A were more influenced by higher concentrations of sulphate ions, forming smaller ettringite needles at a slower pace than for orthorhombic C3A.

It was also concluded that the rate of release of aluminate species into the solution was also accelerated by Na-doping.\[117\] The early-age hydration of ye’elimite phase (the main component of the calcium sulphonoluminate cements) has also been studied using this technique.\[118\] Diluted suspensions of this phase saturated in CSH2, and with variable amounts of CH, were examined. The most voluminous hydration product observed was ettringite. Under these hydration conditions, AFt commonly displayed acicular, filiform and plenty of intergrowths with stellate habits (see Figure 13).

The previous examples used 2D image evolution (microscopy) for obtaining the required information. The current technology allows to rapidly record a set of 2D images rotating the sample and so tomograms can be acquired and reconstructed. Therefore, as images may have nanometer resolution, the term nano-tomography was coined. ACsX-nCT has been recently employed to characterize the microstructure of two hydrated cement pastes.\[119\] Furthermore, it was possible to perform a direct computation of the ultra-small angle-scattering spectra from a single projection image which agreed well with the experimental synchrotron SAXS data obtained for the same pastes. Finally, soft X-ray nanotomography has also been employed for characterizing 2000-year-old Roman seawater concrete.\[101\] The data analysis showed clusters of Al–tobermorite crystals displaying both platy and elongated 1–2 µm crystals typical of geological 11 Å–tobermorite.
Full-field hard X-ray micro-tomography

Full-field transmission hard X-ray computed micro-tomography (here summarized as AChX-µCT) is the high-resolution adaptation of the CAT technique employed in medical applications since the 1970s. The AC-hX-µCT imaging technique is based on the virtual reconstruction of the inner density distribution of a sample from 2D X-ray radiographs collected at many viewing angles, while the sample rotates relatively to the source-detector direction (see Figure 14). We can distinguish two different steps in data handling. Initially, the recorded data must be processed to obtain a digitalized version of the studied object. The final outcome of the reconstruction process is a series of images called slices that, when stacked together after proper alignment, provide a 3D map of the spatial variations of the X-ray linear attenuation coefficient (µ) within the investigated object. Here, the key result is that each slice is composed by a matrix of voxels (volume elements) whose grey values are proportional to the mean value of µ (see Figure 14, bottom) The maximum spatial resolution achievable can reach the sub-micrometre scale for standard applications and is related not only to the technical specifications of the experimental set-up (characteristics of the synchrotron X-ray source and used detector), but also and chiefly to the size of the sample. As a general rule, higher resolutions require smaller samples due to the limited field of view of current X-ray area detectors. In a second stage, the digitalized object must be analysed with the appropriate software(s) depending on the information to be obtained: pore size distribution and connectivity, tortuosity, size and shapes of some particles, leaching effects, etc.

The main advantage of AC-hX-µCT is that it provides three-dimensional visualization of the internal microstructure of untreated samples. This is key as several other techniques for microstructure analysis (MIP, SEM, TEM) are known to produce irreversible changes in the
Figure 13. In-situ soft X-ray images, ACsX-nM, of hydrating C₄A₃S particles in a saturated CH-CSH₂
very diluted solution, showing the ettringite growth. Hydration time is indicated. Scale bars correspond to 1 µm. Reprinted from reference 116 with permission from Elsevier.

pore structure of cement pastes, particularly at small sizes. Furthermore, parameters such as connectivity and tortuosity are completely inaccessible by these techniques but they can be inferred from AC-hX-µCT, although the obtained results can be resolution dependent, see below. Fast data acquisition times are very important to avoid heating of the sample with the possibility to alter the microstructure of the studied sample due to local dehydration processes. The main two disadvantages of AC-hX-µCT are: first, its still low spatial resolution, voxel size of ≈0.5 µm³, relative to the sizes of capillary pores controlling transport properties in mature pastes. Second, sample preparation is delicate as narrow capillaries are required to achieve high resolution and in these conditions the w/c ratio and homogeneity of the pastes are difficult to ensure.

Perhaps it is worth mentioning that initial (independent) contributions from the Europeans, Japanese and Americans were reported within two years. In a seminal work of European researches,[120] AC-hX-µCT was used to study the connectivity and tortuosity of the pore network in OPC pastes. It was also shown that the degree of connectivity of the pore network was very sensitive to both the spatial resolution of the images and the evolution of contrast resolution during ageing of the cement. Some experimental conditions were: Lindemann Glass type capillaries with diameter of 600 µm and a wall thickness of 10 µm; pastes with w/c ratio of 0.5; X-ray
energy variable between 12 and 15 keV; 1001 projections with an angle step of 0.18° and an 
exposure time of 3 s; detector being a CCD camera (2048 pixels) equipped with a 1400 mm field 
of view and a 10 × magnification optical objective. Japanese experts studied not only porosity 
but the degree of pore connectivity and tortuosity,[121,122] which are very important param- 
ters to understand the mechanical performances and the durability of mortars and concretes. In 
these studies a voxel size of 0.5 µm³ was achieved and pastes at different hydration ages (2, 7 
and 28 days) were characterized. These authors also used AC-hX-µCT for studying the intern- 
al microstructure of deteriorated cementitious matrices due to leaching.[123] Finally, American 
researches applied both AC-hX-µCT and PC-hX-µCT to study the durability of concrete by 
analysing in-situ ice formation in entrained air voids within hydrated cement paste.[124] It must 
be highlighted that at X-ray wavelengths, the index of refraction is about 10–100 times larger 
than the absorption index. Thus the phase contrast mode of operation is more sensitive than the 
absorption mode. However, PC-hX-µCT data are harder to interpret quantitatively than those 
arising from AC-hX-µCT, unless phase retrieval procedures are implemented.

More recently, other works used these techniques for further characterization of building 
materials. For instance, AChX-µCT has been used [125,126] to study ASR which is one of 
the most important weathering processes in cement chemistry. To optimize the phase-contrast 
effects, the sample-to-detector distance was set to 200 mm. These authors found three different 
kinds of voids due to the effect of three different mechanisms: (i) cracks from ASR expansion, 
(ii) irregular-shaped voids due to the aggregate particles dissolution, and (iii) bubbles due to 
cement paste preparation. AC-hX-µCT has also been used to compare the early hydration 
of three cementing materials: OPC, CSA and a mixture of these two types of cements.[127] 
A full tomogram took 5 min and data were taken from hydration times between 1 and 12 h, 
the resolution being not high (0.74 × 0.74 µm pixel size). Some consequences of the hydration 
processes on the microstructure were followed including the porosity evolution with hydration 
time at early hydration ages. As an example, Figure 15 (left) shows the single slice evolution

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Figure 14. (Top) Typical system for absorption-contrast hard X-ray micro-computed tomography. (Bot- 
tom) Scheme describing pixel, voxel and slice, with each voxel being associated with an attenuation 
coefficient. From reference 119 © JCI – reprinted with permission.
Figure 15. (Left) Single slice evolution for the three samples (OPC, CSA and a blended OPC/CSA cement) at different hydration times (A: Lindemann glass tube; B: gypsum; C: portlandite; D: C4AF). (Right) Evolution of the grey-level histograms during cement hydration. Reprinted from reference 125 with permission from Elsevier.

of the three samples (OPC, CSA and an OPC-CSA mixture) at selected hydration times. Data can be analysed in a quite sophisticated way but an initial test is to plot the evolution with hydration of the grey-level histograms extracted from a Voxel (see Figure 15, right). These grey values correlate with the absorption coefficient and so with the hydration degree of the different components.[128] AC-hX-µCT has also been used to study emerging eco-cements like alkali-activated binders.[129] Insights into microstructural and pore structure characteristics were obtained including pore tortuosity calculated by a random walker method. AC-hX-µCT has also been used to compare the microstructure evolution of an OPC cement paste with and without a PCE superplasticizer.[130] Selected experimental details were: borosilicate glass capillaries with internal diameter of 600 µm; monochromatic beam of an energy of 14 keV; voxel size of 0.7 µm³, 30 min per single tomogram (about 800 ms per projection), and 20 mm between sample and detector which means that edge enhancement due to (propagation-based) phase contrast also contributed to the recorded signal. A very clear difference in the microstructure evolution between cement pastes with and without superplasticizers was shown.

There have been special applications of AC-hX-µCT in cements. For instance, it has been used with a white (filtered) synchrotron beam with energies ranging between 20 and 80 keV[131] and final voxel size of ≈ 18 µm³. The authors developed a method for extracting sub-voxel mineralogical and chemical information by combining advanced image segmentation with geochemical models of cement alteration. This method relies on determining ‘effective linear activity
coefficients’ for the white beam to generate calibration curves that relate the image greyscales
to material composition which was applied to the determination of diffusion profiles in altered
wellbore cements. Another very recent special application was collecting AC-hX-µCT data on
small cylinder specimens under load at varying degrees of damage.[132] Selected experimental
details were: 5 mm of diameter of the specimens; monochromatic beam of an energy of 30 keV;
voxel size of 6 µm³, and 2.5 h for recording a single tomogram. AC-hX-µCT allowed character-
izing microstructure and internal damage, which could then be related to bulk splitting strength
and fracture energy. Results showed that aggregate surface roughness had little effect on strength
but significant effect on fracture energy.

AC-hX-nCT has also been employed in cement studies. For attaining (dozens of) nanome-
ter resolution, the commonly focusing optics used before the sample are capillary lenses and
the field of view is considerably decreased when compared to AC-hX-µCT. Nanotomographic
reconstruction of a geopolymer binder, formed by hydration of fly ash, allowed very high reso-
lution (voxel size of 30 nm³) observation of the pore structure of the aluminosilicate geopolymer
gel.[133] However, it must be noted that this very high resolution was achieved by decreasing
the FoV approximately to 10 µm. Finally, it also worth mentioning that AC-hX-nCT, voxel size
of 64 nm³, has also been very recently obtained from a laboratory source.[134] The FoV in this
high resolution configuration was 65 µm in the three directions, with 12 h per tomogram. FZP
optics was used to focus the transmitted beam on a scintillator plate in front of a 20 × optical
device, and the energy source was a rotating anode copper tube which produces a polychromatic
beam with a maximum intensity of 8 keV.

3.7.3. Scanning synchrotron radiation microscopies

Synchrotron scanning microscopies comprise a set of techniques with a common feature, a small
monochromatic beam impinged onto a sample which is scanned resulting in an image array. The
image nature will depend upon the property being measured (transmitted X-ray beam, emitted
X-ray fluorescence, transmitted or reflected IR beam etc.). It must be also noted that many stud-
ies recorded more than one signal for the same sample set-up (for instance transmitted X-ray:
image, plus energy variation of the recorded absorption: elemental sensitivity, etc.), thus having
access to complementary information. Several techniques have been used for hydrated cement
characterization depending upon the property being measured, and key papers are discussed
below.

3.7.3.1. Scanning infrared spectro-microscopy

Infrared spectro-microscopy at synchrotrons provides the opportunity to collect spatially resolved IR data for samples with a spatial resolution
close to 3 µm (depending on the wavelength region to be used). Obviously, there are laboratory-
based IR spectro-microscopes but the brilliance of the infrared light produced at synchrotron
radiation is much higher compared to that of thermal sources (about three orders of magnitude)
and therefore it provides very good signal-to-noise ratio even when the microscope’s apertures
are set at the diffraction limit. This technique can operate in transmission, in total reflection and
in attenuated total reflection. The choice of the experimental configuration mainly depends on
the characteristics of the studied sample.

To the best of my knowledge, synchrotron IR spectro-microscopy has not been used in cement
chemistry as widely as other synchrotron techniques in spite of its high degree of chemical
selectivity by tuning to the specific vibration bands that can provide information about different
phases, additives, etc. I can highlight the work on the effect of seeded nucleation on the for-
mation and structural evolution of geopolymer gels.[135] This research showed that the nature
of the seeds affects the structure of the growing gel by affecting the extent of phase separation,
identified by the presence of a distinct silica-rich gel in addition to the main, alumina-richer gel
Figure 16. IR images of a $\alpha$-C$_2$SH single crystal (6 × 5 pixels, 5 µm step size) and after subsequent thermal treatment at different temperatures (19 × 16 pixels, 1.5 µm step size). The images were created by integration of the intensity over a certain absorption band which is represented by the colour of a given pixel yielding an image with white, pink, red, yellow, green and blue (high to low intensity, respectively). For more information, the reader is directed to the original publication. Reprinted from reference 134 with permission from John Wiley and Sons.

phase. Laboratory data could not detect such differences likely due to their poorer resolution. On the other hand, synchrotron IR spectro-microscopy can be used to follow chemical reactions (f.i. on heating) by following the evolution of the vibration bands. This use has been very elegantly employed to follow the evolution of a single crystal (grain) of $\alpha$-C$_2$SH on heating.[136] The exact transformation mechanism of dehydration remained controversial but this work showed conclusive evidences of the formation of Dellaite at about 350°C. Figure 16 shows diffraction-limited IR images of a thin $\alpha$-C$_2$SH crystallite, thickness $\approx$ 3 µm, showing the spatially resolved intensity evolution with temperature of selected vibration bands to highlight the appearance and disappearance of the phases.

3.7.3.2. Scanning transmission X-ray microscopy and tomography In STXM, the hard or soft X-ray beam is focused to a small point by the appropriate optics (commonly FZP). The transmitted X-rays are measured in the detector resulting in spatially resolved images because of the scan of the sample. If coherence properties of the beam are not used, soft X-rays are preferred as the contrast between different parts of a (heterogeneous) sample is enhanced mainly for soft condensed matter. The chemical information of the sample can be simultaneously obtained by acquiring a signal at a given position for multiple photon energies.

Using soft X-rays (energy ranging between 200 and 2000 eV), the effect of polymers on the nanostructure and on the carbonation of calcium silicate hydrates was studied.[137] Images of the transmitted soft X-ray beam as well as XANES (also known in soft X-ray studies as NEXAFS)
spectra for the C, Ca and Si edges were obtained. The authors reported a different behaviour for the absorption and carbonation of polyethylene glycol and hexadecyltrimethylammonium polymers on C–S–H. STXM coupled with NEXAFS spectra for C, Ca, Al and Si edges were used to better characterize Al-tobermorite-rich Roman seawater concrete.[138] The microstructure of a 2000-year-old concrete block submerged under the Mediterranean Sea was comprehensively characterized including the spatial arrangements of Al-tobermorite and the poorly crystalline calcium-aluminium-silicate-hydrate (C–A–S–H) binder (see Figure 17). Finally, scanning transmission soft X-ray microscopy and full-field hard X-ray micro-tomography were used to study the interactions and microstructure of an OPC paste reinforced with polymeric fibres.[139]

3.7.3.3. Scanning fluorescence X-ray microscopy and tomography In scanning fluorescence microscopy, a hard X-ray beam (energy can be close to 10 keV) is also focused by the appropriate optics (commonly FZP). Emitted characteristic X-ray fluorescence radiation is detected with the suitable energy dispersive detector (f.i. a multi-channel silicon drift detector). Scanning the sample with small steps (they can be as small as 20–30 nm) provides the high resolution multi-element images with the elemental compositions (derived from the integrated area of the recorded fluorescence signal).

SFXM was used to characterize the nanoscale distribution of elements within fly ash geopolymers, the binders obtained by alkaline activation of fly ash mainly formed by aluminosilicate gel(s).[140] Hydroxide-activated geopolymer gel was highly heterogeneous with high-content Ca particles within the geopolymer binder matrix. Conversely, silicate-activated geopolymer gel showed a much more homogeneous geopolymer gel binder structure surrounding the unreacted
fly ash particles. Several transition metals (Cr, Fe . . . ) were mapped. In a related recent work,[141] SFXM has been used to quantify the effects of the activator concentration on the microstructure of alkali silicate-activated slag/metakaolin pastes. The space-resolution properties of this technique (a 10.5 keV beam focused down to 60 nm) allowed to distinguish two coexisting gels: calcium aluminosilicate hydrate (C–A–S–H) and sodium/calcium aluminosilicate hydrate [(C,N)–A–S–H] type gels. As an example of an outcome of this type of studies, Figure 18 shows X-ray fluorescence micrographs for selected elements in this system. Finally, this technique has also been applied to study alkali-activated slag binders.[142] A key observation was that iron-rich, titanium-rich and manganese/silicon-rich particles remain stable under the reducing conditions prevailing during alkaline activation. There was no evidence of chemical interaction between these particles and the geopolymer binder.

On the other hand, the combination of techniques always provides a better picture of the studied sample/processes. One selected example is the combined use of soft X-ray SFXM and micro-XANES spectroscopy to determine the spatial distribution of Al and S and to identify the Al- and S-bearing species in compact hardened OPC paste hydrated at 50°C.[143] SFXM data were obtained by focusing a 3.9 keV beam (energy just below the Ca absorption edge) to a size of approximately 3 × 3 µm, and by measuring the emitted fluorescence X-rays with a single-element silicon drift detector. The Al and S K-edges micro-XANES spectra were recorded in fluorescence yield mode using the single-element drift diode detector. The depth of X-ray penetration was slightly less near the Al K-edge (∼3.6 µm at 1559.6 eV) than the S K-edge (∼9.4 µm at 2472 eV). The relatively large escape depths, similar to the penetration depths, mean that the size of the sample probed could be tens of µm², possibly containing several species. A second selected example is the combined use of hard X-ray SFXM and micro-diffraction to study the phase assemblage and microstructure of the hydration products formed in blended OPC cements with high-volume fly ash.[144] The reported data showed that the C–S–H formed in the system containing 50% of fly ash had a similar structure as C–S–H(I) with comparatively lower Ca/Si ratio than the one produced in the OPC system. Moreover, coexistence of C–S–H(I) and
strätlingite was observed in the system containing 80% of fly ash, confirming that the amount of alumina and silicate phases provided by the fly ash is a major factor for the formation of strätlingite (and C–S–H).

One selected example on the combined use of hard X-ray SFXM and micro-EXAFS and micro-XANES spectroscopies is the work on the Ni uptake by OPC pastes including the influence of the inherent heterogeneity of the cement matrix on the Ni speciation.[145] Some key experimental details were: beam size of $5 \times 5 \, \mu\text{m}$, a fixed beam energy of 10 keV for the µ-XRF study, and a variable beam energy (close to the Ni K-edge, 8.3 keV) for the µ-XAS study. Both µ-XRF and µ-XAS data were collected at room temperature in fluorescence mode using a seven-element Ge-solid-state detector. Unfortunately, the thicknesses of the thin sections were not reported.

3.7.3.4. Scanning diffraction hard X-ray microscopy

In section 3.3, the uses of synchrotron powder diffraction for quantifying crystalline phases and to follow phase evolutions are discussed. These works used a large beam (usually larger than a millimetre) yielding an accurate average picture but without spatial resolution. However, for heterogeneous materials, like cement binders, and for some applications, it is invaluable to have the spatial distribution of the different phases (for instance the changes/alteration with depth due to sulphate attack). This can be obtained by focusing the beam down to several micrometric size and scanning the sample in the appropriate direction. The technique, SDXM, works in transmission but not only is beam focusing important (for instance using K-B mirrors) but sample preparation is also very important. For this technique to be useful, a relatively thin (unaltered) cross-section must be prepared along the appropriate direction (see Figure 19). Typical thicknesses for the flat slices used in these studies ranged from 100 to 500 µm. This technique is commonly known simply as synchrotron microdiffraction.

SDXM, aka synchrotron microdiffraction, was used to quantify the orientation distribution of fibrous ettringite crystals after sulphate attack in fractured concretes.[146] Sample preparation is key and a summary of the process follows. A flat surface was impregnated with epoxy resin and then mounted on a glass slide. Then, using a diamond saw, and with kerosene as cooling agent, a 50 µm thin slice was prepared also sealed with epoxy. Finally, the slice was removed from the glass slide and used for the synchrotron characterization (beam energy: 8 keV, beam...
size: $2 \times 2 \mu m$). The analysis revealed that the c axes of the trigonal ettringite crystallites were preferentially oriented perpendicular to the fracture surfaces. SDXM was also employed to determine, with spatial resolution, the changes in the phase composition due to sulphate attack [147] for establishing the durability of cementitious materials under these conditions. Key experimental details were, beam energy: 11.6 keV, beam size: 10 µm, sample thickness: 200 µm. Furthermore, the sulphate attacks are affected by the presence of SCM. In two subsequent works from the same group, the microstructural profile analyses of concrete deterioration after sulphate attack of OPC blended with fly ash [148] and with natural pozzolana, granulated blast furnace slag or fly ash [149] were deeply investigated. In the first work,[148] OPC was mixed with 30 wt% of class F fly ash and hydrated with w/c ratio of 0.5 for 28 days. Afterwards, the sulphate attack was carried out under laboratory conditions for 6 months. Then, the samples were embedded in epoxy resin and polished to thicknesses of 200 µm. During all preparation steps the samples were cooled with petroleum to avoid dissolution of water-soluble phases. Key experimental details were, beam energy: 14.5 keV, beam size: 10 µm. In the second work,[149] the sample specimens were embedded in sulphate-bearing soil ($\approx 1 \text{wt} \% \text{SO}_4^{2-}$) for 19 years. The samples were also embedded in epoxy resin for the preparation of cross-sections which were made parallel to the direction of the sulphate ingress. Selected experimental details were, beam energy: 12.4 keV, sample thicknesses: 500 µm. The beam size was not reported but it was mentioned that the achieved spatial resolution was 30 µm.

Finally, synchrotron microdiffraction can be used to map the phase distributions in any complex binders after successful sample preparation. For instance, this technique has been very recently applied to study pyroclastic aggregate concrete of Trajan’s Markets (1900 years old) as well as their reproductions (experimental archaeology): hydrated lime–volcanic ash mortar that binds decimeter-sized tuff and brick aggregates.[150] The mortar reproduction gains fracture toughness over 180 d through progressive coalescence of C–A–S–H gel binder and crystallization of strätlingite and katoite at $\geq 90$ days, after pozzolanic consumption of hydrated lime was complete. Key experimental details were, beam energy: 10 keV, beam size: $8 \times 2 \mu m$, and sample thickness: 300 µm.

### 3.7.3.5. Scanning diffraction hard X-ray micro-tomography

SDX-µCT [151] is a combination of diffraction (crystalline phase sensitive) and imaging (through tomographic reconstruction) techniques which allows determining the three-dimensional spatial distribution of different phases within heterogeneous samples. The reconstruction scheme for the SDX-µCT technique is depicted in Figure 20. Unless the previous technique, SXDM, flat thin sections are not needed. Furthermore, the appropriate rescaling of the voxel intensity to the total intensity of sample scattering makes it possible to obtain the absolute quantification of the phase proportions in each voxel. SDX-µCT was initially used for 3D monitoring of the evolution of the microstructure and phase formation non-invasively.[152–154] For this particular set of experiments, the authors used a monochromatic beam ($E = 18 \text{keV}$) with a size of $2 \times 4 \mu m^2$. Although this technique gives good insight into the three-dimensional phase arrangement at intermediate and later hydration ages, the long acquisition times ($\sim 8 \text{h}$ for a slice of 500 µm diameter and 2 µm thickness) do not allow the phase mapping at early stages of hydration, as reaction kinetics are too fast for the microstructure to be resolved. Subsequently, AC-µCT was used at early ages (7 h of hydration) and SDX-µCT at later ages (7 days of hydration) for obtaining better insight into the phase and microstructure developments in hydrating pastes.[155]

SDX-µCT, combined with numerical simulations for C–S–H precipitation, was used to investigate the C–S–H topological distribution and modes of precipitation.[156] This technique has been recently used to map the phases present in two hydrating OPC cement pastes (one sample hydrated with pure water and a second one hydrated in the presence of nucleation seeds).

The quantitative description of the phase spatial distribution by radial distribution functions...
allows the discrimination of different nucleation mechanisms.[157] SDX-µCT has also been very recently used to map the C – S – H precipitation in the absence and presence of superplasticizer, (see Figure 21, left). The observed spatial correlation between C – S – H and unhydrated cement particle surfaces indicated that, in the absence of PCE superplasticizers, C – S – H forms by a process of heterogeneous nucleation, on the surface of the dissolving cement particles (see Figure 21, right). Conversely, the lack of significant spatial correlation between C – S – H and the surface of unhydrated particles, when PCE is added to the system, revealed that C – S – H precipitates randomly throughout the available space in the paste.[158]

3.7.4. Hard X-ray coherent diffraction imaging

As it was mentioned above, there is an alternative to image-forming optics where the X-rays scattered by the sample are reconstructed by appropriate algorithms. These set of techniques are commonly named CDI techniques, and the two most common ones are discussed below.

3.7.4.1. Hard X-ray Bragg coherent diffraction nano-tomography

BCDI is also a non-invasive imaging technique which can yield three-dimensional images of individual crystals on the nano-scale through inversion of the diffraction data by a computational method, but it also highly sensitive to crystal defects and strain fields inside crystals seen as phase evolution. The experimental set-up (see Figure 22) is compatible with cement hydrations as very recently reported.[159] In this study, the early hydration (up to 3 days) of microcrystals of calcium monoa-
Figure 21. Scanning diffraction X-ray micro-tomography results. (Left) Phase maps displaying the space distribution of the unhydrated cement particles (red), C\textsuperscript{−}S\textsuperscript{−}H (green) and the combination of the two within a virtual slice through an OPC paste sample hydrating in water and in the presence of a PCE superplasticizer. The colour intensity is proportional to the volume fraction of the given phases in each voxel. The white circle represents the enclosing glass capillary (internal diameter 400 µm). (Right) Radial distribution functions relative to the spatial distribution of C\textsuperscript{−}S\textsuperscript{−}H with respect to the position of the unhydrated particle surfaces, as calculated for: (a) the phase maps of the OPC sample without PCE, at 7 days of hydration; (b) the phase maps of the OPC sample with PCE, at 7 days of hydration; (c) the difference phase map of the OPC sample with PCE at shorter times (7 − 17 h). Reprinted with permission from reference 156. Copyright © 2015 American Chemical Society.

to the change of the degree of crystal ordering, which could occur through ion transfer during hydration. The observed strain, coming from interfacial mismatch effect between high Bragg density and low Bragg density parts in the crystal, remained throughout the experiment. The first Bragg density change during hydration process was due to a big loss of Bragg density and was seen as removal of density, but not phase. The work provided new evidence supporting the through-solution reaction mechanism of calcium monoauminate.

3.7.4.2. Hard X-ray ptychographic forward coherent diffraction nano-tomography PFCDI-nCT is a non-invasive imaging technique based on the (partly) coherent properties of synchrotron radiation and it allows the three-dimensional mapping of the electron density in the studied sample. This variant of CDI does not require crystalline ordering and its quantitativeness combined with a resolution close to 100 nm (for a field of view of about 60–100 µm) makes it very appropriate for studying the hierarchical microstructures in complex materials including cement pastes. The term ‘ptychography’ stems from the Greek word for a fold, related to the Latin origin of the term convolution. Instead of fully illuminating a small sample with a featureless plane wave,

Figure 23. Cross-sections through the reconstructed 3D images of the CA crystal showing the internal Bragg density variations during 3 days of hydration measured by BCDI. (a) unhydrated CA, (b) CA hydrated for 2h, (c) CA hydrated for 24h, (d) CA hydrated for 52h, (e) CA hydrated for 67h. Reprinted with permission from reference 157. Copyright (2015) American Chemical Society.
Figure 24. (a) Schematic representation of the experimental set-up for hard X-ray ptychographic forward coherent nano-tomography. At each incidence angle, coherent diffraction patterns are recorded by a pixilated detector for a number of overlapping scanning positions, which allows the projected complex-valued transmission function to be reconstructed. (b) Example of a single reconstructed phase projection of the epoxy resin impregnated hardened cement paste sample. The black dots indicate the scanning positions at which diffraction patterns were recorded, and the black circles represent the approximate shape of the beam – shown for the first two shells of the circular scan only. The scale bar corresponds to 5 µm. Reprinted from reference 158 with permission from Elsevier.

PFCDI-nCT was applied to image hydrating cement pastes. Figure 24 illustrates the experimental set-up (a) as well as the result (b) for imaging a sample of resin-impregnated, hardened cement paste. The data from multiple known scan positions are inverted to yield a 2D image, whose resolution is limited by the maximum scattering angle where there is signal and by the positioning accuracy of the sample. When combined with a rotation stage/strategy, a 3D tomographic image can be obtained. Furthermore, the high accuracy in measuring the electron density allows accurate segmentation of the data. In a very recent work, PFCDI-nCT has been applied to the microstructural characterization of C–S–H formed by hydrating C₃S. The 3D spatial resolution of the phase contrast images was close to 130 nm, whereas the resolution of the absorption images was poorer, ≈ 250 nm (see Figure 25). It has been observed that the C–S–H density can depend on the particles’ states of hydration. For fully hydrated particles, the estimated density of the outer-product C–S–H was larger than that of the inner-product C–S–H, whereas for the partially hydrated particles, the densities of the apparent outer and the inner products were very similar. The density values of C–S–H ranged from 1.72 g cm⁻³ to 1.96 g cm⁻³, and its water content ranges from 4.3 to 7.6 mol, assuming a fixed C/S molar ratio of 1.75.

4. Outlook

It is difficult to forecast the main research lines in cements using synchrotron tools as these evolve quite rapidly. In any case, there are challenges in the chemistry of cements where developments in the synchrotron characterization techniques may play a leading role. I highlight my shortlist below:

1. Most cement binders are based on amorphous gels or they contain large contents of amorphous materials. We all know that the characterization of amorphous materials is always complicated due to the lack of long-range order and periodicity, and also because of their
Figure 25. Vertical slices of the (a) phase-contrast and (b) absorption-contrast nano-tomograms of the hydrated cement paste, UN: Unhydrated C$_3$S, CH, C$-$S$-$H, Q (quartz capillary) and W (pore solution). (c) 3D renderings of the volume showing how the phases are located with respect to each other. (d) Bivariate histogram of absorption and phase. (e) Radial electron density profile of the particles as a function of distance from surface inward. The effective particle diameters are shown in the legend. (f) Site-specific water content of the C$-$S$-$H at the same slice shown in (a) and (b) with a resolution of 1 µm, where the colorbar is given in units of mol of water. Reprinted from reference 159 which is an open access article published under a Creative Commons Non-Commercial No Derivative Works (CC-BY-NC-ND) Attribution License.
large chemical variability. Advances in synchrotron tools will tackle these issues starting
with the determination of chemical compositions and density values of these gels with very
high spatial resolution by further development of the appropriate (combination) of imaging
techniques.

2. The microstructure quantitative study of cement pastes is very important to understand and
predict their mechanical behaviour as well as chemical durability. In this arena, synchrotron
tools are very well suited as they do not require special sample preparation or sample envi-
ronments that alter the microstructures. Here the challenge is to continue developing the
imaging techniques, and the sample preparation procedures, for entry into the resolution
range lower than 100 nm. A very good resolution, well below 100nm, without a trade-off of
field-of-view is important to properly characterize key pore microstructure details: connect-
tivity and tortuosity. It will be also important to quantify the changes in the microstructure
provoked by the uses of SCM which can vary quite a lot (fly ashes, slags, partially burned
clays, etc.). For this type of application, one of the techniques with the brightest future, in
my opinion, is hard X-ray ptychographic forward coherent diffraction nano-tomography.

3. Most crystalline materials in cement chemistry have known crystal structures. However,
some hydrates have still unknown crystal structures. For instance, C$_2$AH$_8$ has been known
for more than a century and its crystal structure is still not reported although it is known that
it belongs to the AFm type structure. This is due to the combination of its chemical instability
(it loses water very easily) with the lack of single crystals and that it is so far crystallized
with additional coexisting phases. Here, microcrystal structure determination tools could be
key to determine the crystal structure of this type of compounds from powder with grains
smaller than 5 µm but using single-crystal-like techniques.

4. The footprint of cement production is high and should be reduced, but retaining the life
standards. Therefore the development of eco-cements is important to decrease anthropogenic
CO$_2$ emissions but maintaining the quality of our buildings and constructions. This can be
tackled in a number of ways including the partial replacement of OPC by SCM but also by
developing new binders, not based in OPC, like alkaline-activated materials and sulpho-
belite cements. In this direction, synchrotron techniques are being used, and will be used more
intensively in the future, to shorten the time between chemistry formulation developments
and their market appearances. This usage is very important as durability of new binder must
be ensured and to do this, the understanding and quantification of their microstructures are
vital.

5. Finally, it is worth mentioning that under-construction and planned diffraction-limited stor-
age rings (fourth-generation synchrotron sources) will produce smaller beams with higher
flux/brilliance of much higher coherence. These properties will directly impact the points
described above and several others. The interested reader is directed to the special issue
of Journal of Synchrotron Radiation published in September of 2014 which was fully
devoted to the technical developments and science to be carried out in these last-generation
synchrotrons.

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Disclosure statement

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Miguel A. G. Aranda received his Ph.D. from the University of Malaga (Spain) in 1992 in the field of Rietveld analysis for structure determination. During his Ph.D. he carried out three summer research stays at Chemical Crystallography Laboratory (University of Oxford). He carried out his postdoctoral training at the University of Cambridge with Paul Attfield to work on the crystal structure of Cu-based high-Tc superconductors using synchrotron and neutron powder diffraction. He came back in 1994 as Assistant Professor to the University of Malaga where he also was Associate Professor and Professor. In January 2013, he moved to ALBA Synchrotron Light Source as Scientific Director. He has research experience in cements and building materials but also in other fields like: ceramics, pigments, cultural heritage and archaeometry as well as strongly electron-correlated transition-metal oxides, solid-oxide fuel cells and metal-organic-framework materials. His hobbies include walking in the hills, snorkelling, travelling and enjoying good wine and food.

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