Soft X-ray Microscopy Techniques for Medical and Biological Imaging at TwinMic—Elettra

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Abstract: Progress in nanotechnology calls for material probing techniques of high sensitivity and resolution. Such techniques are also used for high-impact studies of nanoscale materials in medicine and biology. Soft X-ray microscopy has been successfully used for investigating complex biological processes occurring at micrometric and sub-micrometric length scales and is one of the most powerful tools in medicine and the life sciences. Here, we present the capabilities of the TwinMic soft X-ray microscopy end-station at the Elettra synchrotron in the context of medical and biological imaging, while we also describe novel uses and developments.

Keywords: X-ray microscopy; soft X-ray; X-ray fluorescence; ptychography; compressive sensing

1. Introduction

X-ray microscopy gathers a cohort of techniques that allows the acquisition of two and/or three-dimensional views of an object at micrometric and sub-micrometric spatial resolutions by the use of X-ray radiation [1,2]. Soft X-ray microscopy, which is a subset of it, employing soft X-ray photons (usually in the 200–2000 eV energy range) as the illumination source, is nowadays a standard technique in synchrotron radiation facilities [3–13]. Even though X-ray microscopy cannot compete with the nanometer or sub-nanometer spatial resolution of electron microscopes, it allows for a more relaxed sample preparation, especially for biological and organic specimens, and for higher penetration depths, thus permitting thicker samples to be analyzed. Moreover, the high sensitivity of X-ray-spectro-microscopy, such as X-ray fluorescence (XRF) microscopy, makes it an essential tool for tracking chemical changes at the micro- and nanoscale.

The TwinMic beamline end-station at Elettra [7] is a soft X-ray microscope originally designed to provide a collection of imaging techniques in a single instrument [14], specifically with an easy switch between scanning transmission X-ray microscopy (STXM) and full-field modes. Afterward, STXM has become the more dominant operation mode, especially due to the development of the first worldwide low-energy X-ray fluorescence (LEXRF) system [15–17].

TwinMic capabilities have attracted the interest of scientists from different research fields, from the material sciences [18], especially from the energy field [19–22], to environmental sciences [23–31], the life sciences [32–46], and cultural heritage [47].

In the present manuscript, we describe the TwinMic imaging and spectroscopic techniques made available especially to the biological and medical user community. We highlight the TwinMic potentialities and the recent achievements in the medical and biological field through suitable research cases, demonstrating its contribution and complementarity with other microscopy approaches.
Finally, we introduce new ongoing developments regarding advanced smart data acquisition solutions and novel data analysis methods.

2. Materials and Methods

2.1. TwinMic Imaging Modes

The present section illustrates in detail the TwinMic Imaging Modes, which are also summarized in Table 1.

Table 1. Summary of TwinMic techniques.

| Technique       | Spatial Resolution [µm] | Typical Acquisition Time/Pixel [s] | Overall Acquisition Time [h] | Field of View [µm] | Detector Type |
|-----------------|-------------------------|-----------------------------------|-----------------------------|--------------------|---------------|
| STXM            | 0.08–2                  | 0.005–0.05                        | 0.1–2                       | 10–80 with piezo motors; 1000 with stepper motors | EMCCD          |
| LEXRF           | 0.4–2                   | 1–10                              | 2–10                        | 10–80 with piezo motors; 1000 with stepper motors | SDD            |
| Ptychography 3  | <0.04                   | 0.1–10                           | 0.05–0.5                    | 10–80              | CCD          |
| Full-field 3    | 0.02–0.05               | few seconds or fractions of seconds | few seconds or fractions of seconds | 20             | CCD          |

1 Recently, with compressive sensing (CS) implementation and sample stage upgrade, we have been able to reach fields of view of about 1 mm × 1 mm. 2 Electron-multiplying CCD. 3 Here, we report the values for full frame acquisition.

2.1.1. Scanning Transmission X-ray Microscopy

STXM is nowadays a standard X-ray microscopy technique, used especially in the soft or milder X-ray regime. It requires suitable X-ray optics, usually zone plate (ZP) diffractive optics, to focus the X-ray photons on the sample plane, where the specimen is usually scanned across the microprobe. In a few cases, the beam can be scanned across the sample [4]. In scanning techniques, the spatial resolution is determined by the probe size.

For each point in the raster scan, the transmitted X-rays are collected by means of suitable detectors (photodiode, ionization chamber, CCD . . . ), allowing one to record the absorption signal. In the case of a two-dimensional detector, as with the TwinMic beamline, in addition to the absorption image, differential phase contrast and darkfield contrast can be acquired simultaneously, providing a complementary imaging mode [48–51]. According to the incident beam intensity and the detector characteristics, where the transmitted signals are intrinsically acquired in a serial way, STXM images may require from a few minutes to several tens of minutes. However, by means of suitable detectors, the scanning operation mode also allows for the simultaneous acquisition of other signals point-by-point in the raster scan, such as XRF photons [15] or photoelectrons [52] emitted by the sample. Thus, besides morphological images, chemical ones can be obtained. Similarly, by collecting a stack of absorption images across the absorption edge of specific elements, after suitable image alignment, one can extract X-ray absorption near-edge spectroscopy (XANES) signals, which allows for element speciation identification [4,8,10].

A schematic view of the TwinMic STXM operation mode is presented in Figure 1. A series of pinholes of different dimensions acts as a secondary source for the microscope. The incoming monochromatic beam is focused on the sample plane through ZP diffractive optics, whose first diffraction order is selected by the order sorting aperture. The size of the focused beam is given by the secondary source size demagnified through the ZP. For the experiments shown here, a 600 µm diameter Au ZP with a 50 nm outermost zone was used. The transmitted X-rays are collected by an electron multiplying CCD (EMCCD) camera DV860 from Andor Technology (128 × 128 pixels, 24 × 24 µm² pixel size) by means of an X-ray–visible light converting system constituted by a P43 phosphor screen, a visible light lens, a mirror tilted by 45º, and an objective lens mounted on the CCD camera itself [50].
Figure 1. Schematic view of the TwinMic STXM mode setup with the microprobe forming zone plate (ZP) on the specimen (S), diffraction order-selecting aperture (OSA), transmission detection system with fast-read-out CCD camera (FRCCD) and visible light converting system (VLCS), and low-energy X-ray fluorescence (LEXRF) detector system based on 8 silicon drift detectors (SDDs) in backscattered configuration. The VLCS consists of a phosphor screen (PS), a lens (L) and a 45°-tilted mirror (M).

Eight silicon drift detectors (SDDs) are mounted circularly and symmetrically, with their axes placed at 20° from the sample plane and their center 28 mm from the focal spot [15,16]. The output signal of the preamplifier electronics (XGLab srl, Bruker Nano-Analytcs), which is placed in the TwinMic vacuum chamber in close proximity to the SDD chip, is then processed through the DANTE Digital Pulse Processor (XGLab srl, Bruker Bruker Nano-Analytics) [53]. A new SDD system consisting of 4 trapezoid chips, each with 8 square SDDs on it, is under development and has been already tested [54–56], and it is sketched in Figure 2. The new system will cover a larger solid angle (5 to 10 times higher) than the current one, allowing for faster measurements and, thus, reduced radiation damage. Once optimized, it will replace the current LEXRF system.

Figure 2. Schematic view of the TwinMic current (A) and future (B) LEXRF system setup.

While LEXRF mapping is a standard TwinMic acquisition mode, XANES spectroscopy is currently under development. It has already been performed on materials science-related topics [18,22,57], where metal concentrations were high enough to detect speciation changes; however, in order to become a standard TwinMic technique, it requires fine tuning and some changes in the beamline acquisition/control software.

The scanning stage where the sample is located consists of two X and Y stepper motors, where a X–Y piezoelectric motor stage (with 80 μm range of movement) is mounted. In the
first version of the sample stage, the scanning was performed exclusively by using the piezoelectric motors, after positioning the sample in the desired area by means of the stepper motors, allowing a movement range of 5 mm in both X and Y [7,17]. A newer version of it was recently installed by replacing the mechanics and the stepper motors, allowing for a larger range of movement (12 mm × 12 mm). This allows for larger samples and/or a larger number of samples to be mounted, avoiding several vacuum breaks and, therefore, reducing the measurements’ deadtimes. Moreover, with the new sample stage, both STXM and XRF can be performed not only by using the piezoelectric motors, as originally designed, but also in a coarser way by using the stepper motors themselves. This allows for larger areas to be scanned, as it is shown in the results section.

2.1.2. Full-Field X-ray Microscopy (TXM)

TwinMic was originally designed to be able to operate both in STXM and TXM mode with an easy switch between them [14]. Finally, the STXM mode resulted in the most requested one, because of the increasing interest of the scientific community to the newly developed LEXRF. Indeed, scanning instruments are well suited to combine imaging and spectroscopy measurements for a given fixed X-ray energy. On the other hand, the TXM mode offers a faster acquisition time, of the order of seconds, and a higher spatial resolution. For this reason, it is usually well suited for monitoring dynamic changes in the sample or for the tomography experiment, where multiple images are taken sequentially.

The TwinMic TXM mode is summarised in Figure 3. A condenser ZP (CPZ) optics illuminates an area of the sample, which is magnified through a high-resolution ZP on an imaging detector. The CZP used at TwinMic is 1 mm in diameter and consists of 256 grating elements in 5 rings. The grating constant is 218 nm for the innermost ring and 126 nm for the outermost ring. Each grating covers an area of 50 × 50 µm². CZPs were fabricated by Zoneplates.com. The transmitted X-rays are collected by a back-illuminated Princeton CCD camera with 1340 × 1300 pixels, a pixel size of 20 × 20 µm², and a 2 s readout time when a full frame is acquired [7,17].

![Figure 3. Schematic view of the TwinMic full field imaging mode setup, with a condenser ZP (CZP) illuminating the specimen (S), an order-selecting aperture (OSA), and a micro zone plate (MZP) that magnifies the image of the specimen into the detector with a directly illuminated CCD camera.](image)

2.1.3. Ptychography and Coherent Diffraction Imaging (CDI)

Ptychography [58,59] is an emerging scanning imaging technique based on CDI [60–63]. It is attracting interest at synchrotron facilities, as it provides not only transmission imaging data, such as STXM, but also phase information simultaneously. The technique provides
very high resolution over large sample areas and is characterized by the high reliance on computationally demanding techniques for the reconstruction of the results.

The single image (probe) is a diffraction image that, via numerical inversion, can provide us with the original transmission image in terms of intensity and phase. This is a complex mathematical problem with multiple solutions. To obtain the right solution, the sample is scanned at a step size smaller than the probe size, ensuring adequate overlapping of subsequent illuminated areas, acquiring a diffraction pattern for each step in the raster scan. Indeed, the redundant information due to the oversampling allows computer algorithms to reconstruct reliably both the sample transmission function and the illuminating probe, recovering the amplitude and phase of both these signals. As in classical SXTM, by reconstructing a stack of images acquired at photon energies across the absorption edge of a specific chemical element, ptychography allows for XANES-type studies, as demonstrated in [64,65], providing sub-micrometric resolution chemical speciation.

In addition to ptychography experiments, the beamline has successfully tested the keyhole-CDI technique [61], which, by being a single-shot technique, is in general, preferable for in situ dynamic studies, such as monitoring the dynamics of electrodeposition [64,65], but which can also be used for biological samples [66].

Figure 4 outlines the direct-detection transmission X-ray ptychography setup at TwinMic, where ZP optics illuminates an area over a sample placed at a certain distance from the focus, while the sample is raster-scanned with a step size smaller than the probe size. Currently, the same previously mentioned back-illuminated Princeton CCD camera used for TXM is also employed for ptychography.

Figure 4. Schematic view of the TwinMic ptychography setup with the zone plate (ZP), the order-selecting aperture (OSA), the specimen (S), and the CCD camera.

Ptychography can be classified as a computational technique as it requires reconstruction similar to that of a computed tomography workflow. Advances in algorithms and computational resources have affected the quality and speed of reconstruction. TwinMic, in collaboration with the Elettra Scientific Computing group, has developed a specialized data acquisition and ptychographic reconstruction in-house software called SciComPty that allows for the state-of-the-art reconstruction and optimization of various experimental parameters. SciComPty [67] was developed and tested with data at energy ranges suitable for medical and biological studies. The system and the in-house computing expertise should allow noncomputational researchers to employ this technique for an easier absorption and phase imaging data analysis.
2.1.4. Compressive Sensing

Originally introduced in the astronomy field, compressive sensing [68,69] is an emerging method where image reconstruction is performed from a limited (ideally small) number of datasets without compromising the final quality. It is based on the assumption that the “most useful” datasets are smartly acquired, while the less significant ones can be easily extrapolated by a fast or small dataset. Inspired by this idea, we have recently developed in TwinMic a set of approaches [70,71] that allow us to perform dynamic scans where it is possible to skip points (sparse) and perform acquisitions with variable parameters (i.e., acquisition time). Such an approach leads to a substantial reduction in the required time, besides reducing the size of stored datasets. This reduction in terms of time, coupled with other technological and scientific advances such as new detectors or new sources, can produce maps of very large areas (one or two orders of magnitude higher than the standard ones), which could not be acquired with the traditional ways. Briefly, we proposed three different methods for optimizing the XRF data acquisition, where only the most significant XRF data are collected, by: (i) applying a mask based on a previously acquired absorption image (and therefore based on the sample morphology); (ii) applying an absorption threshold, and therefore acquiring the XRF signal only on specific regions, which represent the real sample (and not the support or the embedding medium); (iii) by applying a threshold on the XRF signal and therefore acquiring a longer XRF spectrum in the regions where a specific element of interest is present in higher concentrations. Naturally, such methods can also be applied as a combination, allowing an even smarter acquisition strategy.

Of course, these approaches can be extended to other kinds of microscopies or analytical techniques, as suggested or explored by other groups [72–75]. Compressive sensing is thus an imaging strategy that can represent a TwinMic imaging modality.

2.2. Sample Preparation

2.2.1. Balb/3T3 Mouse Fibroblast Cells Exposed to CoFe$_2$O$_4$ NanoParticles (NPs) and Met5A Cells Exposed to Crocidolite Fibres

The Balb/3T3 mouse fibroblasts and Met5A cells were seeded on 100 nm thick silicon nitride (Si$_3$N$_4$) membranes contained in a 6-well plate (Corning, Costar) at the density of $10^5$ cells/well and covered with 3 mL of standard culture medium. After 24 h, the medium was replaced with a treatment medium containing 1000 µM of CoFe$_2$O$_4$ NPs in the case of fibroblasts [43] or containing 5 ug/mL of crocidolite [33,76] in the case of Met5A cells. CoFe$_2$O$_4$ NPs were supplied by Colorobbia S.p.A. (Vinci-Firenze, Italy) as a suspension in 100% diethylene glycol (DEG). The particles show a diameter of 36 nm in the stock suspension but clustering up to 600 nm in the culture cell medium [77]. Crocidolite Asbestos UICC Standard fibres (SPI#02704-AB) were supplied by SPI Supplies Division, Structure Probe, Inc. (West Chester, PA 19381-0656, USA). Dried fibres were suspended in sterile phosphate-buffered saline (PBS) at a concentration of 10 mg/mL. The fibres were sonicated for 1 h and dispersed by a 5–10 mL syringe with a 21-gauge needle [33].

After 24 h, the treatment was removed and the samples were washed twice with PBS without calcium, magnesium, and sodium bicarbonate and fixed with 4% (v/v) of paraformaldehyde solution prepared in PBS buffer [43].

2.2.2. Ovarian Tissue

The ovarian tissue reported here belonged to a sample grouping already studied [36,78] and was obtained from a patient at the Gynaecological and Obstetrics Department of the Institute of Maternal and Child Health IRCCS Burlo Garofolo, Trieste, Italy. The research on this type of sample had been approved by the Institutional Review Board of IRCCS Burlo Garofolo (Trieste, Italy). The tissue sample was prepared as reported in [36,78], where it was fixed in 2.5% glutaraldehyde (Electron Microscopy Sciences, Hatfield, PA, USA) prepared in 0.1 M of sodium cacodylate buffer (pH 7.2–7.4), and a post-fixation
treatment was performed with osmium tetroxide 1% in the same buffer. The sample was then dehydrated with increasing concentrations of ethanol (50%, 70%, 90%, and 100%) and embedded in epoxy resin (DurcupanTM Fluka) [36,78]. A 2 µm thick section of this was placed on a 4 µm thick Ultralene film for XRF analysis.

2.2.3. Human Spermatozoa

The sample reported here belonged to a sample collection already described [79] and was obtained from healthy volunteers. In particular, the specimens were the discarded residuals from fertilization procedures. After centrifugation (10’ at 300× g), the spermatozoa were fixed in paraformaldehyde (4% in phosphate-buffered solution) for 20 min at room temperature. After washing twice in physiologic solution (NaCl 0.9%, 15’ at 300× g), they were re-suspended in 15 µL of physiological solution (~100,000 cells). For XRF imaging, 2 µL of samples was dropped on Si3N4 windows, let to dry, washed in water, and dried again [32].

2.2.4. Rhodnius Prolixus

The insect was transversally cut at the junction between prothorax and mesothorax segments of the body and fixed with formalin. After that, the insects were embedded in paraffin and then cut in a microtome. A 5 µm thick section was placed on a 4 µm thick ultralene film. More details are given in [80].

2.2.5. Coronary Artery Sample

The coronary artery presented in this paper is part of the research project illustrated in [81]. The coronary artery of a wistar rat was removed, cleaned, and placed in 10% neutral-buffered formalin solution. After that, the tissue was embedded in Paraplast Plus (Sigma-Aldrich, St. Louis, MO, USA) and fixed. The sample was sectioned with a microtome to a 5 µm-thick slice and deposited on a 4 µm-thick ultralene film. All procedures were carried out according to international guidelines of animal protection and were approved by the Ethics Committee on Animal Experimentation of the State University of Rio de Janeiro (Process CEA/018/2018).

3. Results

In this section, we present how the abovementioned TwinMic imaging techniques can be applied in the medical and biological fields, providing new insights in specific scientific questions.

3.1. STXM Coupled with LEXRF

3.1.1. Nanotoxicology

NPs, such as cobalt ferrites, are increasingly being used in the biomedical field, as drug-delivery agents, biosensors, or imaging contrast agents [82,83]. The study of the complex NP–cell interaction is crucial to better understand their uptake mechanism and cytotoxicity effects.

To this aim, the capabilities offered by the TwinMic microscope are particularly suited to study the behavior of CoFe2O4 NPs at the single-cell level [43,77]. STXM sub-micron resolution, combined with LEXRF trace element sensitivity, can provide the simultaneous correlation between cellular changes in shape and morphology and the elemental distributions of both cells and NPs.

Figure 5 shows an example of LEXRF mapping performed on mouse fibroblast cells exposed to NPs. Cells are well visible by the distribution of C, O, Na, and Mg, with a higher level of O and Na in the areas corresponding to the cell nuclei. The co-localization of Co and Fe reveals the distribution of NPs, which appear to be preferentially located in the perinuclear/cytoplasmic region of the cells. Moreover, NPs often appear to coalesce between them, forming aggregates or vesicles. More details can be found in [43,77,84].
Figure 5. Absorption (Abs) and differential phase contrast (DPC) images of mouse fibroblast cells cultured on Si3N4 membranes and exposed to a 500 µM concentration of CoFe2O4 NPs, depicted with the corresponding XRF maps of C, O, Na, Mg, Fe, Co, and Scattering (E0). The maps were collected at 1.5 keV over an area of 40 µm × 70 µm with a 400 nm step size, with a 4 s/pixel acquisition time for XRF. Scale bar is 10 µm.

3.1.2. Reproductive Medicine

The combination of STXM with LEXRF allows for an efficient localization of specific elements in biological specimens. In reproductive medicine, elements such as Na or Mg have been shown to be directly related to the functionality of the biological system. The TwinMic setup offers the capability to localize these elements in oocyte and spermatozoa (Figures 6 and 7). At the moment, we cannot provide a quantitative analysis of the elements present in the biological specimens, which requires suitable reference standards.
with similar composition, hard to find in the soft X-ray regime. However, comparisons between samples can still be made by evaluating the relative variation of a specific element.

**Figure 6.** Absorption (Abs) and differential phase contrast in X (X mom) and Y (Y mom) images of primary oocyte in an ovarian tissue, depicted with the corresponding XRF maps of O, Na, and Mg. The maps were collected at 1.5 keV over an area of 70 µm × 70 µm (STXM images) or 60 µm × 60 µm (XRF maps) with 300 nm and 1 µm step sizes, respectively, with a 7 s/pixel acquisition time for XRF. Scale bar is 10 µm.

**Figure 7.** Absorption (Abs) and differential phase contrast in X (X mom) and Y (Y mom) images of spermatozoa deposited on Si₃N₄ windows, depicted with the corresponding XRF maps of O, Na, and Mg. The maps were collected at 1.5 keV over an area of 80 µm × 80 µm with a 400 nm and 1 µm step size for STXM images and XRF, respectively, and a 10 s/pixel acquisition time for XRF. Scale bar is 10 µm.
In a previous study related to artificial reproductive techniques, we indeed demonstrated that STXM can be of help in analyzing the quality of follicles inside ovary tissue samples and the kind of damage resulting from a slow freezing protocol of cryopreservation, by monitoring the morphology of the cells and the distribution of several physiologically relevant elements such as Na, Mg, and O [36]. In particular, it was found that among other effects, low-quality follicles, frequent in the freeze–thawed tissues, show a high Na level in the ooplasm [78].

Figure 6 shows the absorption and differential phase contrast images of an oocyte in a human ovarian tissue slide together with the corresponding distribution of oxygen, sodium, and magnesium, which was collected in the frame of this research.

Analogously, STXM coupled with LEXRF has been successful in revealing elemental changes in spermatozoa treated with photobiomodulation therapy [85] and in capacitation procedures [32], where elements such as Na and Mg seem to play an important role. Additionally, elemental changes were revealed when sperm cells were exposed to oxidative stress conditions [79]. Figure 7 depicts an example of STXM and LEXRF mapping performed on healthy spermatozoa. The data confirm the capability of our technique in localizing specific elements inside sperm heads and tails.

3.2. Ptychography

Figure 8 of this section shows expected imaging results using far-field ptychography [86] at TwinMic for biological studies. In particular, it shows a reconstruction of diffraction data acquired at 1.02 keV, achieving a resolution better than 50 nm for a sample area larger than 60 × 60 μm², where the specimen is a group of chemically fixed human mesothelial cells (Met5A) exposed to crocidolite fibres. The high quality of the obtained image allows us to better recognize the subcellular structures and to identify the nanofibres internalized by the cells. The spatial resolution achieved with this setup can be seen in Figure S1, showing the reconstruction of a tungsten Siemens Star test reference object with a 20 μm external diameter. As can be seen from Figure S1, panel C, a spatial resolution better than 50 nm could be reached.

![Figure 8](image-url)

**Figure 8.** Panel (A) shows the ptychographic phase reconstruction of chemically fixed mesothelial cells. This is an example of a large field of view (>60 × 60 μm²) examined with a high resolution (approx. 50 nm). Panel (B) and (C) show enlarged ROIs, as indicated in panel (A).
The same type of scan could be subsequently performed across an energy range that covers a specific absorption edge, eventually allowing for XANES analysis.

As ptychography is more complex and less diffused than older standard techniques such as STXM and full-field, such images are unique and of particular interest for the research community.

3.3. Compressive Sensing

Even though compressive sensing XRF can be applied to any kind of sample, it appears to be particularly useful for biological/medical samples, where the specimens are often of irregular shapes or discontinuous by nature (such as cell or alveolar structures in lungs) and with chemical elements of interest located in specific limited areas. On the other hand, for such kind of samples, in order to collect statistically significant data, it may be necessary to scan large areas (millimeters wide) or several samples/patients.

In this section, we thus present two examples that allow us to appreciate the benefits of the compressive sensing approach in biology and medicine.

The first example applies to a specific field of growing interest in biomedical research due to recent and pressing concerns over viruses and pathogens, especially in relation to pandemics and the climate crisis [87,88]. In particular, the study of insects, and more widely, of other arthropods and invertebrates, can bring fundamental insights when dealing with pathogens causing diseases potentially harmful or fatal for humans [89]. Insects are indeed often vector/carriers for the transmission of known pathogens and might also foster the spread of novel virulence factors into existing ones [90,91]. Moreover, many invertebrates possess highly specific virus–host interactions and immune responses, as well as a variety of regulatory mechanisms either to eliminate toxic compounds from their cells, or to balance the uptake of specific elements and intracellular sequestration [92].

The TwinMic setup, combined with compressive sensing acquisition, allows for a more efficient STXM-LEXRF analysis for insect systems, providing both the scan of the entire specimen and the spatial distributions of elements with micrometric resolution.

The example in Figure 9 shows results of a study on Rhodnius prolixus that may be of significance for vector control studies, particularly toward the control of the Chagas Disease [80,93]. LEXRF maps allow the study of tissue/organ-specific accumulation patterns, as well as element distributions in various insect’s life stages, which may have significant implications for toxicology and medicine. More generally, this kind of information can help in understanding insect–virus interactions and the modality of transmission with vertebrates.

This case exemplifies how compressive sensing can vastly improve time-effective acquisition, which is very advantageous for the scanning of large areas or a large sample set. Indeed, by acquiring the XRF signal only in the areas with an absorption signal below a specific threshold, only 21% of the total area was mapped without losing essential information, allowing one to save around 79% of the total time required with a standard LEXRF rectangular mapping. The absorption threshold was defined by acquiring a low-resolution STXM scan with step size of 5 µm. The overall complete STXM scan (Figure 9) coupled with selective LEXRF mapping (Figure 10) lasted 42 h.
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**Figure 9.** Absorption (Abs) and differential phase contrast (DPC) images of a section of a Rhodnius prolixus head [80] acquired at a 1.7 keV excitation energy over an area of 1200 µm × 720 µm with a 1.5 µm step size and probe size, and 50 ms acquisition time. Scale bar is 50 µm.

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The distributions of C, O, Na, Mg, Al, and Fe are shown in Figure 10, which can be correlated to the insect morphological features shown in Figure 9 [80].

The second example of compressive sensing LEXRF concerns the research field of imaging combined with element localization in extended organs, such as arteries and aortic arch [81].

XRF has been shown to be a powerful tool to investigate aortic systems in connection to several medical treatments and diseases, e.g., atherosclerotic plaques [94, 95] radiotherapy and chemotherapy treatments [96] or calcification effects [37, 97]. Moreover, the same technique can reveal interesting connections between the distribution of specific elements and physical and mechanical properties of aortic media, as presented in the work of Z. Qin and coauthors [98].
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Figure 10. LEXRF maps of C, O, Na, Mg, Al, and Fe acquired at a 1.7 keV excitation energy with a 1.5 µm step size and probe size, and 2 s acquisition time. The STXM images were acquired over an area of 1200 µm × 720 µm with a 1.5 µm step size and probe size, while the XRF signal was collected only on selected areas, based on a preselected absorption threshold value in the absorption signal. Scale bar is 50 µm.

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The internal structure of the coronary artery wall is quite complex with different layers of tissue [81]. Collecting the LEXRF information from the entire section instead of a small area may bring several advantages to the medical investigation. However, one of the main issues in fluorescence mapping of such a large area is related to the long acquisition time. The compressive sensing method can thus play a key role in overcoming this limitation. Figure 11 shows the absorption and differential phase contrast images of a whole section of an aortic arch of a hypertensive rat [81], while Figure 12 depicts the corresponding LEXRF signal of C, Mg, and Na elements. The bottom row of Figure 12 also shows a sub-region of whole mapped area, to better highlight the coronary artery structure and elemental distribution. Again, by acquiring the XRF signal only in the areas with an absorption signal below a specific threshold, only 25% of the total area was mapped without losing essential information and saving around 75% of the total time required with a standard LEXRF rectangular mapping.
Figure 11. Absorption (Abs) and differential contrast image (DPC) of a section of coronary artery from a rat [81], acquired at a 1.5 keV excitation energy with a 2.5 μm step size, over an area of 2100 μm × 2000 μm. Scale bar is 100 μm.
Figure 12. XRF maps of C, Na, Mg, and scattering signal, collected on the coronary artery section shown in Figure 11, at 1.7 keV, with a step size of 2.5 µm and a 1 s acquisition time per pixel, only on the areas with an absorption signal below a specific pre-selected threshold. The bottom row depicts a sub-region of the absorption (Abs) and differential phase contrast (DPC) images shown in Figure 11 together with the corresponding LXERF maps of Mg and Na (Panels Mg_K 2 and Na_K 2, respectively). Scale bar in map C is 100 µm, while in panel Abs, it is 50 µm.
4. Discussion and Conclusions

In the present paper, the TwinMic imaging and spectroscopic potentialities in biology and biomedicine were highlighted by showing relevant examples in the life sciences field. The possibility of deploying different imaging modalities coupled with X-ray fluorescence spectroscopy makes TwinMic a very attractive tool for the life sciences community.

Although TwinMic has been successfully employed to provide answers or information for different research topics, there is a continuous trend to improve its capabilities in terms of: (i) sensitivity, by means of new XRF detector systems and new imaging detectors, such as CMOS ones; (ii) spatial resolution, by means of more reliable sample stages, higher-resolution optics systems, and last but not least, by improving the ptychography imaging mode; (iii) chemical element accessibility, by extending its energy range until 4.2 keV, as foreseen in the next years to come.

TwinMic upgrades will also be pushed forward by new possible requirements or wishes from the life sciences community.

Another successful key point of TwinMic, which has not been stressed in the current paper, is that its setup modularity and flexibility (i.e., motors stages and sample holders) allow it to be easily complemented with other synchrotron techniques, such as FTIR spectromicroscopy [42], XRF at higher energies [37,38], and microtomography [34], often by analyzing the very same specimens.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/app11167216/s1, Figure S1: (A) Ptychographic reconstruction of a 20 µm diameter tungsten Siemens Star test pattern. (B) Zoom of the area indicated with the red box in Panel A. (C) Profile graph extracted along the red line indicated in Panel B, showing an achievable spatial resolution better than 50 nm.

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Informed Consent Statement: Informed consent was obtained from all subjects involved in the studies.

Data Availability Statement: The data presented in this study are available on request from the corresponding author. The data are not publicly available due to privacy reasons including an embargo period according to the facility’s scientific data policy.

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