# SMX011 - Superior data quality achieved with the XtaLAB Synergy-i, microfocus Mo source

#### Introduction

Mo Ka radiation is ideal for evaluating large crystal samples, crystals that contain several heavy atoms and crystal samples that are densely packed with heavy atoms. Some examples of such crystals include: minerals, metal clusters and other inorganic materials like perovskites. By studying these sample types using Mo Ka radiation, it is possible to reduce absorption effects and achieve better data quality overall. In addition to studying highly absorbing samples, it is also possible to obtain high-resolution datasets using Mo Ka radiation and perform charge density measurements. Through sophisticated charge density refinements, it is possible to determine the bond connectivity and electron placement for a particular crystal structure in more detail.

The XtaLAB Synergy-i (Figure 1) is a cutting-edge diffractometer equipped with the latest-technology detector, optimized for its small pixel size and sensitivity, and can be configured with up to two bright microfocus sealed tube sources. The Mo microfocus source has its own benefits, and a typical data collection on an organometallic crystal is discussed herein.



Figure 1: The XtaLAB Synergy-i.

# **Experimental**

An organometallic complex containing a first-row transition element was mounted on a nylon loop (Figure 2) and a standard dataset for publication purposes was collected. The data collection details are shown in Table 1.

Table 1: Data collection details.

Crystal-to-detector distance	40 mm
Exposure time	10 seconds
Total collection time	1 hour 48 minutes
Scan width / No. of frames	0.5° / 640
Completeness	99.9% (0.83 Å)
Average I/σ(I)	40.0

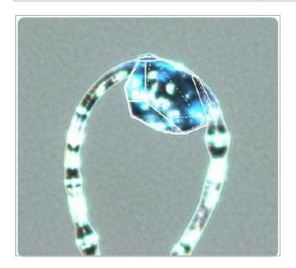


Figure 2: Face-indexed organometallic crystal (0.26 × 0.18 × 0.15 mm).

#### **Data collection results**

Figure 3 illustrates a typical diffraction pattern measured on the XtaLAB Synergy-i with the HyPix-Bantam detector. Highintensity reflections can be detected beyond a resolution of 0.83 Å.

The data was processed using the CrysAlis<sup>Pro</sup> software package, which produced an excellent Rint value. CrysAlis<sup>Pro</sup> also contains powerful, easy-to-use tools for face-indexing that makes it easy to apply face-indexed absorption correction models to every set of data collected using this instrument.

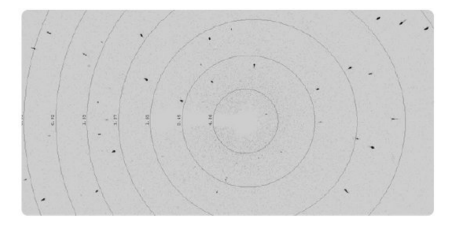


Figure 3: Diffraction image.

 Table 2: Crystal data and refinement statistics.

Chemical formula	$M_{2}C_{13}C1_{2}H_{20}O_{22}$
Space group	Monoclinic, P2 <sub>1</sub> /c
a (Å), α (°)	10.103(1), 90
b (Å), β (°)	10.8119(3), 142.77(3)
c (Å), γ (°)	18.237(2), 90
V (Å3) / Z	1205.3(8) / 2
R <sub>int</sub> [merged data]	2.17%
R1 [merged data]	3.51%
Largest residual peak/hole	0.44 / -0.75



Figure 4: Organometallic structure.

### Conclusion

Publishable results were obtained from this sample in a reasonable time frame, which can be attributed to the cuttingedge technology of the of the XtaLAB Synergy-i. The high flux of the Mo microfocus source, the true digital photon counting technology of the detector, the efficient 4-circle Kappa goniometer and the all-important CrysAlis<sup>Pro</sup> software are all responsible for the superior data quality.

## **Related products**





#### **HyPix-Bantam**

Extremely low noise, compact, detector based on direct X-ra y detection technology.

#### XtaLAB Synergy-i

An entry-level single crystal X-ray diffractometer for Chemic al Crystallography configured with microfocus sealed tube t echnology and a direct X-ray detection detector.