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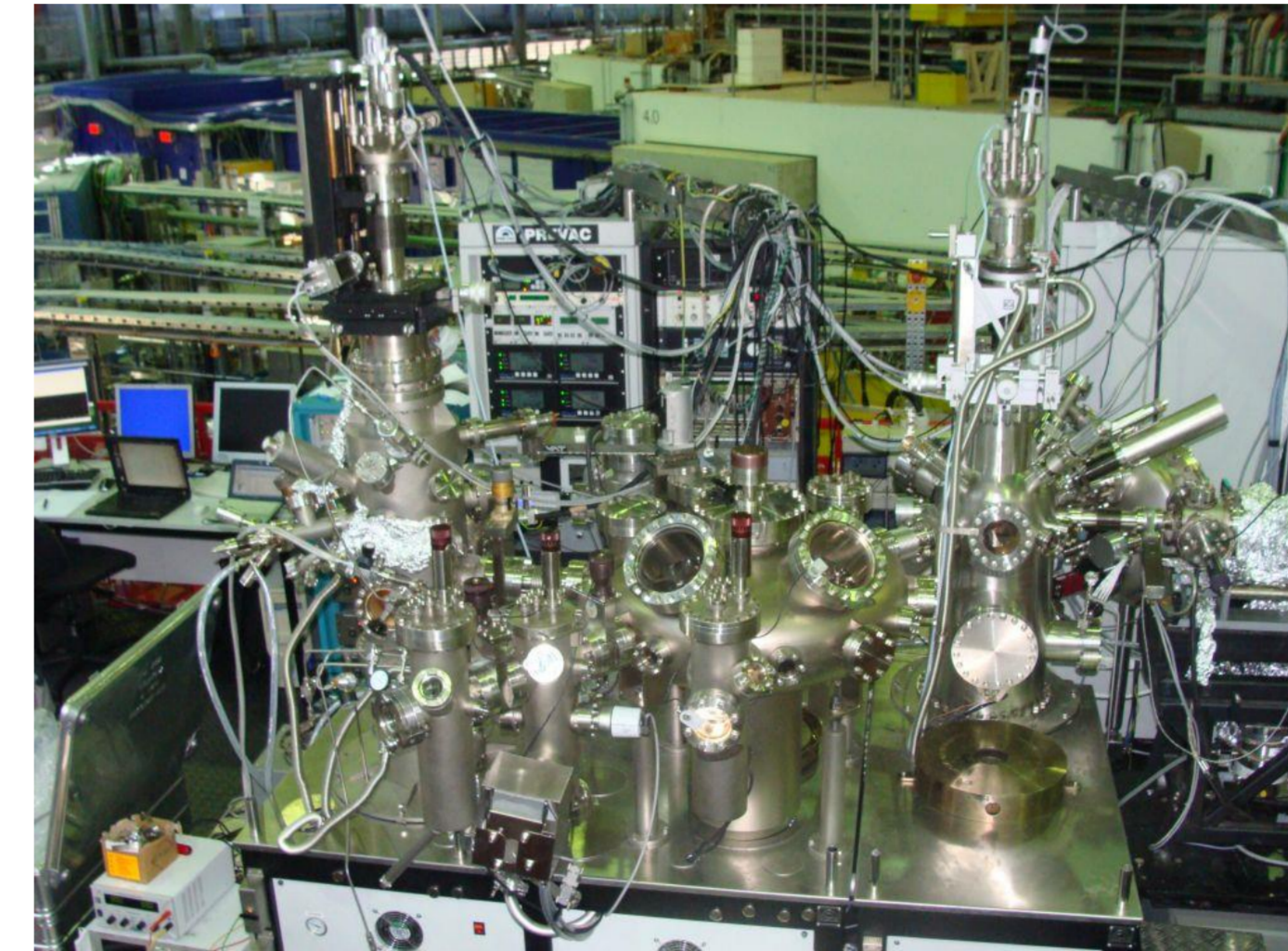
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Motivation

- Pentacene and graphene are promise candidates for organic electronics:
 - behaviour on the interface between them is very important
 - flat orientation of the first pentacene layer is necessary
- Fundamental understanding of the interaction on graphene/metal interface:
 - „weak“ or „strong“
 - Pentacene as a probe molecule
 - Comparison of EG/Ni(111) vs. EG/Au/Ni(111) and h-BN/Au/Ni(111)
- Possible interaction of pentacene with metal substrate through graphene - ?
 - Presence of „surface phase“ similar to chemisorbed pentacene on Ag(111) [Käfer et al, CPL, 442 (2007) 376–383]
- Additional challenge:
 - Evaluation of C K-edge NEXAFS spectra for organic/graphene systems

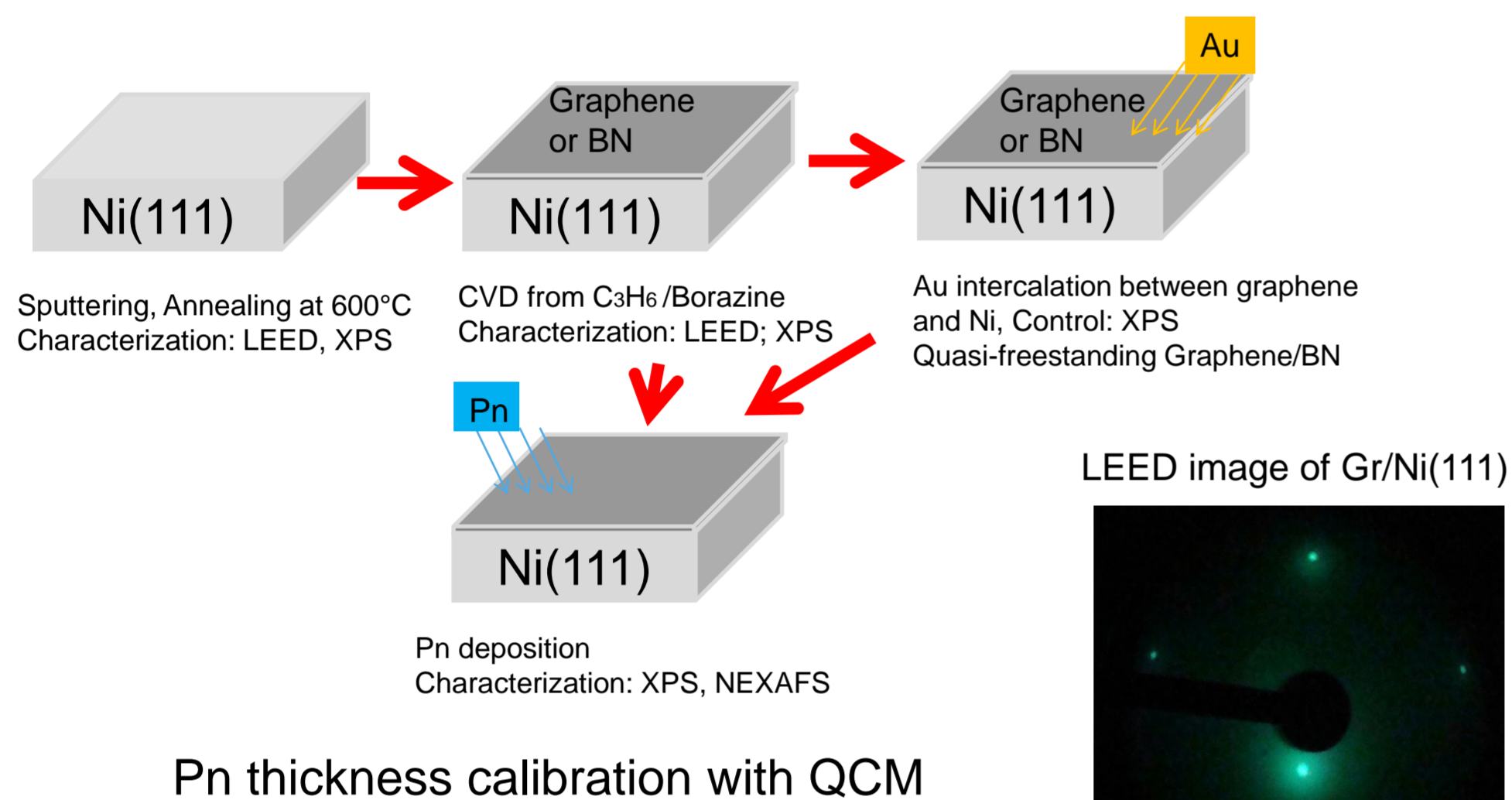
XPS/NEXAFS endstation @ HESGM (BESSY II)



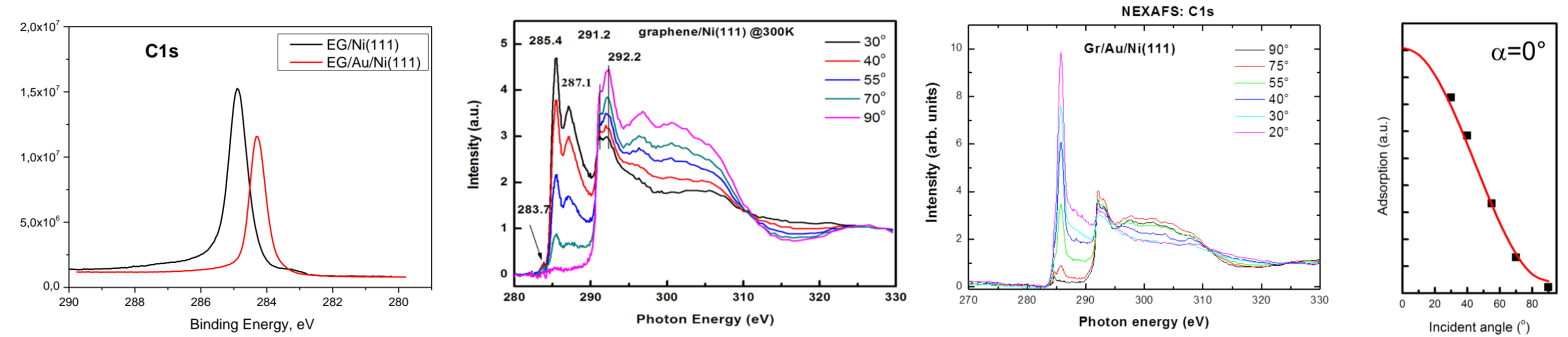
Partial electron yields (PEY) mode for NEXAFS with home-made MCP detector. SRXPS with VG Scienta R3000 analyzer.

Sample preparation and layout of the experiment

Used systems: EG/Ni(111), EG/Au/Ni(111) and h-BN/Au/Ni(111)



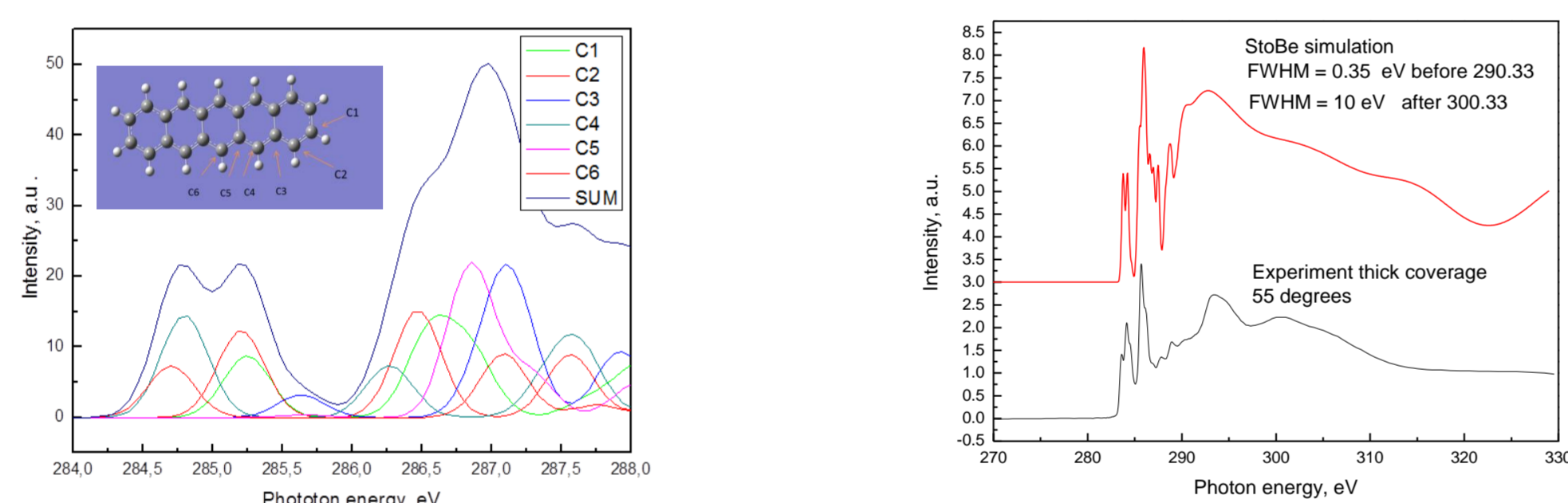
XPS and NEXAFS: Comparison of Gr/Ni and Gr/Au/Ni



XPS (left panel) and NEXAFS characterizations of single layer of graphene prepared by C₂H₄ cracking on Ni(111) surface before and after Au intercalation (central panels). The state at 287.1eV denotes strong hybridization between graphene with Ni(111).

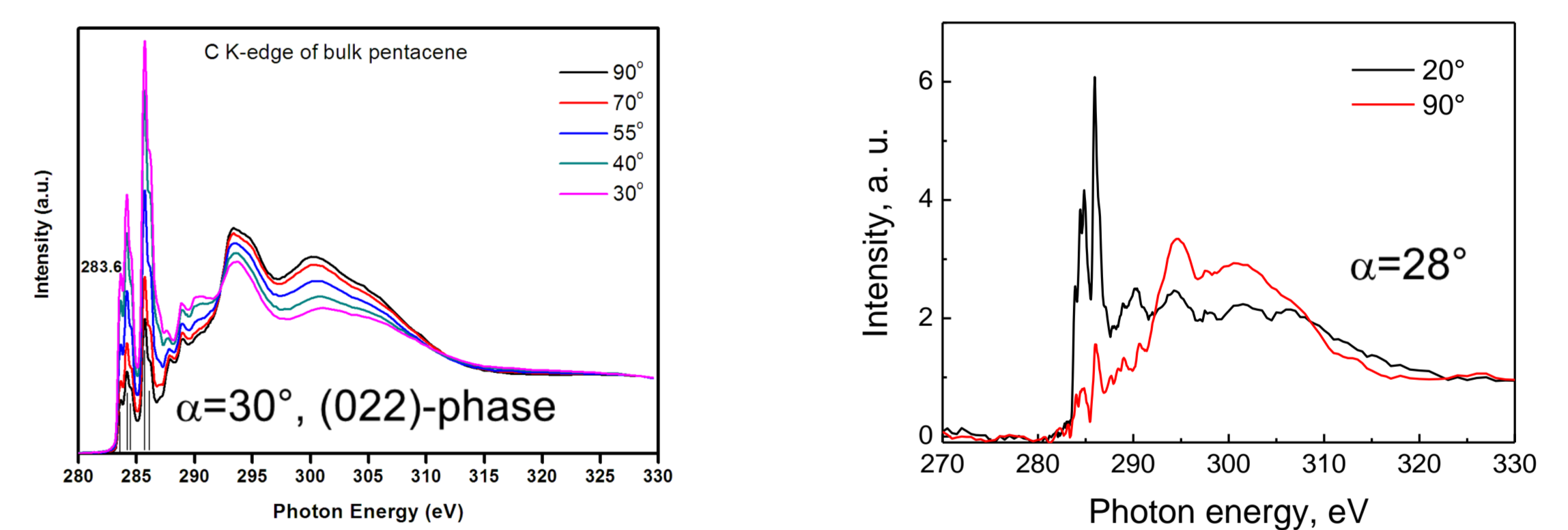
Angle dependence of NEXAFS Intensity (right panel) shows a highly parallel graphene layer on Ni(111).

Simulation of NEXAFS spectra of Pn molecule (in gas phase)

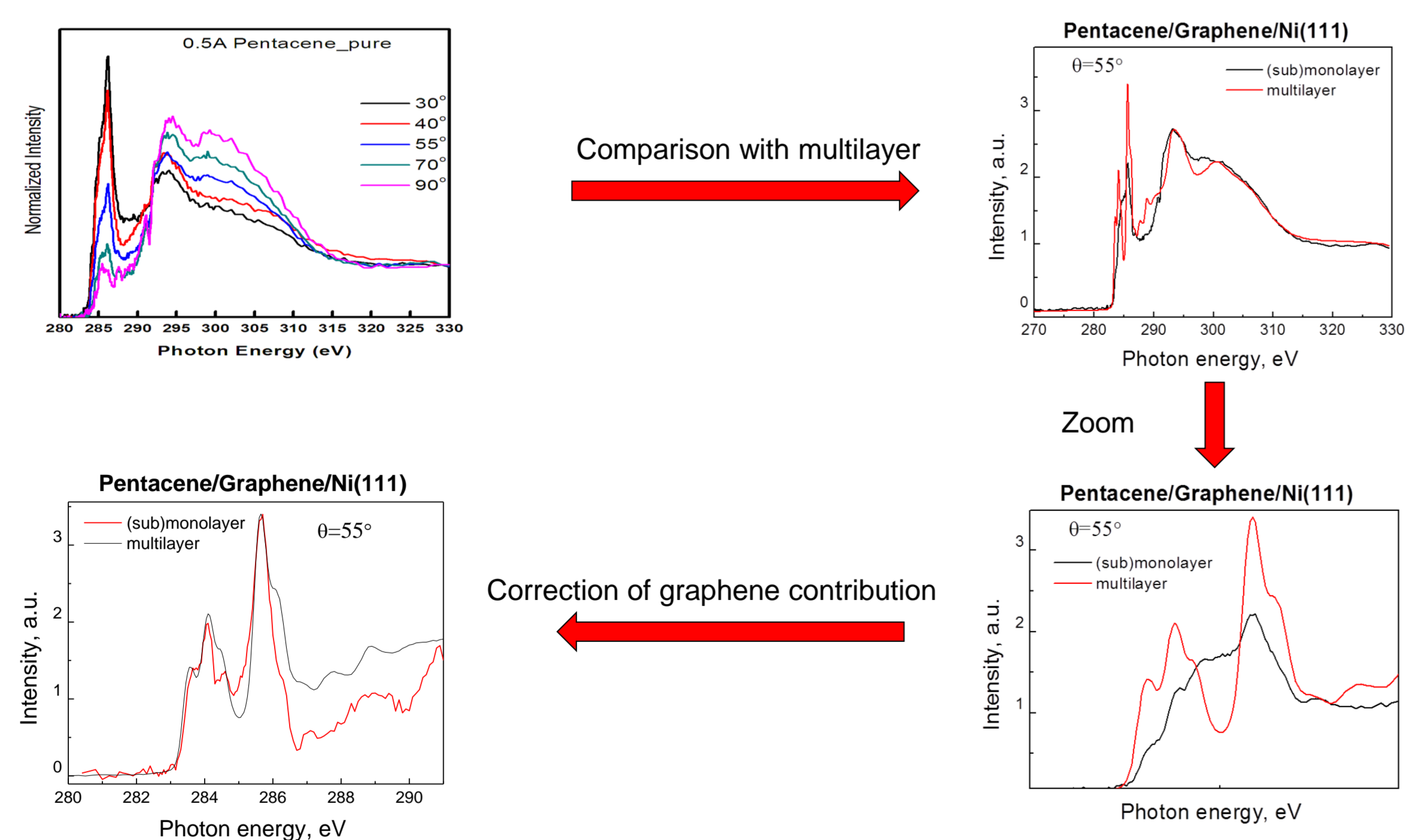


Atomic specific XAS calculations for carbon K-edge of Pn molecule using StoBe package.

Pentacene multilayer on graphene and h-BN

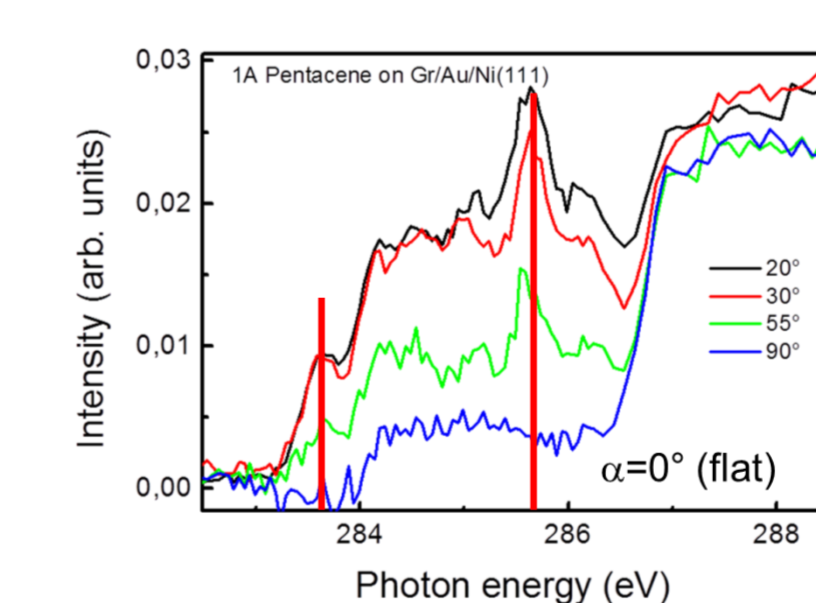


Pn submonolayer on EG/Ni(111): correction of graphene contribution

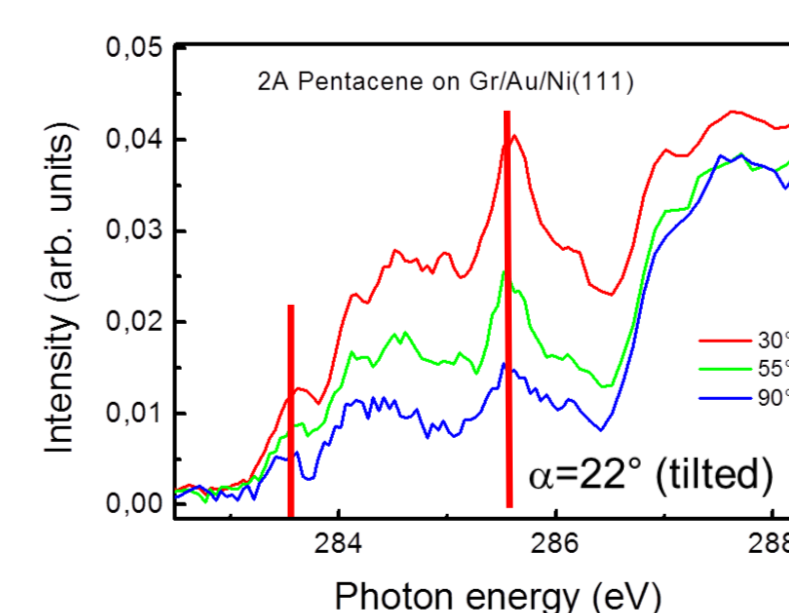


No hybridization – „weak“ interaction! Flat orientation of the first Pn layer!

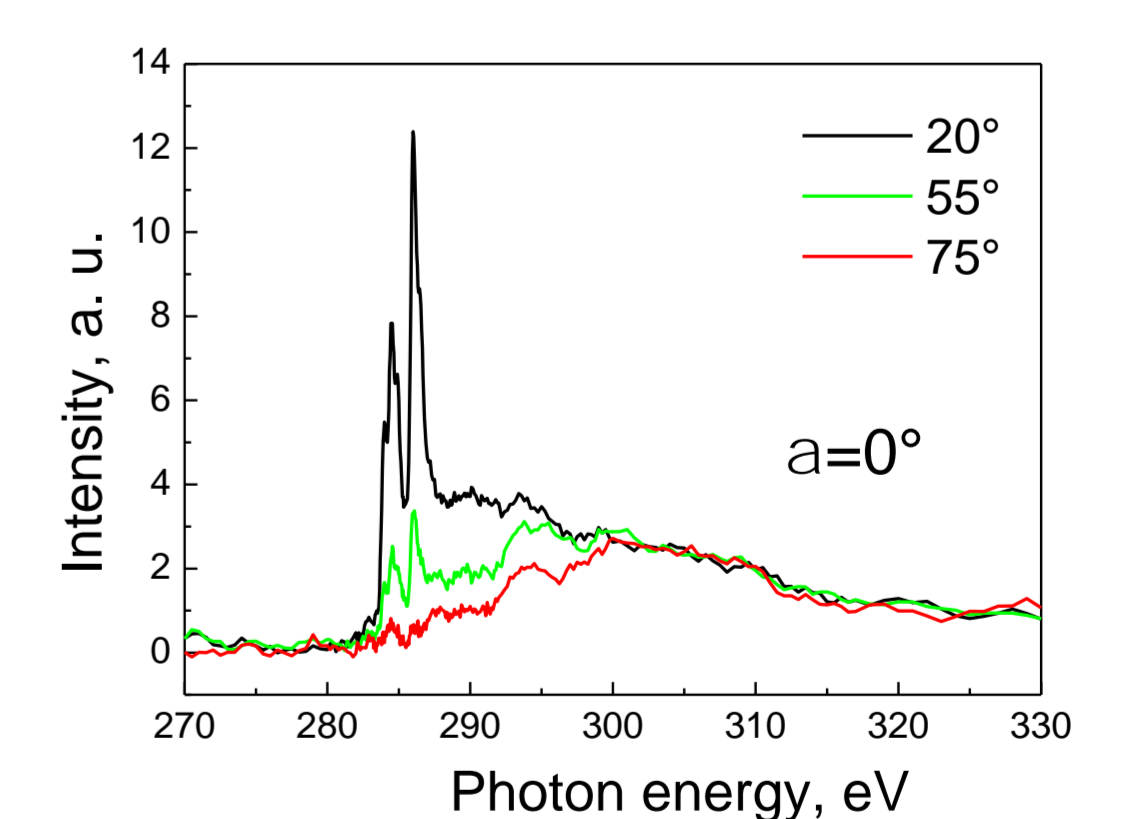
EG/Au/Ni(111) Submonolayer



Growth of the second layer



Pentacene on h-BN: (sub)monolayer



Conclusions

- Flat orientation of pentacene at submonolayer coverages – independent on substrate configuration
- Direct growth of tilted phase already for the second layer
- Weak interaction between pentacene and graphene
- No „surface phase“ was observed – no interaction with metal substrate through EG layer