


Information to be requested from all CA17104 participants:

	
Indicate your Working Group(s) in COST Action17104:	WG4
First Name:	Petko
Surname:	Alov
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Link to webpage with biography:	

Link to webpage with group description:	http://biomed.bas.bg/en/departments/qsar-and-molecular-modelling/
Orcid ID or Scopus ID	
Linkedin	
Expertise relevant for this COST Action:	<i>In silico</i> and bioinformatics approaches in pharmacology and toxicology
Available facilities to conduct work, relevant for this COST Action:	HPC cluster and specialized software for molecular modelling (e.g. MOE, GOLD, Schroedinger, free software)
Materials/Methods that could be shared with other members of this COST Action:	In silico models /predictions of pharmacological and toxicological effects of bioactive molecules

NOTE: By submitting this form to the Grant Manager of CA17104, I agree that this information can be used within the scope of this COST Action (e.g. may be included on the webpage of CA17104).