

**Institute of Pharmaceutical Technology and  
Regulatory Affairs  
Faculty of Pharmacy  
University of Szeged**

# **I. Symposium of Young Researchers on Pharmaceutical Technology, Biotechnology and Regulatory Science**

**Szeged, Hungary**



**31<sup>th</sup> January  
2019**



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**OP-9**

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## **API – excipient interactions in non-biodegradable solid matrix systems**

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Solid dosage forms are still the most preferred types of medicines in the pharmaceutical market. Due to the emerging trend on personalized medicine, pharmaceutical industry faces a new challenge on providing matrix systems with tailorable properties. To fulfil this request a novel approach of pharmaceutical design may be applied with a more detailed investigation of physico-chemical property-based interactions between the drug and the applied excipients. The main aim of this research work is the better understanding of these interactions and fulfilling the requirements of the ‘Functionality related properties of materials’ concept of Quality by Design.

A line of chemically similar APIs and matrix forming agents were mixed and directly compressed with an instrumented IMA Kilian SP300 tablet press. The interactions formed within tablets were studied by FT-IR and NIR spectroscopy, and a custom-made device was used to perform dissolution tests to obtain information about the effects of interactions on the drug liberation kinetics.

The spectral information revealed that hydrogen bonds are being formed between the drug and excipients even in solid state, while investigations during dissolution tests have proven that the strength of interactions have increased due to the formation of polyelectrolyte complexes which affects not just the speed of drug liberation but also the quantity of the liberated drug.

According to the findings it can be concluded that in addition to the physico-chemical properties of the drug delivery system, the drug liberation is fundamentally influenced by the chemical interactions formed between APIs and excipients.