Taste Masking Coating of Mini-Tablets with Cellulose **Derivatives: Formulation and Analytical Challenges**



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Introduction

An important step in patient compliance is palatability enhancement of drugs by improving or masking their taste. Within the methods available, covering the taste by applying a coating layer can be considered as easy to implement. However, one of the major challenges during the development of taste masking coatings is to have a fast feedback on the coating performance during the laboratory trials. We developed an easy taste-masking coating formulation and utilized a screening method to assess the masking property using low volume for dissolution testing with simulated saliva.[1-3]

Materials and Methods

Caffeine mini-tablets (round, 3 mm, 20 mg) were produced on a rotary press (Romaco Kilian Pressima). The taste-masking formulation is based on an enteric polymer (hypromellose acetate succinate: Shin-Etsu AQOAT[®] AS-HG) and a pore former (hypromellose: Tylopur[®] 606). Results were compared with a taste-masking coating premix commercially available. Mini-tablet coating was performed on a coating pan (Solidlab 1 Bosch) with the formulations given in the table. To mimic the oral cavity, the volume of dissolution media was adjusted to the amount of saliva available in the mouth (approx. 5 mL [1]) and quinine (bitter taste perception at 0.000008 M, 2.5 µg/mL) was selected as reference API concentration threshold value for successful taste masking. For low volume dissolution analysis, one coated caffeine mini-tablet was placed in the Ultra-Turrax tube drive vessel (5 mL simulated saliva pH=6.2, 400 rpm). A sample was drawn after 5 min (n=6) and analyzed in a Perkin Elmer Lambda 25 spectrometer (λ =275 nm). Regular dissolution test was carried out in an Erweka DT720 dissolution tester with UV analysis (USP, n=2x25 mini tablets, paddles, 50 rpm, 37 °C, 0.1M HCl, 900 mL, λ = 275 nm).

19.9

42

0.01

Study Outline

Taste-masking coating of minitablets

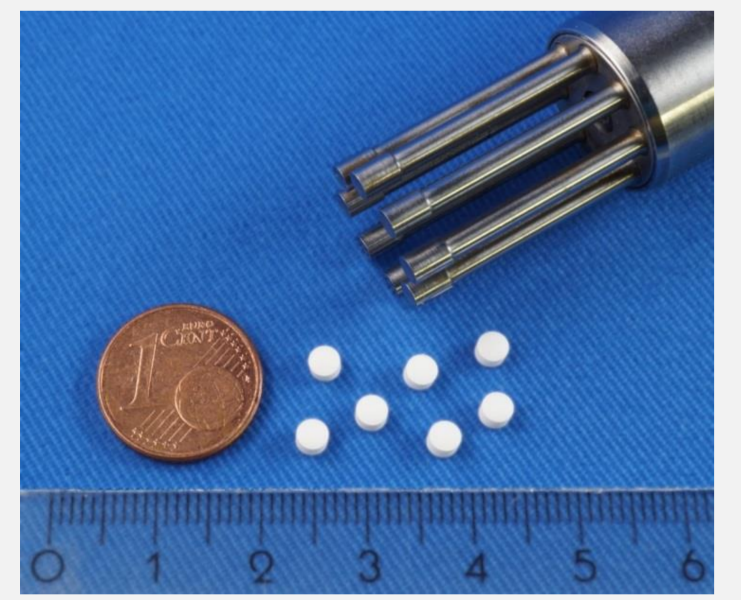
Small volume testing with simulated saliva

> Comparison with quinine & dissolution

Mini-tablets Formulation / Attributes

Material	w/w [%]	Attributes
Caffeine	20.0	Av. Weight [mg]
L-HPC (NBD-021)	15.0	Hardness [N]
Lactose	63.75	Friability [%]
(Flowlac 90)		Disintegration in
Silicon dioxide (Aerosil®)	0.25	water 37°C [min.]
Magnesium stearate [*]	1.0	
Total	100.0	

* Magnesium stearate added before tableting



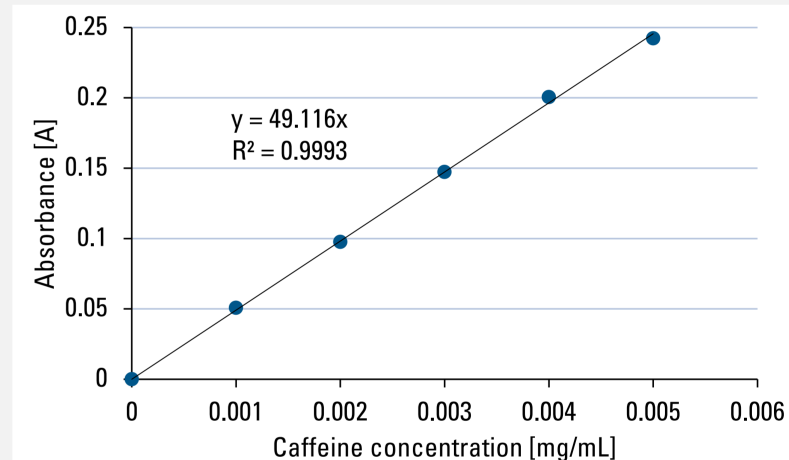
Coating Formulations / Parameters

Material	Content [%]			
	F1	F2	F3	F4
HPMCAS AS-HG (Shin-Etsu AQOAT®)	6.3	3.5	0.7	-
Hypromellose (Tylopur [®] 606)	0.7	3.5	6.3	-
Ammonia (as NH ₃)	0.08	0.045	0.009	-
Eudragit [®] EPO ReadyMix	-	-	-	15.0
Purified Water (H ₂ 0)	92.92	-	92.92	85.0
Total	100.0	100.0	100.0	100.0

F1-F3: Shin-Etsu AQOAT[®] AS-HG was dispersed in H₂O followed by NH₃ (removed during drying). Finally Tylopur[®] 606 was added.

Low-Volume Dissolution: Analysis

Caffeine concentration below 2.5 µg/mL (0.0025 mg/mL) in the low volume dissolution test is considered as a successful taste-masking. Suitability of UV analysis was assessed via a calibration curve.



3 mm mini tablets and multi-tip tool

F4: Eudragit[®] EPO ReadyMix was dispersed into $H_{2}O.$

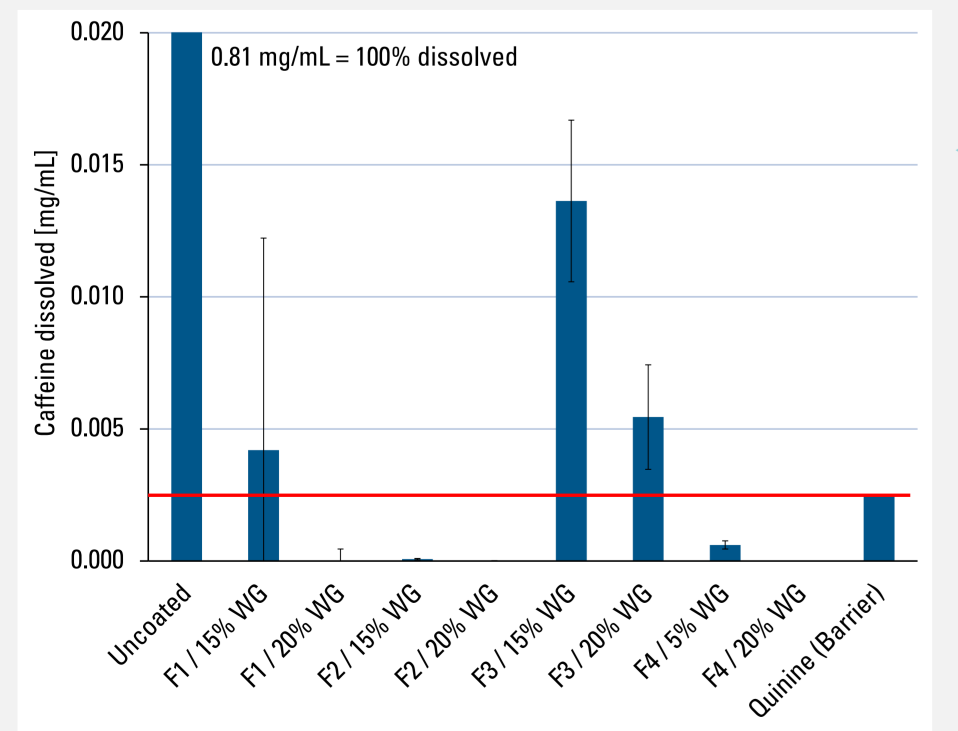
Coating Parameters	F1/F2/F3	F4	
Machine	Solid lab 1 (Bosch)		
Nozzle	0.5 mm Schlick ABC		
Inlet temperature [°C]	54	52	
Inlet air flow [m³/h*kg]	40	40	
Spraying rate [g/min]	5.0	4.0	
Core temperature [°C]	32	33	
Weight gain [%]		Up to 20	

Calibration curve for caffeine in in simulated saliva



IKA Ultra-Turrax tube drive

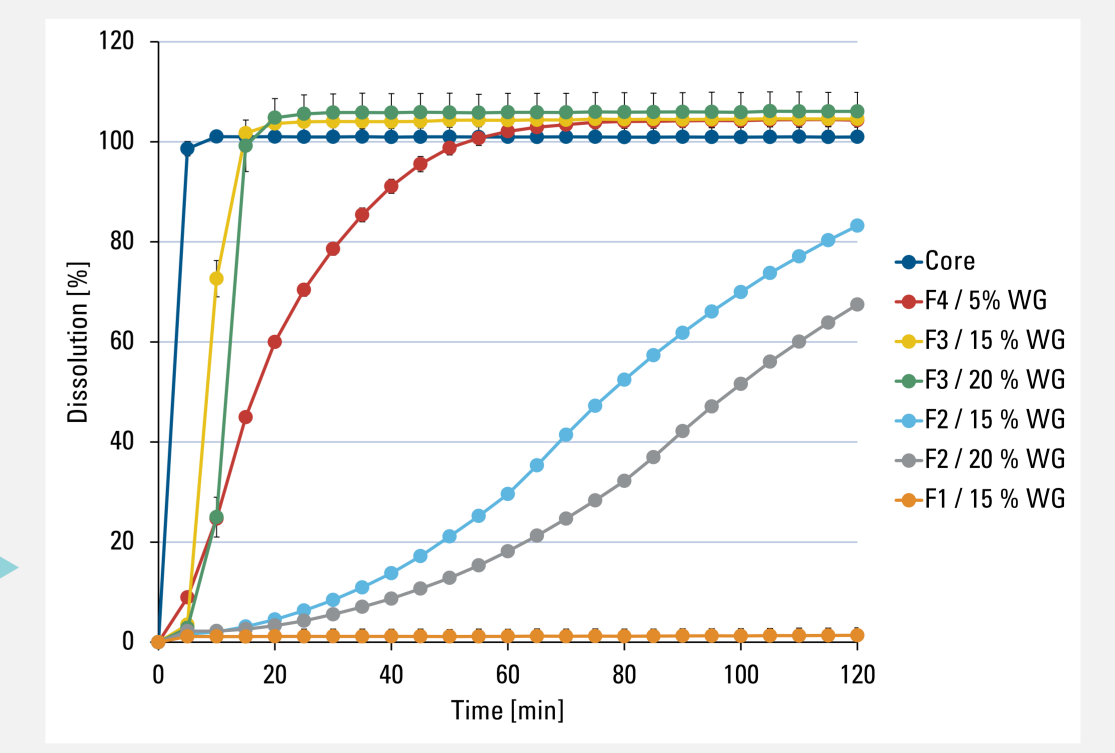
Dissolution Results



Low volume dissolution in simulated saliva:

- Formulations F1 and F2 minimum 15% coating weight gain (WG) needed for taste masking
- F3 slightly above defined threshold
- Taste masking achieved with 5 % coating weight gain for formulation F4

Standard dissolution in 0.1 M HCI



Caffeine concentration after 5 minutes low-volume dissolution test (n=6).

- F1: no dissolution observed
- F2: delayed release of API at pH=1.2
- F3: short lag time of around 5 min. then immediate dissolution
- F4: fast dissolution of the API

Dissolution profile in pH=1.2, 0.1M HCl, n=2x25 mini-tablets

Summary

- Mini-tablet coating was performed without any agglomeration issue.
- Taste masking performance of coatings could be easily assessed by UV analysis and low volume dissolution (5 mL) in simulated saliva (quinine threshold).
- Immediate release of API in gastric fluid was achieved with F3 and F4. Delayed release with F2. Enteric properties with F1.
- Shin-Etsu AQOAT[®] (HPMCAS AS-HG) is suitable for delayed release taste masked dosage forms with adjustable dissolution profile.

References

- 1. Gustafson, T., Taste masking assessment, AAPS **2017**.
- 2. Sona, P.S and C. Muthulingam, Formulation and Evaluation of Taste Masked Orally Disintegrating Tablets of Diclofenac Sodium, Int J Pharmtech Res., 2011, 3, 819-826. Shin-Etsu technical information, 2016, A-045/H-013, A-048/H014. 3.