

Fighting multi-drug resistant tuberculosis: paving the way with different approaches

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The Disease: Tuberculosis

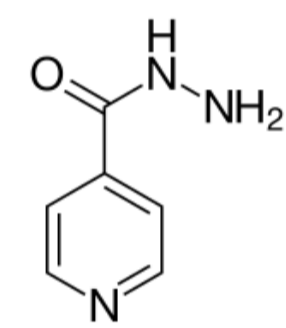
The causative agent: *Mycobacterium tuberculosis*

10 million new cases in 2018

1.5 million deaths in 2018

(Global TB facts, WHO Global Tuberculosis Report 2019 (refers to 2018 situation))

Pharmacological therapeutic approaches



ISONIAZID (INH)

One of the first-line and most effective drugs to treat tuberculosis

Mutations in the multifunctional catalase-peroxidase enzyme *KatG* ⇒ Resistance to Isoniazid

3.5 % of new cases and 18 % of previously treated

Multidrug-resistant (MDR) (isoniazid and rifampicin resistance) or rifampicin-resistant tuberculosis

The development of new effective and low toxicity antitubercular compounds is urgent

TARGET TUB
PTDC/MED-QUI/29036/2017

Synthesis and characterization of a series of INH derivatives

THE COMPOUNDS

ID	Name	Structure
INH-C ₂	N'-acetylisonicotinoylhydrazide	
INH-C ₁₀	N'-decanoylisonicotinoylhydrazide	
N33	(E)-methyl 4-((2-isonicotinoylhydrazono)methyl)benzoate	
N34	(E)-N'-(4-phenoxybenzylidene)isonicotinoylhydrazide	
N33red	methyl 4-((2-isonicotinoylhydrazinyl)methyl)benzoate	
N34red	N'-(4-phenoxybenzyl)isonicotinoylhydrazide	

Previous promising results:

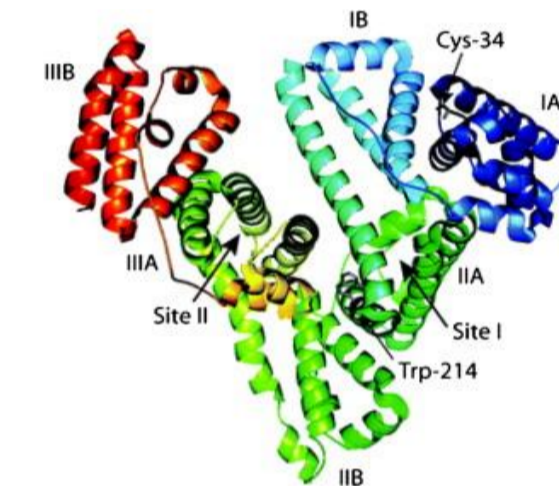
N'-decanoylisonicotinoylhydrazide (INH-C₁₀) 6x more potent than isoniazid against the *KatG* mutated strain, S315T

Evaluating the Therapeutic Potential

Experimental approaches

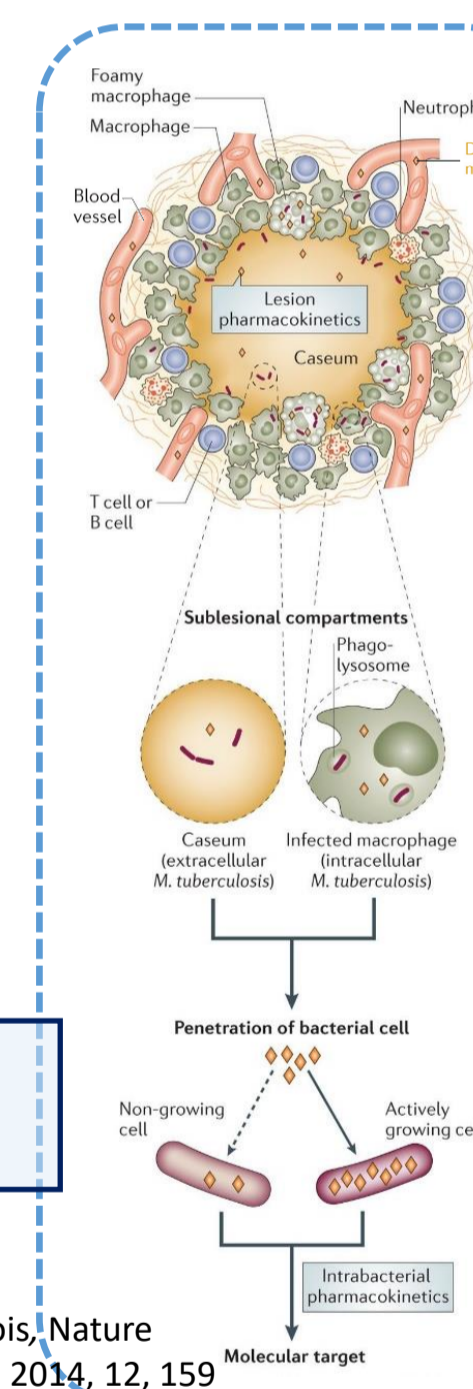
- Ability to bind Human Serum Albumin
- Ability to interact with cellular barriers – membranes
- Toxicological effects on human cell lines

Human Serum Albumin (HSA)



Makes the transport of compounds through the blood stream

The barriers to be crossed



Determination of:

- Dissociation constants (K_d)
- Binding site

Interaction of INH derivatives with systems mimicking:

- plasma membrane of mammals
- cell wall of *Mycobacterium tuberculosis*

V. Dartois, Nature reviews 2014, 12, 159

Cell Viability Assays and Fluorescence Spectroscopy

Evaluating toxicological effects

- Hep G2 Cells
Immortal human liver cancer cell line
- Mtt Assay
assay for assessing cell metabolic activity

RESULT COMPILATION

Compound	K_d (M)	K'_d (M)	Cell Viability IC ₅₀ (μM)
INH	2.72×10^{-3}	5.38×10^{-3}	> 200
INH-C ₂	2.30×10^{-3}	3.50×10^{-3}	n.a.
INH-C ₁₀	4.59×10^{-5}	2.16×10^{-4}	>25
N33	1.66×10^{-4}	1.35×10^{-4}	>25
N34	1.26×10^{-4}	2.30×10^{-4}	>200
N33red	5.07×10^{-3}	1.23×10^{-3}	>200
N34red	1.50×10^{-4}	4.69×10^{-4}	48.5

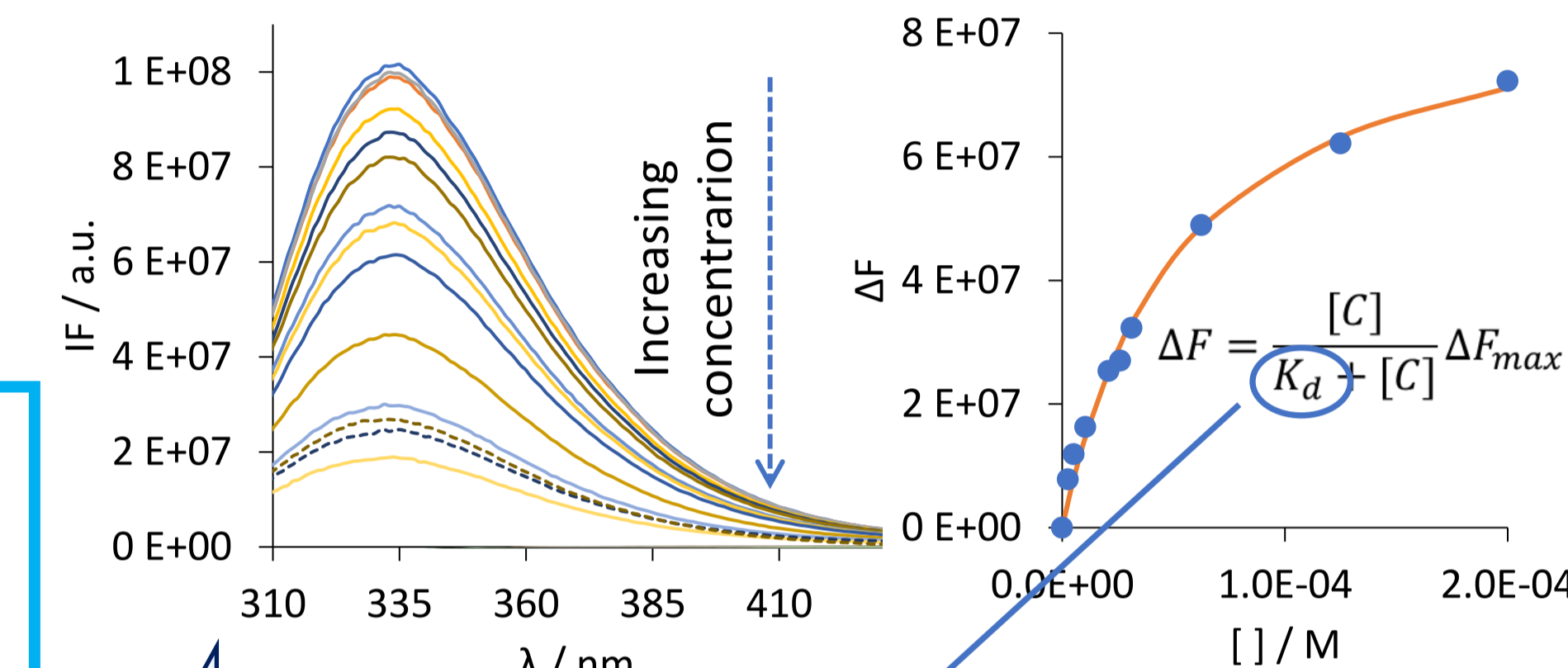
$K'_d - K_d$ in the presence of warfarin

All compounds interact more strongly with HSA than INH, in particular INH-C₁₀, with a K_d 2 orders of magnitude stronger

All compounds are less toxic than drugs currently in pharmaceutical use

Viable alternatives to current therapeutics against MDR tuberculosis

Following the interaction Compound-HSA



Dissociation Constant

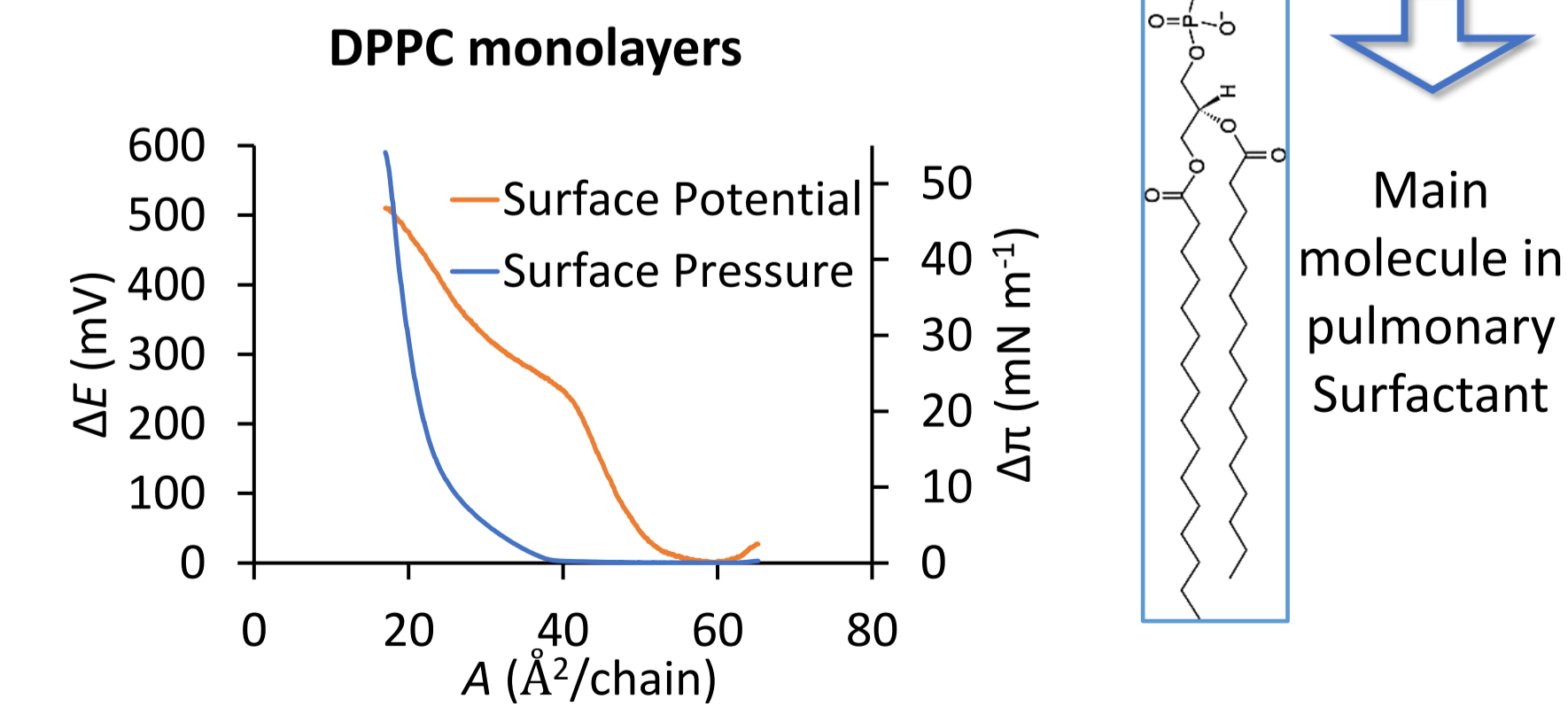
How strong is the Compound-HSA binding

Use of site markers

Binding Site

Warfarin
Binds to Sudlow site I

Monolayer compressibility curves



Next steps

How will the compounds interact with DPPC membranes? Or models of plasma membrane of mammals? Or mycolic acids?

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