

1 PhD scholarship in the MOI III Manchot Graduate School

The Manchot Graduate School „Molecules of Infection“ offers an innovative and structured graduation and research program in a stimulating and interdisciplinary environment of academic biology and medicine.

The scholarship includes a monthly payment of **1.800 € for a period of 42 months** and additional funding for consumables as well as the participation in national and international congresses.

Project 11 (P11): Modulation of cutaneous immune responses by *Candida albicans*

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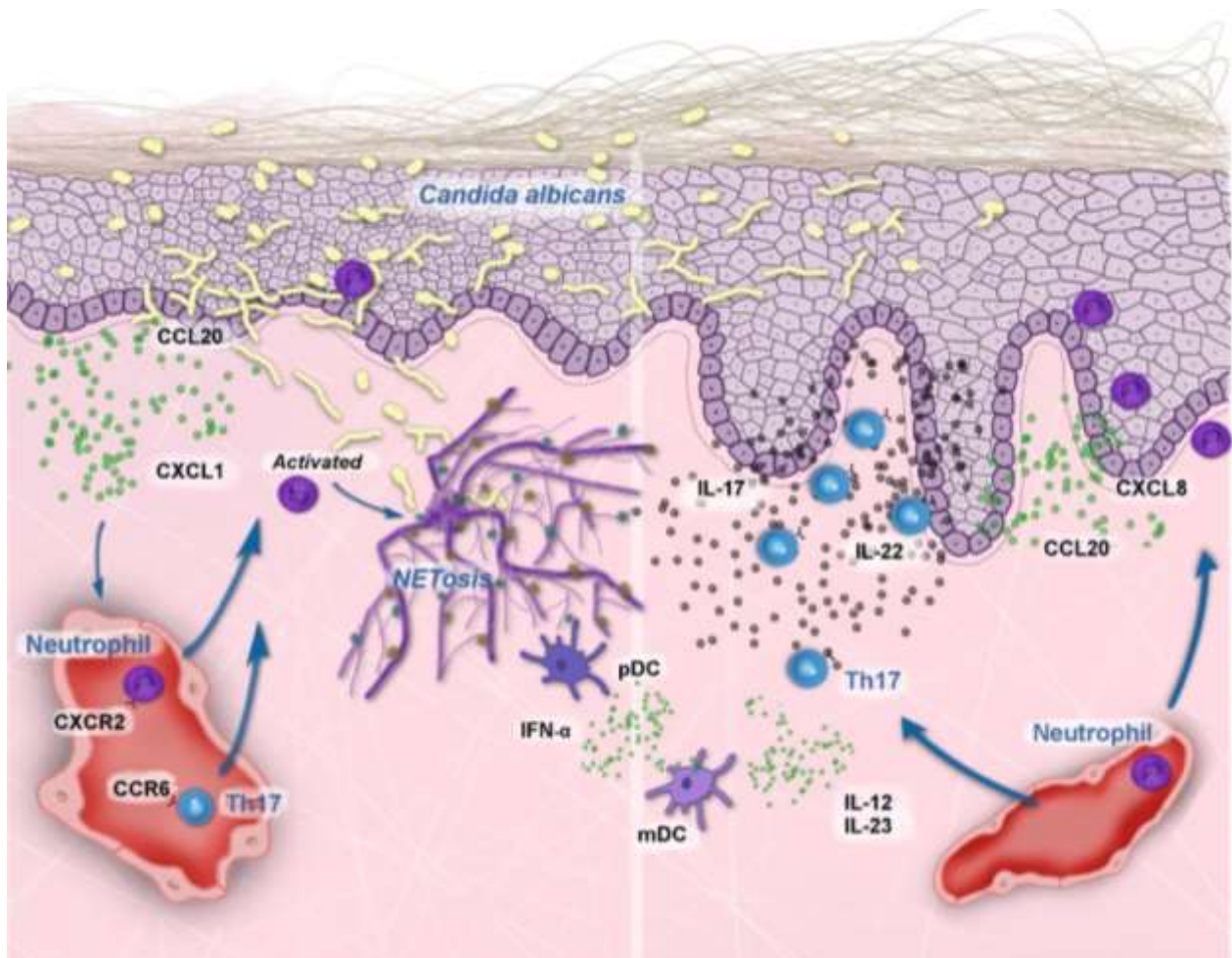
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Findings of the EU-funded project „**Microbes in Allergy and Autoimmunity Related to Skin (MAARS)**“ demonstrate that microbes (bacteria, viruses and fungi) often interact with the host's immune system and influence inflammatory immune responses. In this multi-national consortium, preliminary results indicate a role for *Candida albicans* in the modulation of immune responses within the skin of psoriasis patients.

A principle clinical and so far unsolved question is, why are only distinct skin sites affected by psoriatic inflammation (predilection sites). In psoriasis patients, the skin of extensor parts of large joints as well as the big body folds in the inguinal, submammary and rima ani are affected. These areas often show colonization with Candida species. Immunologically yeasts, like *Candida albicans*, induce T_H17-directed immune responses. Interestingly, genome wide association studies show, that the IL-23/ IL-17 signaling pathway is overrepresented in psoriasis patients and that therapeutic inhibition through neutralization of IL-23 or IL-17 is highly effective in treating the disease.

Hence, the hypothesis of the present project is, that colonization with *Candida albicans* in the large body folds acts as an initial trigger of psoriatic inflammation. During the MOI II-funding period a biobank of human skin specimens of psoriasis patients, Candida-colonization and a *Candida albicans*-modulated psoriasis-like skin inflammation mouse model was established.

In the present MOI III-project, we aim to unravel the *Candida albicans*-associated factors, that promote the development of a psoriatic inflammatory response. The *Candida albicans*-host interaction will be investigated at the level of keratinocytes, neutrophil granulocytes and T cells *in vitro* and *in vivo*.



Further information for your application you may find under the following links:

[01 MOI III Application Instructions 2016](#)

[02 MOI III Application Form 2016](#)

[03 MOI III Application Check list 2016](#)

[04 MOI III Confidential letter of recommendation 2016](#)

If there are futher questions please feel free to contact me via bernhard.homey@med.uni-duesseldorf.de

Looking forward to hearing from you.

Best regards,

Bernhard Homey