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Challenges and opportunities in nasal subunit vaccine delivery : mechanistic studies using ovalbumin as a model antigen

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CHALLENGES AND OPPORTUNITIES IN NASAL SUBUNIT VACCINE DELIVERY

mechanistic studies using ovalbumin as a model antigen

Proefschrift

Ter verkrijging van
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Door

Bernard Adam Slütter,
geboren te Warnsveld
in 1983

Promotoren:

Prof. W. Jiskoot

Prof. J.A. Bouwstra

About the cover:

The application of a nasal vaccine could be accomplished using a spray, which is already common practice for several small molecular weight drugs. The formulation of vaccines however, still needs investigation. Should we formulate antigens in particles, perhaps?

Figure: Cislunar Aerospace, San Francisco, California.

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