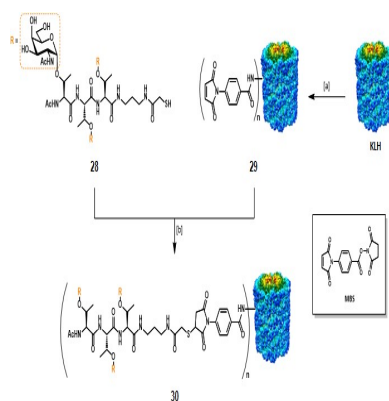
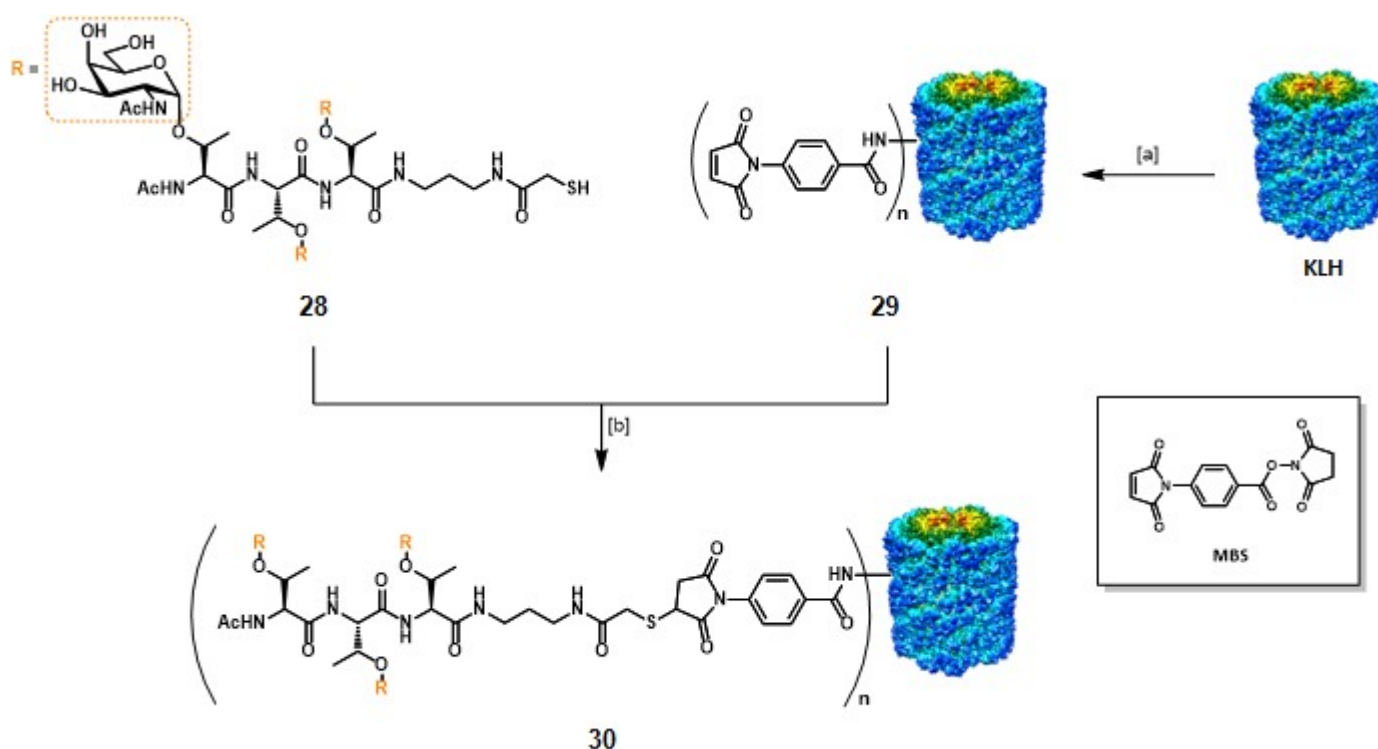




Clustered mono-TACA vaccines based on protein-carriers

Description

In the frame of clustered monomeric vaccines, Danishefsky's research group reported the synthesis and immunological evaluations of a clustered Tn-based conjugate vaccine (**30**), where a cluster of three threonine-linked Tn antigens (**28**) was conjugated to the KLH carrier protein (Scheme 4). Kagan et al., 2005



Scheme 4. Synthesis of clustered Tn-KLH conjugate vaccine **30** by Danishefsky and co-workers. Conditions: [a] MBS, DMF, r.t., 30 min.; [b] DMF, r.t., 3 h.

The clustered B-cell epitope **28**, containing a terminal sulfhydryl group was obtained by chemical synthesis. KLH protein was treated with m-maleimidobenzoyl-N-hydroxysuccinimide ester (MBS) to

obtain maleimidated KLH **29**, which subsequently underwent thiol-maleimide coupling to give vaccine **30**. The epitope ratio (n) was determined by high-pH anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD) assay for carbohydrate after acid hydrolysis ; two batches of **30** were obtained, with an epitope ratio of 201:1 and 648:1, respectively. The clustered vaccine **30** was tested in comparison with its monomeric version ; C57BL/6 female mice were immunized three times with 3 μg of Tn-based vaccine plus 10 μg of QS-21. Previous reports by Kurosaka *et al.* (A monoclonal antibody that recognizes a cluster of a disaccharide, NeuAc $\pm$ 2 $\rightarrow$ 6GalNAc, in mucin-type glycoproteins) and Nakada *et al.* (Epitopic structure of Tn glycoprophorin A for an anti-Tn antibody) suggested that a clustered display of STn and Tn antigens allows an improved recognition by mAbs MLS 102 and MLS 128, respectively. Kurosaka *et al.*, 1988 Nakada *et al.*, 1993 Danishefsky and co-workers demonstrated that clustered Tn-KLH conjugate **30** (Scheme 4) is consistently a better form of Tn for inducing high titers of antigen-specific IgG and IgM, which are able to bind two sources of naturally expressed Tn (i.e. dOSM : desialylated ovine submaxillary mucin, and LSC : a human colon cancer cell line that express Tn but not MUC1).

Following the same design approach, the Danishefsky group carried out a clinical trial on 20 patients with biochemically relapsed prostate cancer by administering clustered TF-KLH conjugate vaccine **31** (Figure 4). Slovin *et al.*, 2005

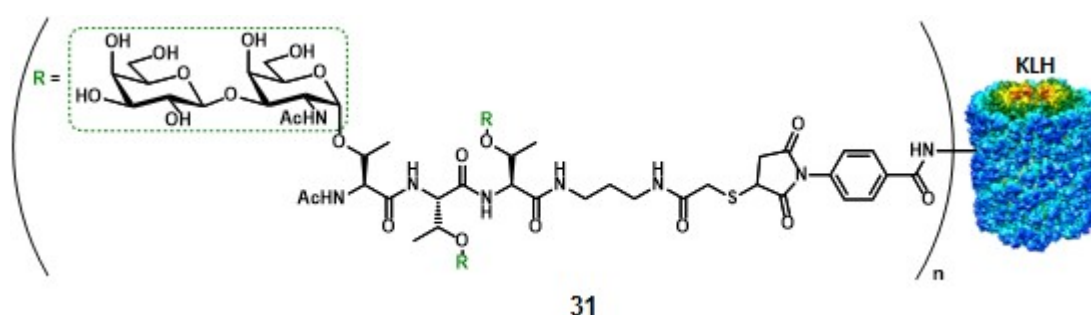


Figure 4. Clustered TF-KLH conjugate vaccine by Danishefsky's research group.

Two batches of **31** were obtained : with an estimated epitope ratio (n) of 466:1 (batch A) and 579:1 (batch B). Four cohorts, each composed of five patients, were vaccinated with different doses of **31** along with 100 μg of QS-21 : 1 μg (batch B), 3 μg , 10 μg and 30 μg (batch A). The toxicity of this vaccine formulation was comparable to that observed in previous trials, where other TACA-KLH conjugates plus QS-21 were administered. All patients developed TF-specific IgM and IgG (mainly IgG1 and IgG3, determined by ELISA). Overall, the IgM and IgG titers at the 1 μg and 3 μg doses were at least as high as at the higher doses. Once again, clustering appears to be a valid way to mimic the way simple antigens, expressed on epithelial but not normal tissues. Scher *et al.*, 2004

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