

ALLERGY, GENETICS

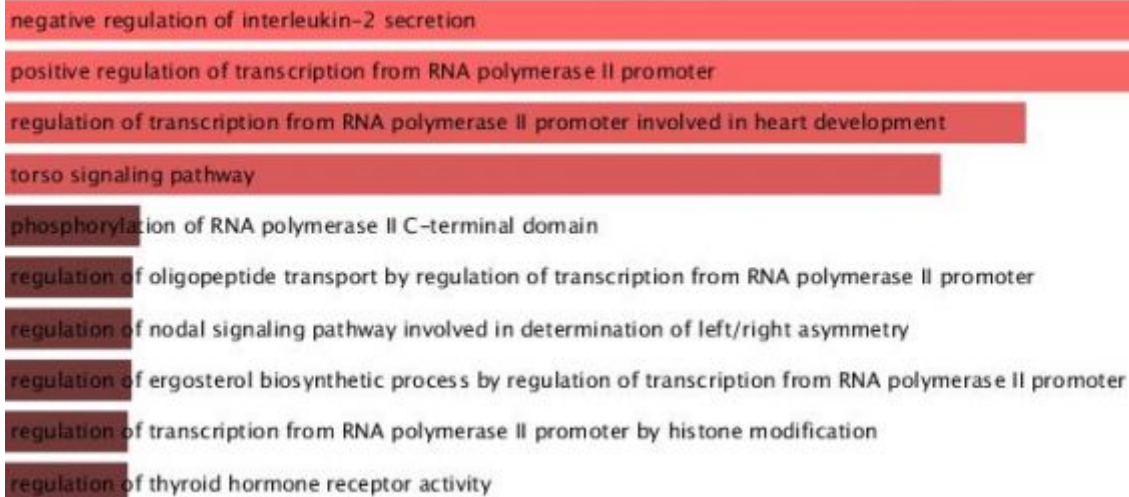
AN UPDATE OF IL2 FUNCTION

8.11.2017

More than 7 years ago, I wrote a blog post that there is [nothing new under the sun](#) predicting the next asthma genetics study for 2020 to include 100K asthmatics.

Ok, I am wrong the [paper](#) appeared already a week ago with 360K asthmatics while basically doubling the number to 136 independent risk variants. It's not an asthma only study as the authors had a rather loose definition of asthma or rhinitis or eczema - is is more about the allergy/atopy complex.

It will be a long time reading and replicating the data while my first interest was to examine the affected gene list as shown in table ST15. Using the online [Enrichr database](#) I get the following result from the GO ontology



This is basically the same result as the authors see in their table ST20.

The highest combined score is with a negative regulation of IL-2. What does that mean? A genetically disturbed pathway predicted by a gene ontology network?

IL2 was discovered in [1976 by Robert Gallo](#) by growing T-cells in culture for more than nine months by stimulating lymphocytes with phytohemagglutinin. ((IL1 was described back in [1972 by Charles Dinarello](#)). Gallo identified T-cell growth factor (TCFG), now known

as interleukin-2 (IL-2) as being absolutely important for a protective immune responses ([Science 1976](#)).

The next important paper was also published in [Science 1984](#) showing the following figure

1,25-Dihydroxyvitamin D₃: A Novel Immunoregulatory Hormone

Abstract. *The hormonal form of vitamin D₃, 1,25-dihydroxyvitamin D₃ [1,25(OH)₂D₃], at picomolar concentrations, inhibited the growth-promoting lymphokine interleukin-2, which is produced by human T lymphocytes activated in vitro by the mitogen phytohemagglutinin. Other metabolites of vitamin D₃ were less effective than 1,25(OH)₂D₃ in suppressing interleukin-2; their order of potency corresponded to their respective affinity for the 1,25(OH)₂D₃ receptor, suggesting that the effect on interleukin-2 was mediated by this specific receptor. The proliferation of mitogen-activated lymphocytes was also inhibited by 1,25(OH)₂D₃. This effect of the hormone became more pronounced at later stages of the culture. These findings demonstrate that 1,25(OH)₂D₃ is an immunoregulatory hormone.*

The importance of 1,25-dihydroxyvitamin D₃ [1,25(OH)₂D₃] in mineral and skeletal metabolism is well established (1). However, the widespread distribution of receptors for 1,25(OH)₂D₃ in tissues not regarded to participate in mineral metabolism has made it apparent that 1,25(OH)₂D₃ plays a wider biologic role

than was previously thought (2). We and others have found that 1,25(OH)₂D₃ receptors are present in normal human monocytes and malignant lymphocytes but are absent from resting T and B lymphocytes (3). Nevertheless, T and B lymphocytes obtained from normal human subjects express the 1,25(OH)₂D₃ receptor when the cells are activated in vitro by mitogenic lectins and Epstein-Barr virus, respectively (3). The mitogenic lectin phytohemagglutinin (PHA) stimulates T lymphocyte proliferation (4) and induces the production of various lymphokines, including interleukin-2 (IL-2), which is important for the growth of T cells (5). In the present investigation we report that 1,25(OH)₂D₃ suppresses IL-2 and inhibits the proliferation of PHA-stimulated lymphocytes.

Peripheral mononuclear leukocytes were isolated by Ficoll-Hypaque gradients from blood samples obtained from normal adults. The cells were cultured for 2 days in medium containing PHA (1 percent) alone or in the presence of increasing concentrations of 1,25(OH)₂D₃ (10⁻¹³M to 10⁻⁸M). The IL-2 content of the media was determined by means of a bioassay that involves titration of the media on the proliferation of the murine cell line CTLL-2; proliferation of the CTLL-2 cells depends strictly on the presence of IL-2 (5). Using this assay method, we found that the media from

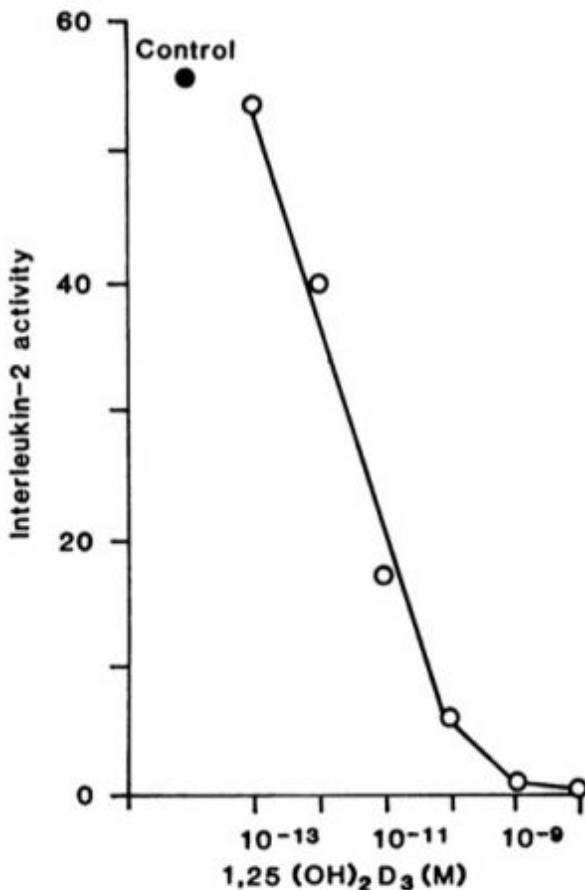


Fig. 1. Effect of 1,25(OH)₂D₃ on IL-2. Human peripheral blood mononuclear cells (1 × 10⁶).

So we have in allergy 2 hits on IL2: by genetics (as show in the new study) and by early vitamin D supplementation (as I reviewed earlier). Maybe we need more clinical studies like the [Zhang 2016](#) study:

In a general population-derived birth cohort, we found that in infants who developed food allergy, cord blood displayed a higher monocyte to CD4(+) T cell ratio and a lower proportion of natural regulatory T cell (nT(reg)) in relation to duration of labor. CD14(+) monocytes of food-allergic infants secreted higher amounts of inflammatory cytokines (IL-1 β , IL-6, and tumor necrosis factor- α) in response to lipopolysaccharide. In the presence of the mucosal cytokine transforming growth factor- β , these inflammatory cytokines suppressed IL-2 expression by CD4(+) T cells. In the absence of IL-2, inflammatory cytokines decreased the number of activated nT(reg) and diverted the differentiation of both nT(reg) and naïve CD4(+) T cells toward an IL-4-expressing nonclassical TH2 phenotype.

Suppressed IL2 is a key for allergy development shown also in experiments by [Bonnet 2016](#):

We previously demonstrated that Tregs can be selectively expanded and activated by low doses of IL-2 (ld-IL-2) inducing immunoregulation without immunosuppression and established its protective effect in autoimmune diseases. In this study, we evaluated the ability of ld-IL-2 to control allergy in an experimental model of food allergy. Ld-IL-2 induced Treg expansion and activation that elicited protection against clinical manifestations of food allergy in two mouse models with OVA and peanut. This clinical effect was lost in Treg-depleted mice, demonstrating the major contribution of Tregs in ld-IL-2 efficacy. Mechanistic studies further indicated that protection from allergy could be explained by a Treg-dependent local modification of the Th1/Th2 balance and an inhibition of mast cell recruitment and activation. Preventive and therapeutic effects of ld-IL-2 were observed over a 7-mo-period, highlighting its long-term efficacy. This study demonstrated that ld-IL-2 is efficient to prevent and to treat allergic immune responses, and thus represents a promising therapeutic strategy for managing allergic diseases.

What we do need now for allergy prevention is a [trials of newborns with low dose IL2](#) and

also more trials refining the IL2 antagonist application of vitamin D.

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