**Pre-diabetes associated M.tb-specific T cell dysfunction**

Pre-diabetes mellitus (PDM) is characterised by higher than normal levels of glucose that are below the threshold for diagnosis of type 2 diabetes mellitus. The prevalence of PDM is steadily increasing globally, and is estimated to increase to almost half a billion by 2030. The role of type 2 diabetes mellitus as an important risk factor for development of active Tuberculosis (TB) disease has been extensively studied. However, there is limited knowledge on the effect of PDM on immune responses in latent mycobacterium tuberculosis (M.tb) infected (LTBI) individuals.

CD4 and CD8 T cells are important players in immunity against M.tb, highlighted by increased risk of developing TB during T cell dysfunction. Kumar *et al.* aimed to determine if PDM results in dysfunction of M.tb specific CD4 and CD8 T cell responses in LTBI individuals.

Kumar *et al.* showed that individuals with PDM-LTBI had significantly decreased CD4+ Th1 T cell functionality compared to LTBI only individuals. This dysfunction was characterised by decreased mono-functional (IL-2, IFN-g) and poly-functional (IL-2±IFN-g±TNF-a±) M.tb– specific T cell responses.
Dysfunction extended to CD4+ Th17 cell functionality, where PDM-LTBI individuals expressed lower levels of CD4+ *M. tb*-specific IL-17A and IL-17F responses. They also observed a similar level of IFN-γ and IL-17 dysfunction in CD8+ *M. tb*-specific immune responses.

Kumar et al. also investigated the impact of PDM on CD4+ Th2 responses. Surprisingly, PDM-LTBI individuals had significantly reduced impairment in production of IL-4 only but not IL-5 and IL-13 in response to non-specific, *M. tb*-specific and flu-specific antigenic stimulation. This suggests a complete dysfunction of IL-4 production in PDM-LTBI infected individuals, which requires further investigation.

In summary Kumar and colleagues, illustrate that PDM is associated with *M. tb*-specific dysfunction in Th1/Tc1 and Th17/Tc17 T cell subsets, as well as overall dysfunction of IL-4 production by T cells. Highlighting the need for research that interrogate the role of PDM as a potential risk factor for active TB, as the prevalence of PDM is steadily increasing.


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