



**With insufficient NAD+ / elevated NADH and / or insufficiency of Mg2+ / Zn2+ / K+ / Cl-, HSD3B1,2 reactions have decreased activity.**

**Low activity / NADPH insufficiency downregulates the entire pathway, reverses HSD17B1,2,3,4 and AKR1C2,3,4. Exercise / resistance training promotes IFN-gamma. IFN-gamma promotes NADPH elevation.**

**Cortisol: cortisone senses low NAD+:elevated NADH, with elevated NADPH:low NADP. Cortisol release is triggered by limbic system response and/or low phosphatidylserine - sensing [low glucose / glycogen, low NAD+:elevated NADH | low P5P]. Cortisol promotes catabolic pathways and inhibits IFN-gamma, glycogen synthase, negatively regulating the IFN-gamma alterations.**

**The primary sex hormone (PSH) senses elevated NAD+:low NADH, with elevated NADPH:low NADP.**

**PSH inhibits cortisol, promoting IFN-gamma and glycogen synthase. PSH promotes anaplerosis to the TCA cycle at alpha-KG and promotes creatine synthesis via AGAT. Creatine promotes GLUT4 and glucose uptake.**

**These combined alterations increase flux through the TCA cycle, eventually converting nitrogen metabolites such as glutamate and glutamine into glycogen, enhancing glycogen storage and IFN-gamma activity.**

**Dietary input and exertion insufficiency dysregulates hormones, glycogen levels and IFN-gamma activity.**

**Figure 4. Pathway Diagram: The role of IFN-gamma in hormonal dysregulation.**  
**ME/CFS: Correcting Chronic Mitochondrial Dysfunction**  
 Author: Joshua Leisk, [DRAFT / INCOMPLETE - may contain errors]