Pellagra: 4 D’s and 8 Points

Abstract
Pellagra has largely been forgotten. This is unfortunate as important lessons are to be learnt for the diseases and social consequences of poverty (and of affluence) that often involve dietary nicotinamide and nicotinamide adenine dinucleotide (NAD) homeostasis. NAD disruption can occur not only from poor diet but from increased consumption of NAD from genotoxic and other stresses. High doses of nicotinamide lead to inhibition of NAD-consuming enzymes and excessive induction of nicotinamide-n-methyl transferase (NNMT) with consequent effects on the methylome giving a mechanism for a new hypervitaminosis-B3.

The history of Pellagra has been largely forgotten even if the 4 ‘D’s’ of Dementia, Dermatitis, Diarrhoea and Death are still taught to medical students. Few realise that a festinating gait, fasciculations of the tongue or myoclonic encephalopathy were first described in the pellagra epidemics and other close mimics of many neurodegenerative and neuropsychiatric diseases were seen.

Pellagrins were prone to both dysbiotic and acute infections explaining the gut manifestations and the high incidence of TB. It was widely believed to be hereditary and certainly ran in families.

The original 18th century European epidemics affected poor peasants on monophagic maize based polenta diets. The early 20th century American epidemic predominantly affected poor blacks (and whites) often working as semi-slave sharecroppers thrown into poverty with the collapse of the cotton market and eating maize, molasses and small quantities of low quality pork.

Pellagra was a systems failure causing premature ageing with evidence of mitochondrial failure, oxidative stress and proteinopathy. Pellagra was curable particularly after the discovery of nicotinamide in the 1940s and is normally sourced from animal products such as Vitamin B3. There were many undiagnosed and untreated cases as the exaggerated sunburn rash (Casal’s necklace) was often not present (“pellagra sine pellagra”) particularly in those with pigmented skin.

Point 1
A disorder that mimicked many neurodegenerative conditions as now classified can have a single and simple dietary cause even when there is evidence for (epi-) genetic involvement, dysbiotic microbiomes, mitochondrial and oxidative stress, and proteinopathy. Dietary nicotinamide with back-up from the degradation of tryptophan on the kynurenine pathway are the precursors to NAD critical for mitochondrial energetics as NADH (and other dehydrogenase reactions), anabolism as NADP(H), and NAD consumer pathways in a “NAD world”. Stress from toxins, whether chemical or microbial, requiring DNA or tissue repair by NAD consuming enzymes such as poly ADP ribose polymerases (PARPs) and SirTunis could have the same pathological result if the dose of nicotinamide is not increased.

Point 2
Pellagra may be being missed. The better known clinical manifestations were recognised as being the “tip of the iceberg” in the epidemics in Europe and the south-eastern confederate states of the USA. Pellaga may be endemic in the millions in poverty who are meat and milk deprived masquerading as Kwashiorkor (“juvenile pellagra”) or “environmental enteropathy” or as poor cognition particularly in black populations who are resistant to the sunburn rash. A community screening test that would not be difficult to develop should be a priority.

Point 3
Tuberculosis (TB), dysbiotic diarrhoeal illnesses and high death rates from acute infections, such as smallpox and measles, disappear as societies increase their meat and nicotinamide intake. This is no mystery as nicotinamide and its analogues, such as Isoniazid, are TB antibiotics and many bacterial toxins interact with NAD-consumer pathways – so being NAD replete would improve host resistance. Topically, COVID-19 interferes with the
tryptophan uptake pathway and therefore NAD levels and T cell and Interferon responses through the Angiotensin-converting enzyme (ACE2) receptor through which it enters cells (that also malfunctions with mutations that lead to Hartnup disease that includes a pellagra-like syndrome). Some clinical manifestations, such as on gut and on cognition or long COVID, could be “forms fruste” or new versions of pellagra.7

Point 4
Reduction in many infections coincided with “meat transitions” and triggered demographic and epidemiological switches toward infertility and auto-immune, allergic and other diseases of modernity. Immune intolerance with changes in T cell subset ratios come about from less use of Indoleamine 2,3-dioxygenase (IDO) and the tryptophan to kynurenine “immune tolerance” pathway as in house production of nicotinamide is no longer necessary.9 Dietary modification of nicotinamide or tryptophan in diet, perhaps in concert with other vitamins such as Vitamin D, could affect the incidence of auto-immune conditions such as Multiple Sclerosis.6

Point 5
Meat transitions and “modernity” have, and by magic, reduced the incidence of premature ageing, dementia, and death. Stunted lives were features of pellagra and in all species there are well described links between NAD metabolism, stem cell health and ageing, alongside resistance to infection, so no magic or much medicine is required. NAD levels fall with age and even further with many diseases of ageing and could respond to supplementation.10-12

Point 6
Early pellagra-ologists, such as Lombroso, have been right in sensing that pellagrians were atavistic examples of the term given that increasing meat intake (let alone nicotinamide supplementation) is linked to induction of the enzyme NNMT. NNMT detoxifies nicotinamide but consumes valuable methyl groups and nicotinamide overload might over-inhibit NAD-consumer enzymes that are metabolic master molecules.13 Nicotinamide’s methylated derivative resembles the dopaminergic neurotoxin MPTP and may, like nicotinamide, be a “double-edged sword”.

Conclusion
Pellagra’s history is well worth remembering given that nobody systematically makes sure that it or “pellagra sine pellagra” is ever “owned” by any one specialty but it should have remained a public health concern doubtless helped by supplementation but not helped by never being a universal policy. Nicotinamide’s potential toxicity in high meat economies was also never monitored over the longterm.

Dietary dosage or nicotinamide supplements may need to be boosted when individuals have certain mutations or are under stress, whether genotoxic or anoxic/metabolic – or restrained if there really is a hypervitaminosis B3 contributing to diseases of affluence. The environmental cost of optimising meat intake would be mitigated by affluent countries eating and consuming meat to the extent of the “third world” both channelling meat to the wealthy. Earlier extremes between the Old World and the New World that had few natural animal domesticates were corrected by the Columbian exchange enabling the rise of the West. However the global South was also unlucky in its meat supply, particularly Africa with a lack of animal domesticates and unhelpful human and veterinary infections in the tsetse fly belt with trypanosomiasis and rinderpest.