

Coronavirus mRNA Vaccine Information

mRNA COVID-19 Vaccines are Fast Tracked

mRNA vaccines were developed in under one year through the federal government program Operation Warp Speed.[1] These experimental products have received Emergency Use Authorization (EUA) from the FDA.[2][3] EUA allows for the utilization of unapproved medical products that may be effective versus one that must demonstrate substantial effectiveness. EAU also does not allow for adequate time to thoroughly evaluate safety outcomes in general or at risk populations because the rushed trials do not include those with multiple comorbid conditions, pregnant/breastfeeding women, children under 16 or provide for long term follow up.[4][5][6][7]

mRNA COVID-19 Vaccines are New Technology

mRNA vaccine technology has never been licensed or approved for use before in The United States. mRNA technology encountered a questionable safety profile in 2017 when it was found that the technology triggered significant side effects.[8] Beyond the direct safety concerns of a never approved new technology, we must also pay attention to the long term immune effects elicited by experimental biological products. Binding versus neutralizing immune effects make a difference in how we respond to viral infection.[9] If a vaccine produces high binding antibodies, the individual is at greater risk upon subsequent exposures as the antibody does not block infectivity, this is known as pathogenic priming.[10] This effect has been observed with a rushed to market dengue vaccine in a human population and in multiple animal studies. [11][12][13][14]

mRNA COVID-19 Vaccines Do Not Protect Against Transmission or Infection

mRNA vaccine trials were not designed to determine if the experimental products prevent reduction in severe COVID-19 infection, hospital admission, ICU admission, death, or transmission. The primary end points in determining efficacy were “subjects without evidence of infection” and “subjects without evidence of prior infection”. These products may reduce mild symptomatic infection in the individual recipient while simultaneously increasing asymptomatic infection and transmission.[15][16][17]

mRNA COVID-19 Vaccines are Free from Liability (PREP Act)

Everyone including government, corporations, and individuals involved in causal relationship with the design, development, clinical testing or investigation, manufacture, labeling, distribution, formulation, packaging, marketing, promotion, sale purchase, donation, dispensing, prescribing, administration, licensing, or use of mRNA vaccines are provided immunity from liability under the federal Public Readiness and Emergency Preparedness Act (PREP Act).[18][19] Individuals may file product injury and death claims with the federal Countermeasures Injury Compensation Program (CICP) where CICP medical staff will determine if they have a covered injury. Since the CICP program’s inception in 2010 90% of claims have been rejected.[20]



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Vaccine mandates, whether from government or private industry, have no place in a free society. We must protect and preserve prior, free, informed consent and medical autonomy.