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CURRICULUM AND EDUCATION ARTICLE

The Double-Edged Sword: How Evolution Can Make or Break a Live-Attenuated Virus Vaccine

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Abstract Even students who reject evolution are often willing to consider cases in which evolutionary biology contributes to, or undermines, biomedical interventions. Moreover, the intersection of evolutionary biology and biomedicine is fascinating in its own right. This review offers an overview of the ways in which evolution has impacted the design and deployment of live-attenuated virus vaccines, with subsections that may be useful as lecture material or as the basis for case studies in classes at a variety of levels. Live-attenuated virus vaccines have been modified in ways that restrain their replication in a host so that infection (vaccination) produces immunity but not disease. Applied evolution, in the form of serial passage in novel host cells, is a “classical” method to generate live-attenuated viruses. However, many live-attenuated vaccines exhibit reversion to virulence through back-mutation of attenuating mutations, compensatory mutations elsewhere in the genome, recombination or reassortment, or changes in quasispecies diversity. Additionally, the combination of multiple live-attenuated strains may result in competition or facilitation between individual vaccine viruses, resulting in undesirable increases in virulence or decreases in immunogenicity. Genetic engineering informed by evolutionary thinking has led to a number of novel approaches to generate live-attenuated virus vaccines that contain substantial safeguards against reversion to virulence and that ameliorate interference among multiple vaccine strains. Finally, vaccines have the potential to shape the evolution of their wild-type counterparts in counter-productive ways; at the extreme, vaccine-driven eradication of a virus may

create an empty niche that promotes the emergence of new viral pathogens.

Keywords Live-attenuated · Vaccine · Virus · Biomedicine · Evolution · Adaptation · Reversion to virulence

Overview

Even students who balk at the very mention of the word evolution can often be persuaded to consider examples of microbial evolution, and thus to become familiar with the processes common to all biological evolution. Having experienced the inescapable logic of the theory of evolution through these “non-threatening” examples, such skittish students are sometimes then willing to accept the role of evolution in the diversification of other forms of life. Students are often especially receptive to learning how evolution can be used to generate beneficial organisms for biomedicine and biotechnology, a subset of the more general category of “applied evolution” (Bull and Wichman 2001), as well as how evolution can sabotage biomedical interventions (e.g., evolution of antibiotic resistance, http://www.pbs.org/wgbh/evolution/library/10/4/1_104_03.html).

In the following review, I focus on the role of evolution in the successes and failures of one such intervention: live-attenuated virus vaccines. These vaccines are live viruses that have been modified in ways that restrain their replication in a host so that infection (vaccination) produces immunity but not disease (Moser and Leo 2010). Thus, killed and protein subunit vaccines are excluded from consideration, as are vectored vaccines in which an immunogenic protein from one virus is inserted into another species of virus (Girard et al. 2006; Durbin and Whitehead 2010). Many of the vaccines currently licensed

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for use in humans, including the Sabin oral poliovirus vaccine (OPV), the combined measles-mumps-rubella vaccine, and the chickenpox (varicella virus) vaccine, are composed of live-attenuated viruses (Table 1). These vaccines have been a tremendous boon to human health (e.g., Widdowson et al. 2009; Roush and Murphy 2007; Zhou et al. 2005; Levine and Robins-Browne 2009, but see (Francis 2010) for some of the failures and inequities in vaccine deployment). To list just a few examples, variola virus, the agent of smallpox, has been eradicated from the world, poliovirus is approaching global eradication (Nathanson and Kew 2010), measles virus has been eliminated from the United States (Roush and Murphy 2007) (but see, Centers for Disease Control and Prevention 2011), and the incidence of chickenpox has dropped dramatically in the United States (Hambleton and Gershon 2005). However, live-attenuated vaccines are not without their risks, which derive primarily from the further evolution of vaccine viruses themselves, as well as the impact of vaccine-driven immunity on the evolution of their wild-type counterparts.

My intention here is to explore both the positive and negative impacts of evolution on the success of live-attenuated virus vaccines and thereby provide cases of microbial evolution with great relevance to human health. I have used many of the subsections below, modified to incorporate appropriate levels of detail, in a wide variety of college classes ranging from Introductory Biology for Non-Majors through to Advanced Virology. Students have found them useful and interesting in their own right and I have often used them as an entrée to a more general overview of evolutionary theory.

Table 1 Live-attenuated virus vaccines currently licensed for use in the United States (www.fda.gov)

Live-attenuated virus(es)	Trade name
Adenovirus types 1 and 4	None
Influenza	Flumist
Measles	Attenuvax
Measles and mumps	M-M-Vax
Measles, mumps and rubella	M-M-R II
Measles, mumps, rubella and varicella	Pro-Quad
Mumps	Mumpsvax
Rotavirus	ROTARIX
Rotavirus	RotaTeq
Rubella	Meruvax II
Vaccinia	ACAM2000
Varicella	Varivax
Yellow fever	YF-Vax
Zoster (Varicella)	Zostavax

Applied Evolution: Generating Live-Attenuated Vaccines by Serial Passage

To date, all live virus vaccines licensed for use in humans in the United States (Table 1) have been generated by one of two “classical” approaches: (a) the Jennerian approach, first utilized by Edward Jenner, in which a related virus from a non-human animal is used to protect against a human virus or (b) the Pastorian approach, first employed by Louis Pasteur, of serial passage (Fig. 1a) (Bazin 2003). Both methods leverage the evolutionary principle that fitness is always specific to a particular environment, and thus high fitness in one environment may come at a cost of low fitness in another.

In serial passage, adaptations that enhance replication of a virus in a novel host or environmental condition may lead to attenuated replication in the natural host or condition. Adaptation is accomplished by infecting a live animal or a flask of cultured cells with a wild-type virus, allowing the virus to replicate until it reaches a useful concentration, harvesting the virus, and then using these progeny virions to initiate another infection in a new animal or culture (Fig. 1a). Pasteur was the first to use this technique, passaging rabies virus in rabbits in order to attenuate it for dogs and humans (Plotkin 2011). Most subsequent live-attenuated viruses have also been generated by passage through a novel host or hosts. The vaccine for yellow fever virus, a pathogen that is maintained in African and South American primates under natural conditions, was created via passage through Asian macaques, mouse embryo tissues and chick embryo tissues (Fig. 1b). The vaccine for measles virus, which in nature exclusively infects humans, was generated by passage through chick embryo cells (Anonymous 1980), and the vaccine for varicella, another virus that infects only humans during natural transmission, was achieved through passage in guinea pig cells (Fig. 1b). Additionally, adaptation to replication at lower temperatures is often a desirable evolutionary outcome. The cold-adapted influenza virus was passaged in chicken kidney cells in temperatures descending from 36 to 25°C (Fig. 1b), with the goal of generating a virus that could replicate in the upper respiratory tract but not deeper in the lungs (Maassab and DeBorde 1985).

Because evolution relies on the random occurrence and subsequent spread of rare beneficial mutations, the number of passages and the number of conditions needed to achieve satisfactory attenuation varies considerably (Fig. 1b). For example, the varicella vaccine virus was generated by just 28 passages in three different cell lines, while the yellow fever vaccine required hundreds of passages in five different conditions (Fig. 1b). At first blush, it may seem startling that viruses can undergo substantial evolution in just ten or twenty passages. However, viral evolution is

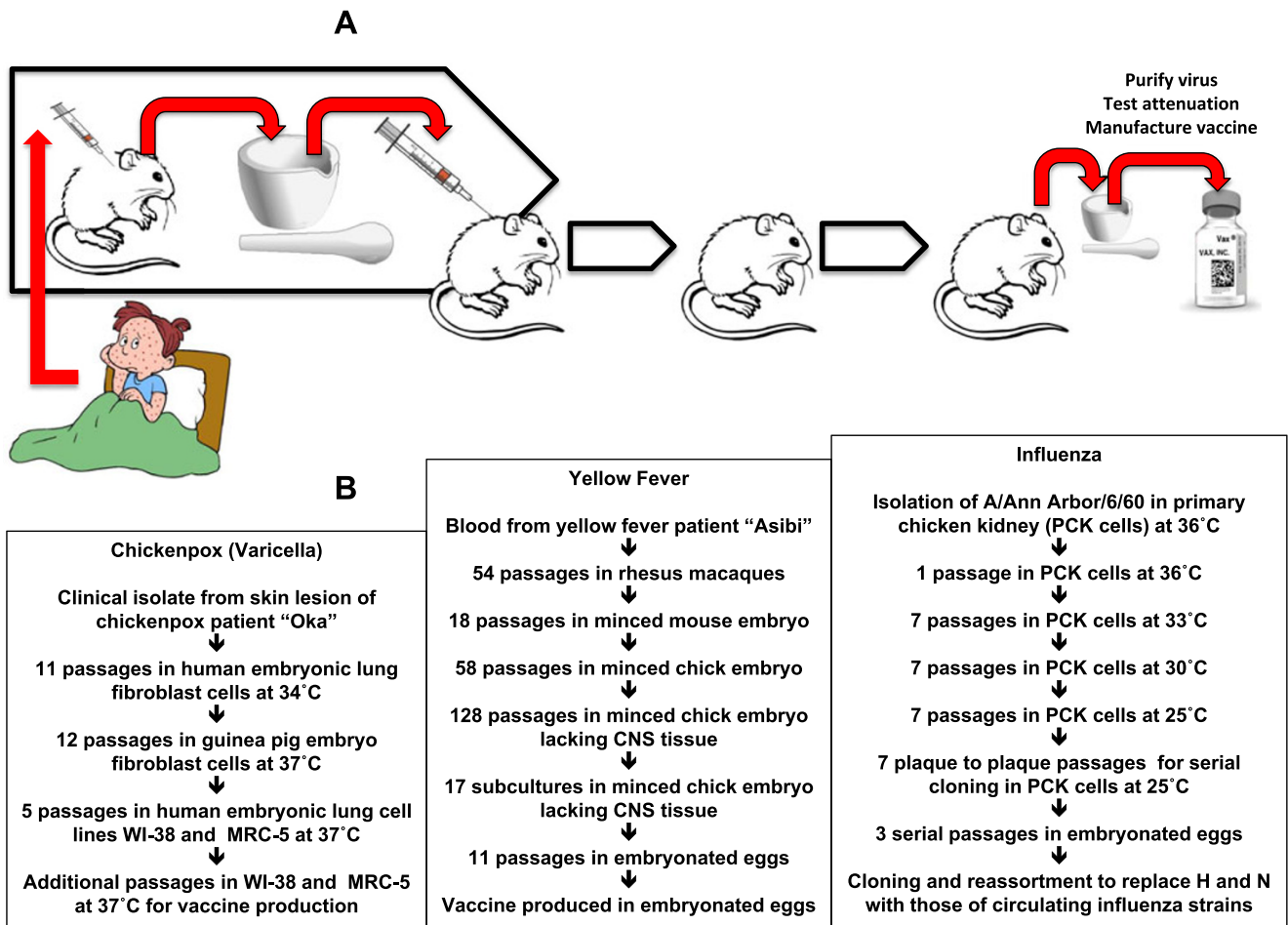


Fig. 1 a The Pastorian approach for attenuation by serial passage b Passage history of three live-attenuated virus vaccines derived by serial passage: the yellow fever virus 17D vaccine strain (Monath et

al. 2008), the cold-adapted influenza A/Ann Arbor/6/60 vaccine strain (Maassab and DeBorde 1985), and the chickenpox (varicella) Oka vaccine strain (Hambleton and Gershon 2005; Takahashi et al. 2008)

known to proceed at a breakneck pace due to the unique features of viral genetics and viral population dynamics (Moya et al. 2004; Domingo et al. 2008).

Viruses have genomes composed of RNA or DNA. Due to the lack of proofreading in RNA-dependent RNA polymerases, RNA viruses have extraordinarily high rates of mutation. The average RNA virus genome experiences one mutation per 10,000 nucleotides per replication, a rate of mutation almost a million times faster than the average bacterial genome (Sanjuan et al. 2010). Additionally RNA genomes tend to be very small, ten kilobases or less (Moya et al. 2004), and thus on average RNA genomes accumulate approximately one mutation per genome per replication. The rate of mutation in RNA genomes is so high that RNA viruses may exist as a *quasispecies*, a swarm of related genetic variants whose interactions determine attributes of the population (Lauring and Andino 2010). DNA viruses show a much greater range in genome size, from 3.2 kilobases (hepatitis B virus; Liang 2009) to 1.2 megabases

(mimivirus; Claverie et al. 2009). Intriguingly, the rate of mutation in DNA viruses shows an inverse correlation with genome size, and small single-stranded DNA genomes may mutate at a rate comparable to RNA genomes (Sanjuan et al. 2010; Duffy et al. 2008). The pace of adaptation depends not only on genomic mutation rate but also on the rate of replication and final population size of those genomes (Moya et al. 2004). Most viruses replicate rapidly within their hosts and some can achieve quite staggering population sizes: for example, three days after an infectious mosquito bite, West Nile virus reaches titers greater than 10^{11} plaque-forming units per ml of serum in certain bird hosts (Komar et al. 2003), a number that greatly exceeds the seven billion humans currently alive and that is roughly equivalent to the number of stars in the Milky Way. Thus, viruses easily outstrip the evolutionary pace of any other living organism, and this allows the evolution of live-attenuated vaccines over a relatively small number of serial passages.

Vaccines Gone Wild: Reversion to Virulence in Live-Attenuated Vaccines

In the classical methods of attenuation, a vaccine virus's phenotype determines its genotype. Passage proceeds until satisfactory attenuation is achieved, irrespective of how many passages or how many mutations this may entail. Indeed, with the notable exception of the OPV polio vaccine strains (Kew et al. 2005), the mutations responsible for the attenuation of most live-attenuated vaccines have not been fully characterized. However, it is clear that in some live-attenuated viruses, the number of attenuating mutations is quite small. Each of the attenuated poliovirus types 1, 2, and 3 strains in OPV carries only a few (two to six) major attenuating mutations (Kew et al. 2005). Given the rapid mutation rates of all viruses, but particularly RNA viruses, it is not surprising that some vaccine viruses revert to virulence. About one in every 750,000 children receiving the first dose of OPV experiences vaccine-associated paralytic poliomyelitis attributable to reversion of one of the three strains (Kew et al. 2005). This propensity to revert is one of the reasons that OPV was replaced with the inactivated poliovirus vaccine in the United States, hence OPV is not listed in Table 1. Such reversion may occur as a result of back-mutation of attenuating mutations, compensatory mutations elsewhere in the genome, or as discussed in the next section, recombination. Conversely, the more attenuating mutations in the genome of a live-attenuated vaccine, the less likely reversion will be. To date, the cold-adapted influenza vaccine, with attenuating mutations in four of the eight genome segments (Maassab and DeBorde 1985), has never reverted to virulence in a vaccinee (Tosh et al. 2008).

Recently, quasispecies heterogeneity per se has also been implicated in changes in virulence (Kenney et al. 2011; Sauder et al. 2006). Sauder et al. (2006) passaged mumps vaccine virus in either monkey kidney or chicken embryo fibroblast cells. They found that these passaged strains showed additional attenuation in rats and that this phenotypic change was associated with changes in quasispecies heterogeneity rather than fixation of any individual mutation.

Several factors can increase the likelihood that a live-attenuated vaccine will revert to virulence. First, the loss of a subset of attenuating mutations during vaccine manufacture can accelerate reversion. Often the cell lines approved for vaccine manufacture differ from the cell lines in which the serial passage of the vaccine was conducted, thus changing the selection pressures on the virus. Complete reversion to virulence would be detected during safety testing and such lots would be eliminated, but if attenuation is enacted by the epistatic effect of multiple mutations (Burch et al. 2003), and a subset of these are lost during vaccine manufacture, then complete reversion would require fewer subsequent mutations. A recent study by

Victoria et al. (2010) searched for back-mutation of attenuating mutations in lots of eight live-attenuated virus vaccine strains, including OPV, using deep sequencing, and found none. However, previous studies of the OPV have detected variants lacking particular attenuating mutations in vaccine lots (Kew et al. 2005).

Second, reversion is more likely the longer vaccine virus replication persists and the higher the virus titer achieved in an individual vaccinee. Extended replication offers more mutational fodder and more time for natural selection to occur within an environment (the host). Such selection is likely to favor virus variants that replicate to higher titers or in a greater range of tissue types than the original vaccine virus; i.e., viruses that are more similar to the wild-type phenotype. Both overall levels of replication and breadth of tissue tropism may be positively correlated with virulence. Valsamakis et al. (1999) experimentally passaged a measles vaccine virus in human tissues engrafted into mice and observed reversion to virulence. Moreover OPV has been shown to replicate to high titers over long periods of time in some immunodeficient vaccinees, resulting in reversion to virulence (Kew et al. 2005; Odoom et al. 2008). These immunodeficient individuals also excrete polioviruses over long periods of time, increasing the likelihood of fecal-oral transmission to unvaccinated individuals in the community.

Such transmission from vaccinees to unvaccinated individuals is a third contributor to reversion to virulence. Although live-attenuated vaccine transmission was initially considered a bonus of live vaccines, a “free” vaccination, it is now clear that person-to-person transmission imposes strong selection for a return to the wild-type phenotype. Recognizing this, polio researchers divide vaccine-derived poliovirus (VDPV) isolates into three categories: (a) iVDPVs derived from immunodeficient patients, (b) cVDPVs that show evidence of sustained person-to-person transmission, and (c) aVDPVs from non-immunodeficient patients or environmental sources that are not associated with an outbreak (Kew et al. 2005). Predictably, virulent vaccine-derived viruses have caused substantial outbreaks in countries where so few children are vaccinated with OPV that herd immunity, the protection of susceptible individuals that occurs when pathogen transmission is “short-circuited” by vaccinated individuals (Fine et al. 2011), is never generated (Kew et al. 2005). Surprisingly, virulent revertants, presumably derived from long-term OPV excretors or from contact with individuals from countries that continue to use OPV, are also detected in wastewater in countries where only the inactivated poliovirus vaccine is utilized (Roivainen et al. 2010; Zurbriggen et al. 2008), suggesting that cessation of vaccination could result in re-emergence of virulent poliovirus in these regions.

OPV is affordable, and vaccination prevents not only polio disease but also poliovirus transmission, making it

an excellent vaccine for the eradication of poliovirus. However, its tendency to slip the leash of attenuation and spawn virulent, vaccine-derived polioviruses through back-mutation as well as recombination (discussed below) has led to the “oral polio vaccine paradox” (Heinsbroek and Ruitenbergh 2010): the same agent that eradicates a disease may serve to reintroduce it! One solution to this paradox is a concurrent, worldwide conversion to inactivated polio vaccine (Heinsbroek and Ruitenbergh 2010; Minor 2009), which would require an unprecedented feat of global coordination.

Hopeful Monsters: Recombination Among Live-Attenuated Vaccine Strains and Between Vaccine and Other Circulating Viruses

In several licensed (Table 1) and candidate (Durbin and Whitehead 2010) live virus vaccines, multiple strains of a single species or multiple species are administered concurrently. Moreover, vaccine viruses may be administered simultaneously with a natural infection of a conspecific or heterospecific virus. Both situations create potential for recombination between viral genomes. Virus genomes recombine by one of three general mechanisms: (a) break and repair in DNA genomes, (b) polymerase template switching in RNA genomes, and (c) reassortment of segments in segmented RNA genomes (Condit 2007). All three mechanisms require co-infection of a single cell by two different parental genotypes. Because many vaccine viruses undergo limited replication relative to wild-type viruses (e.g., Hambleton and Gershon 2005; Monath 2005; Monath et al. 2008; Whitehead et al. 2007; Condack et al. 2007; Miki and Chantler 1992), it would initially seem unlikely that more than one virion would enter a given cell. Moreover, viruses possess a diverse array of mechanisms to prevent additional viruses from entering an infected cell, a process known as superinfection exclusion (Zou et al. 2009 and references therein). However, it is increasingly clear that for some viruses, coinfection occurs at rates greater than expected by chance when viral density is very low. Such coinfection enhancement suggests either that a small fraction of appropriate host cells are actually susceptible to infection (Dang et al. 2004; Cicin-Sain et al. 2005; Smith et al. 2008) or that viruses may possess mechanisms to enhance coinfection at low viral densities (Joseph et al. 2009).

Recombination contributes to the reversion to virulence and subsequent circulation of OPV strains. Such recombination can occur between vaccine viruses, between vaccine and wild-type polioviruses, and between vaccine polioviruses and other species of enteroviruses (Kew et al. 2005). Recombination can occur at multiple locations in the

genome, and multiple recombination events can contribute to the formation of a single genome. An extreme example of this process is a penta-recombinant VDPV, showing five crossover sites between types 2 and 3 polio vaccine strains, that was isolated from a child suffering from poliomyelitis (Zhang et al. 2010). Recombination between vaccine and wild-type viruses has also been documented for numerous virus vaccines for veterinary pathogens, including canine parvovirus (Mochizuki et al. 2008), infectious bursal disease virus (He et al. 2009), bovine herpesvirus 1 (Thiry et al. 2006), and infectious bronchitis virus (Estevez et al. 2003). Recombination is troubling not only for its contribution to virulence, but also because of its potential to generate entirely new species of viruses—western equine encephalitis virus, for example, is the product of recombination between eastern equine encephalitis virus and a Sindbis-like virus (Weaver et al. 1997; Hahn et al. 1988).

Too Much of a Good Thing: Competition and Facilitation Among Strains in Multi-Strain Vaccines

Vaccine viruses can exhibit quite different dynamics when administered in combination with other vaccine viruses than when administered alone. At one end of the spectrum, the individual components of a multi-strain vaccine can show “interference,” in which one or more strains replicate to lower levels or stimulate a poorer immune response in a combined vaccine than a single-strain vaccine. Interference can be the product of immunodominance, direct competition for cellular or viral replication machinery, or immune-mediated apparent competition. Such interference has long been recognized; Sabin and his colleagues (Sabin et al. 1960) pointed out that the individual components of OPV were more efficacious when administered individually (as monovalent vaccines) than when administered together in a trivalent vaccine. Subsequently, multiple instances of interference among live-attenuated vaccine viruses have been documented (Nascimento Silva et al. 2011 and references therein). At the other end of the spectrum, facilitation, in which the replication or immunogenicity of attenuated viruses is enhanced in combination, could also occur. While this phenomenon has not been documented in vaccine viruses to the best of my knowledge, previous studies have certainly shown facilitation between unrelated viruses during concurrent infection (Cicin-Sain et al. 2005; Agrawal et al. 2002; Allan et al. 2007; Allan et al. 2000), thus a similar interaction between vaccine viruses is possible. While the outcomes of interference and facilitation have been described, the mechanisms driving these dynamics are poorly understood. However, as the number of new live-attenuated vaccines targeted to the already overscheduled child continues to increase, it is becoming

increasingly important to gain a mechanistic understanding of the ecological interactions among these attenuated viruses.

Lessons Learned: Evolutionary Considerations in the Rational Design of Live-Attenuated Vaccines

Reverse genetics has offered a new approach for the generation of vaccines, namely rational vaccine design. In this approach, genotype determines virus phenotype. Potentially attenuating mutations are chosen based on available knowledge of the molecular biology of the virus. These mutations are engineered into a recombinant genome that is inserted into a cell, where it generates a recombinant virus. The attenuation of the resulting virus is assessed in a relevant model, and if it is either over- or under-attenuated, then the mutation may be modified or additional mutations may be added.

While rational vaccine design is not a case of applied evolution, to be successful it must be informed by evolutionary thinking. In particular, due consideration must be given to preventing reversion to virulence. This may be ensured by the generation of large mutations (particularly deletions), numerous mutations with epistatic interactions, or chimeric viruses in which the structural proteins of a well-attenuated virus are replaced with those of the virus targeted for immunization. These strategies have been employed in the generation of candidate vaccines for dengue (Durbin and Whitehead 2010) and chikungunya viruses (Kenney et al. 2011). Alternatively, virus genes responsible for counteracting host antiviral responses may be deleted (Kim et al. 2011), or genes in the genome may be rearranged to deoptimize translation and genome synthesis (Lim et al. 2006). As an additional safeguard against reversion, mutations that specifically prevent transmission may also be sought (e.g., Kim et al. 2011; Hanley et al. 2003). Rational vaccine design may also minimize competition among multiple strains of the same virus (multivalent vaccines) by introducing the same mutation into each of the strains, as has been done with candidate vaccines containing all four serotypes of dengue virus (Blaney et al. 2006).

In addition, evolutionary thinking has led to novel approaches to enacting attenuation. As one example, Vignuzzi et al. (Vignuzzi et al. 2008), reasoning that quasispecies diversity enhances RNA virus fitness, have generated polioviruses in which the fidelity of the polymerase is significantly enhanced. Not only are such viruses attenuated, but, because their rate of mutation is lower, they are less able to revert to a less faithful polymerase. In another example, researchers have taken advantage of the tendency of viruses to exhibit codon pair bias: within all of the options in the degenerate genetic code, viruses tend to

use a subset of pairs of codons to encode specific pairs of amino acids. Deoptimizing these codon pairs by inserting different codons that retain the original amino acid sequence results in attenuated influenza (Mueller et al. 2010) and polioviruses (Coleman et al. 2008). This strategy involves hundreds of mutations across the genome and has been called, poetically, “death by a thousand cuts” (Coleman et al. 2008). Clearly, it will stand as a significant barrier to reversion to virulence. Thus, evolutionary thinking is leading to entirely new ways to engineer viral genomes to achieve attenuation.

Under Pressure: How Selection by Vaccine-Generated Immunity Can Shape Evolution of Wild-Type Viruses

Viruses evolve in response to natural herd immunity, and it stands to reason that sufficiently high levels of vaccine-induced immunity could also shape virus evolution (Boni 2008). Evolution of wild-type viruses in response to deployment of live-attenuated vaccines has been observed in several veterinary viruses, including avian metapneumovirus (Catelli et al. 2010; Cecchinato et al. 2010) and avian influenza virus (Park et al. 2011; Lee et al. 2004). The effects of such vaccine-driven evolution on virus virulence and transmissibility in unvaccinated animals have yet to be determined. However, theoretical studies have raised the concern that “imperfect vaccines,” vaccines that do not induce sterilizing immunity but rather modulate disease or transmission, could select for increasing virulence in the wild-type virus targeted by vaccination (Gandon et al. 2003; Mackinnon et al. 2008; Gandon and Day 2007; Andre and Gandon 2006; Ganusov and Antia 2006; Massad et al. 2006). Intriguingly, Manuel et al. (Manuel et al. 2010) recently reported that imperfect vaccination can *prevent* reversion to virulence in a simian-human immunodeficiency virus (SHIV). They generated a SHIV carrying an attenuating mutation and then infected both naïve and vaccinated monkeys with this virus. The attenuated SHIV reverted to virulence through back-mutation in the naïve, but not the vaccinated, monkeys. The investigators attribute this effect to the overall decrease in SHIV replication in the vaccinated monkeys, suggesting that imperfect vaccines may act as a brake on the evolution of their wild-type counterparts.

The ultimate goal of many vaccination programs is the eradication of specific wild-type viruses. This goal has been achieved for variola virus, the agent of smallpox, is within reach for poliovirus (Kew et al. 2005) and perhaps measles virus (Castillo-Solorzano et al. 2011; Moss 2009), and remains a tempting target for many existing and candidate vaccine programs. While there is no question that eradication represents a quantum leap for public health, some thought must be given to the empty niche that eradication

creates. Recent increases in monkeypox have been attributed to the eradication of, and cessation of vaccination against, smallpox (Rimoin et al. 2010). Moreover, Cocksackie A virus (Rieder et al. 2001) and sylvatic dengue virus (Vasilakis et al. 2011) could potentially emerge to fill the niches left vacant should polio and human dengue virus, respectively, be eradicated. Evolutionary analysis will be needed to predict the impacts of eradication on future viral emergence.

Conclusion

In 1937, Max Theiler, the father of the yellow fever vaccine, wrote “One of the most striking phenomena to the student of virus diseases is the occurrence of variants.” (Theiler and Smith 1937). Of course evolutionary biologists are equally fascinated by variants, and it is my hope that this review has shown how evolutionary biology and vaccinology have been intertwined since the inception of vaccination and how they must remain linked for vaccine design and deployment to proceed safely and effectively. The information and references herein may be useful for design of lectures or introduction to case studies that demonstrate the contributions of evolutionary biology to biomedical advances, including the generation of most of the virus vaccines in use today and the creation of novel strategies for vaccine design. This review can also be used to emphasize the danger of ignoring evolution when deploying live-attenuated vaccine viruses, since without due safeguards, evolution may reshape the virulence or transmissibility of these agents. Moreover, virus vaccines themselves can and have influenced the evolution of naturally occurring viruses; the consequences of vaccination as a force of selection will continue to unfold as new vaccines are created and old vaccines become available to a larger proportion of the global population.

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