We compared cellular and humoral immunity to vaccinia virus (VV) in individuals exposed to 3 different orthopoxviruses: 154 individuals previously vaccinated with VV, 7 individuals with a history of monkeypox virus infection, and 8 individuals with a history of variola virus infection. Among individuals vaccinated 1-20 years prior, 9 (14%) of 66 individuals demonstrated VV-specific interferon (IFN)-γ enzyme-linked immunospot (ELISPOT) assay responses; 21 (50%) of 42 had lymphoproliferative (LP) responses, and 29 (97%) of 30 had VV-specific neutralizing antibodies. One year after monkeypox virus infection, 6 of 7 individuals had IFN-γ ELISPOT responses, all had VV-specific LP responses, and 3 of 7 had VV-specific neutralizing antibodies. Of 8 individuals with a history of variola virus infection, 1 had a VV-specific IFN-γ ELISPOT response, 4 had LP responses against whole VV, 7 had LP responses against heat-denatured vaccinia antigen, and 7 had VV-specific neutralizing antibodies. Survivors of variola virus infection demonstrated VV-specific CD4 memory cell responses and neutralizing antibodies >40 years after infection.

The last known case of variola virus infection in the United States occurred in 1949 in Texas [1]. The discontinuation of routine immunization against variola virus in the United States in 1972 and the worldwide eradication of natural infection with variola virus in 1979 provide a unique opportunity for the study of long-term immunological memory to a virus without circulating homologous antigen exposure. The potential use of remaining variola virus stocks as a bioterrorism agent has prompted investigators to measure immunological memory in previously vaccinated individuals. These prior studies revealed the presence of vaccinia virus (VV)–specific memory T and B cell responses [2–4], although the level of protection against natural variola virus infection that this may confer remains unclear, and comparison to individuals infected with variola virus has not been reported previously. Protective immunity conferred after surviving natural infection with variola virus is thought to be lifelong [5], although correlates of that protection have not been examined with contemporary immunological tools. Improved understanding of human immune responses to poxviruses, including variola virus, could have important implications for the development of safer VV-based vaccines. We describe here the human memory immune responses after vaccination with VV and also after natural infection with monkeypox or variola virus.

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METHODS

Study Population
We studied 4 cohorts of individuals from whom blood samples were collected between April 2002 and October...
2005. Volunteers were recruited through the Veterans Affairs New York Harbor Healthcare System (Institutional Review Board–approved protocol 00385) and the University of Massachusetts Medical School (UMMS; protocol H-10849 or H-3348).

**History of VV vaccination.** All 154 individuals had written documentation of the date of VV vaccination. For individuals who had had ≥2 vaccinations, the date of the last vaccination served as the reference date for the determination of time since last vaccination. Eighty-six individuals were vaccinated ≤2 years prior, 2 were vaccinated 3–20 years prior, and 66 were vaccinated >20 years prior.

**History of monkeypox virus infection.** A year after a monkeypox outbreak in Wisconsin in 2003 [6], 7 individuals with a history of monkeypox virus infection were enrolled. Five individuals with laboratory-confirmed monkeypox virus infection were identified through their physician. Two additional individuals were identified through these 5 individuals: one individual was the wife of a confirmed case patient who had had contact with an ill prairie dog and who had symptoms of monkeypox virus infection but did not have laboratory confirmation. The second individual was a veterinary-clinic employee who was exposed to an ill prairie dog during the outbreak period and who developed a typical pox rash and headache within 21 days after contact but did not have laboratory confirmation. Using a standardized questionnaire, we collected information on clinical symptoms, history of VV vaccination, exposure history, and results of diagnostic laboratory tests. Diagnosis of monkeypox was categorized as confirmed, probable, or suspect, as defined by the Centers for Disease Control and Prevention [7].

**History of variola virus infection.** We identified 8 individuals with a history of variola virus infection through advertisements in Indian-American newspapers. Volunteers were questioned in a face-to-face interview about the nature of the variola virus infection and their history of VV vaccination and were surveyed for the presence of facial pockmarks and VV-vaccination scars. The facial scar survey for evidence of prior variola virus infection was conducted as described elsewhere [8]. A probable case of variola virus infection was defined as having occurred in an individual with a history of variola illness before 1975 and the presence of ≥5 facial scars [8]. The main confounder in the facial scar survey is a history of severe chickenpox, which also can leave residual scars. However, a facial scar survey conducted for 12 months in Bangladesh, after the eradication of variola virus infection in 1975, indicated that chickenpox rarely results in ≥5 facial pockmarks [8]. Loss of scars over time has been reported almost exclusively when variola virus infection occurred at 6–12 months of age and does not appear to increase with time [8]. Thus, 1 volunteer who reported having had variola virus infection at <1 year of age (she and several members of her family had variola virus infection) was categorized as having a “probable” history of variola virus infection, since facial scarring was absent. Because 3 subjects had a history of both variola virus infection and VV vaccination and because these events occurred within 1–2 years of one another, we used the year of variola virus infection as the reference year.

**Poxvirus-naive individuals.** We enrolled 15 healthy individuals living in the United States, 18–33 years of age, who did not have a history of VV vaccination. These individuals also had the pertinent negative military and travel history, and pock lesions and vaccine scars were absent.

**Laboratory Assays**

Blood specimens obtained for serum and peripheral blood mononuclear cells (PBMCs) were harvested from sodium citrate cell-separator tubes (Becton Dickinson). Samples were processed within 24 h of blood donation. Separated PBMCs were counted and resuspended with 10% serum from blood-group AB donors in RPMI 1640 medium, at a concentration of 2 × 10^6 cells/mL, for assays using fresh or cryopreserved samples as described elsewhere [3]. The New York City Board of Health strain of VV, derived from the Dryvax vaccine, was used in all assays. Enzyme-linked immunospot (ELISPOT) and lymphoproliferation assays were conducted at New York University (NYU) and UMMS; cytotoxic T lymphocyte (CTL) and neutralizing-antibody assays were conducted at UMMS.

**Interferon (IFN)–γ ELISPOT assay.** We quantified VV-specific IFN-γ–producing T cells by using the IFN-γ ELISPOT assay on both fresh and cryopreserved PBMCs. The results of the ELISPOT assays did not differ significantly between the fresh and frozen specimens (data not shown). The assays were conducted as described by Kennedy et al. [3] and Borkowsky et al. [9], with the following modifications. VV was added at 2 μL/well at an MOI of 2 virions/cell. Phytohemagglutinin (PHA [Sigma]; 1:100 dilution and a final concentration of 10 μg/mL) at 10 μL/well served as a positive control well. Results represent the mean value of triplicate wells, expressed as IFN-γ spot-forming units per 10^6 PBMCs. On the basis of results from 15 poxvirus-naive individuals, a positive cutoff of >15 sfu/10^6 PBMCs was used.

**Lymphoproliferation assay.** The lymphoproliferation assay was a modification of the methods described by Valentine et al. [10]. Fresh PBMCs were isolated by the Ficoll-Hypaque method and then were washed and suspended in RPMI 1640 medium. Cells were counted and adjusted to a concentration of 10^6 cells/mL, and then 0.1 mL of the cell suspension was added to a 96-well (U-bottom) plate containing quadruplicate wells of a 1:5000 dilution of VV, cytomegalovirus, PHA, and a control medium prepared at twice the final concentration in RPMI 1640–20% heat-inactivated serum from blood-group AB.
donors, 2% penicillin, and 2% streptomycin. The VV dilutions were prepared from a stock solution of 10⁸ pfu/mL (the same stock that was used to prepare the VV dilutions for the ELISPOT assays). This virus preparation did not stimulate lymphocytes from unvaccinated healthy individuals.

In assays using vaccinia antigen (VacAg), the VacAg was prepared by heat inactivation and was tested for residual live virus by means of a plaque assay. A single lot of virus-free antigen was used at dilutions of 1:40 and 1:80, for comparison. Stimulation of 2 × 10⁵ cells for 5 days was done as described elsewhere for VV [11, 12].

The stimulation index (SI) was calculated by dividing the median counts per minute from 4 wells containing cells exposed to antigen by the median counts per minute for cells incubated with medium alone. A positive cutoff was defined as an SI of ≥3.

**VV-specific CTL assays.** Blood specimens were received within 16 h of donation, and separated PBMCs were cryopreserved within 24 h of blood donation. B lymphoblastoid cell lines were prepared from PBMCs from each donor by transformation with Epstein-Barr virus [3, 11]. Cryopreserved donor PBMCs were thawed, washed, and suspended in 5 mL of RPMI 1640 with 10% heat-inactivated fetal bovine serum and then were counted and prepared in accordance with well-defined protocols [3, 11]. In brief, CTL assays were performed by the infection of 0.2–1 million target cells (autologous BLCL) with VV 1 day before labeling for 60 min with 0.25-mCi ⁵¹Cr (New England Nuclear). Target cells were washed, counted, and resuspended to 15,000 cells/mL, and 0.1 mL of target cells was added per well in 96-well (U-bottom) plates. Effector cells were counted and washed and then were added to each well at effector-to-target cell (E:T) ratios of 90, 30, and 10, in triplicate. Target cells with 0.1 mL of medium served as minimum lysis controls. Plates were incubated at 37°C for 4.5 h. Well supernatants were harvested by use of the Skatron supernatant collection system and were counted in a Packard gamma counter. Target cells lysed with 0.1 mL of RENEX (detergent) served as maximum lysis controls [4]. At each E:T ratio, a percent-specific lysis of VV-infected target cells was calculated as the difference between the percent-specific lysis of VV-infected target cells and the percent-specific lysis of uninfected target cells. Lytic units (LU) per 10⁶ PBMCs were determined from the percent-specific lysis at each E:T ratio, by use of the exponential fit method based on software provided by Proteins International [13]. A normal LU value was between 1 and 1000, and the LU value represents a semiquantitative measure of the cell-mediated cytotoxicity observed in 10⁶ PBMCs. In prior clinical studies, a positive CTL response was defined as ≥5 LU, and 5 LU indicates an ~5-fold increase above the background lysis level [11].

**Neutralizing-antibody assay.** Humoral immune responses were assessed by serial plaque-reduction neutralization titer (PRNT) assays [12]. Samples obtained at blood-cell harvest were tested for the presence of VV-specific neutralizing antibodies, by means of 2-fold serial dilutions of heat-inactivated serum. VV at a concentration of 1 × 10⁶ pfu/mL was added to multiple dilutions of serum (1:10 through 1:3840) and to positive (New York City Board of Health strain) and negative control samples. The resulting mixtures were incubated for 1 h and then were plated on Vero 76 cells. Plaques were stained with neutral red and counted at 72 h, and the reduction in plaques was plotted against the dilution factor. Antibody titers (the reciprocal of the dilution) resulting in plaque neutralization of 50% (i.e., PRNT₅₀) were calculated from the plot. A positive cutoff was defined as a titer ≥1:20.

**RESULTS**

**Immune responses after VV vaccination.** VV-specific IFN-γ ELISPOT assays were positive for 64 (42%) of the 154 vaccinees, and lymphoproliferative (LP) responses were positive for 56 (67%) of 83 vaccinees tested (figure 1). Of the 83 vaccinees tested by both the ELISPOT and lymphoproliferation assays, 30 (36%) had both VV-specific IFN-γ–producing lymphocytes and proliferative memory responses; 26 (31%) demonstrated proliferative memory responses without the detection of IFN-γ–producing cells; and the remaining 27 (33%) demonstrated neither IFN-γ–producing lymphocytes nor proliferative responses.

CTL responses were detected in 27 (41%) of 66 vaccinees tested. The median number of years since last vaccination was 1 for those with a CTL response, compared with a median of 35 years (range, 1–65 years) for those without a CTL response. All 27 vaccinees with CTL responses also had VV-specific proliferative responses, and 21 also had VV-specific IFN-γ–producing lymphocytes.

The proportion of VV vaccinees with a detectable VV-specific memory T cell immune response decreased over time (table 1). At >20 years after vaccination, 9 (14%) of 66 vaccinees tested had detectable IFN-γ–producing lymphocytes, and 21 (50%) of 42 vaccinees had VV-specific LP responses. CTL responses were not detected in any subjects, but 29 (97%) of 30 vaccinees had a neutralizing-antibody titer ≥1:20. VV-specific IFN-γ–producing lymphocytes were detected in 1 individual 46 years after this person’s last VV vaccination, and LP responses to virus were detected in another individual 56 years after vaccination.

Overall, 60 (91%) of 66 vaccinees had detectable neutralizing-antibody levels (titer ≥1:20); strong neutralizing-antibody responses (titer ≥1:1280) were detected in 3 individuals ≥35 years after their last VV vaccination.

**Immune responses after monkeypox virus infection.** Seven individuals with suspected or confirmed monkeypox virus infection were identified in Wisconsin (4 women and 3 men;
Figure 1. Distribution of laboratory measures of cellular and humoral immunity in individuals with a history of vaccinia virus vaccination, variola virus infection, or monkeypox virus infection. IFN, interferon; PBMCs, peripheral blood mononuclear cells.
Table 1. Cell-mediated and humoral immune responses among individuals vaccinated with vaccinia virus (VV), by type of immunological test and interval since last vaccination.

<table>
<thead>
<tr>
<th>Years since last VV vaccination</th>
<th>IFN-γ ELISPOT assay</th>
<th>Lymphoproliferation assay</th>
<th>Cytotoxic T cell lysis</th>
<th>Neutralizing antibody</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;2</td>
<td>54/86 (63)</td>
<td>34/40 (85)</td>
<td>27/36 (75)</td>
<td>31/36 (86)</td>
</tr>
<tr>
<td>3–20</td>
<td>1/2 (50)</td>
<td>1/1 (100)</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>&gt;20</td>
<td>9/66 (14)</td>
<td>21/42 (60)</td>
<td>0/30 (0)</td>
<td>29/30 (97)</td>
</tr>
</tbody>
</table>

**NOTE.** Data are no. of positive individuals/total no. tested (%). IFN, interferon; ELISPOT, enzyme-linked immunospot; ND, not determined.

Immune responses after variola virus infection. We identified 8 individuals with a history of variola virus infection before 1975 (table 3). The median age was 63 years (range, 50–78 years), and 5 (63%) were men. The variola virus infections occurred in India (7 individuals) and Bangladesh (1 individual) between 1928 and 1961. The median number of years since variola virus infection was 57 (range, 44–77 years), and the median age at the time of variola virus infection was 6 years (range, <1–16 years). Three individuals reported having received a VV vaccination and had a vaccination scar; 5 individuals denied a history of VV vaccination and did not have a vaccination scar. Six individuals reported that other household members also had variola virus infection.

One year after monkeypox virus infection, all 7 subjects had VV-specific LP responses; 6 subjects had detectable IFN-γ ELISPOT responses, and 2 subjects had CTL responses (figure 1). Three (43%) of 7 subjects had a VV-specific neutralizing-antibody titer ≥1:20 (table 2). The 2 individuals who had a positive response for all 3 cellular immune-response assays did not have detectable serum neutralizing antibody to VV. The absence of neutralizing-antibody titers was confirmed with a second assay, described by Kennedy et al. [3].

Table 2. Characteristics and assay results for 7 individuals with a history of monkeypox virus infection and with cell-mediated and humoral immunity 1 year after infection, United States, 2004–2005.

<table>
<thead>
<tr>
<th>Individual</th>
<th>VV vaccination</th>
<th>Clinical symptom⁵</th>
<th>III days, no.</th>
<th>Assay result</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Typical lesions, no.</td>
<td>Headache</td>
<td>Fever</td>
<td>Chills</td>
</tr>
<tr>
<td>1</td>
<td>Yes (&gt;43 years ago)</td>
<td>7</td>
<td>−</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>No</td>
<td>24</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>3</td>
<td>No</td>
<td>20</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>4</td>
<td>No</td>
<td>1</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>5</td>
<td>No</td>
<td>&gt;100</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>6</td>
<td>Yes (year unknown)</td>
<td>12</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>7</td>
<td>No</td>
<td>50</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

**NOTE.** CTL, cytolytic T lymphocyte; ELISPOT, enzyme-linked immunospot; LAD, lymphadenopathy; LU, lytic unit; nAb, neutralizing antibody; PBMCs, peripheral blood mononuclear cells; SI, stimulation index; VV, vaccinia virus.

⁵ A plus sign (+) indicates presence of a symptom, and a minus sign (−) indicates absence of a symptom.

⁶ Value exceeds positive cutoff level for assay.
Figure 2. Facial scar survey of individuals with a history of variola virus infection. A, 50-year-old Bangladeshi man who had variola virus infection 44 years ago and 2 vaccinia virus (VV) vaccinations. Assay results were as follows: interferon (IFN)-γ enzyme-linked immunospot (ELISPOT), 2.7 sfu/10⁶ peripheral blood mononuclear cells (PBMCs); lymphoproliferation, stimulation index (SI) of 38; cytotoxic T lymphocyte (CTL), 2.2 lytic units (LU); and VV-specific neutralizing-antibody titer, 1:80. B, 62-year-old Indian man who had variola virus infection 60 years ago and no history of VV vaccination. IFN-γ ELISPOT, 0 sfu/10⁶ PBMCs; lymphoproliferation, SI of 5; CTL, not available; and VV-specific neutralizing-antibody titer, 1:80.

DISCUSSION

By using contemporary assays, we provide the first description of residual cell-mediated and humoral immunity in individuals who survived variola virus infection, compared with individuals with a history of monkeypox virus infection or VV vaccination. Individuals who survived an episode of variola virus infection were thought to have lifelong protection against reinfection [5], although a reinfection rate of 1 in 1000 cases has been reported in India, with an average interval of ~15–20 years between attacks [14, 15]. We found high titers of neutralizing antibody against VV in 4 of 5 unvaccinated survivors of variola virus infection who had had the infection >40 years ago, suggesting that the antibody response after natural variola virus infection is long lasting. In contrast, none of the 5 unvaccinated individuals with a history of variola virus infection had IFN-γ production detected by ELISPOT assay, and only 1 had an LP response to VV. Stimulation with heat-denatured VacAg led to a more robust expansion of VV-specific CD4 cells; such cells were undetectable when whole VV was used. This discrepancy may reflect differences in concentration and types of epitopes present in VacAg and whole VV preparations; alternatively, it may reflect differences in processing and presentation of antigen in cultured cells.

One year after monkeypox virus infection, VV-specific IFN-γ–producing T cells were present at a level comparable to levels in those who had received a VV vaccination 1–2 years prior, which is within the time frame of maximal vaccine-induced protection against variola virus. T cell LP responses to VV also were robust in these survivors of monkeypox virus infection. The presence of VV-specific IFN-γ–producing T cells and LP responses in individuals with monkeypox virus infection who had never received VV vaccination reflects cross-protective immunity between VV and monkeypox virus [16]. Only 3 of the 7 survivors of monkeypox virus infection demonstrated VV-specific neutralizing-antibody titers ≥1:20, including 2 without a history of VV vaccination. This finding may be explained by the serotype-specific nature of neutralizing-antibody responses, compared with the more cross-reactive T cell responses to epitopes in orthopoxviruses [17, 18]. In contrast to our findings, other studies have found higher antibody responses in patients with previous monkeypox virus infection, by use of ELISA [16, 19].

Characterized as IFN-γ–producing CD4 cells (CD8 cells were not detected) and was not detected in 6 VV vaccinees, who were vaccinated a median of 40 years prior, or in 3 pox-naive individuals (data not shown). These results suggest that the T lymphocytes proliferating in response to VacAg were long-lasting CD4 memory cells and that survivors of variola virus infection maintained a low but extant population of VV-specific CD4 memory cells, despite the lack of ongoing antigen exposure to variola virus or VV.
Years since variola virus infection & Facial scars, no. & VV vaccination & ELISPOT, sfu/10^6 PBMCs & CTL, LU & Lymphoproliferation, SI & Heat-inactivated VacAg & nAb titer

<table>
<thead>
<tr>
<th>Individual (age in years, sex)</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (62, male)</td>
<td>60</td>
<td>~50</td>
<td>–</td>
<td>0</td>
<td>NA</td>
<td>5^b</td>
<td>6^b</td>
<td>1:80^b</td>
</tr>
<tr>
<td>2 (55, male)</td>
<td>51</td>
<td>25</td>
<td>–</td>
<td>1.3</td>
<td>3.2</td>
<td>2.4</td>
<td>2</td>
<td>1:10</td>
</tr>
<tr>
<td>3 (69, male)</td>
<td>63</td>
<td>25</td>
<td>–</td>
<td>1.3</td>
<td>0</td>
<td>1.3</td>
<td>16^b</td>
<td>1:40^b</td>
</tr>
<tr>
<td>4 (71, female)</td>
<td>64</td>
<td>20</td>
<td>–</td>
<td>2.7</td>
<td>0</td>
<td>2</td>
<td>24^b</td>
<td>1:1280^b</td>
</tr>
<tr>
<td>5 (64, male)</td>
<td>61</td>
<td>~100</td>
<td>–</td>
<td>1.3</td>
<td>.4</td>
<td>1</td>
<td>29^b</td>
<td>1:160^b</td>
</tr>
<tr>
<td>6 (78, female)</td>
<td>77</td>
<td>0</td>
<td>+</td>
<td>0</td>
<td>.7</td>
<td>153^b</td>
<td>131^b</td>
<td>1:80^b</td>
</tr>
<tr>
<td>7 (50, male)</td>
<td>44</td>
<td>~50</td>
<td>+</td>
<td>2.7</td>
<td>2.2</td>
<td>38^b</td>
<td>86^b</td>
<td>1:80^b</td>
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<tr>
<td>8 (61, female)</td>
<td>45</td>
<td>~50</td>
<td>+</td>
<td>25.3^b</td>
<td>.9</td>
<td>20^b</td>
<td>8^b</td>
<td>1:320^b</td>
</tr>
</tbody>
</table>

NOTE.  CTL, cytotoxic T lymphocyte; ELISPOT, enzyme-linked immunospot; LU, lytic unit; NA, not available; nAb, neutralizing antibody; PBMC, peripheral blood mononuclear cell; SI, stimulation index; VacAg, vaccinia antigen; VV, vaccinia virus.

^a A plus sign (+) indicates presence of a symptom, and a minus sign (−) indicates absence of a symptom.

^b Value exceeds positive cutoff level for assay.

Future studies could address the issue of lifelong immunity.
induced by variola virus infection by challenging survivors of variola virus infection with Dryvax vaccine or with experimental vaccines, to assess in vitro immune responses. These studies could help delineate the components of postchallenge antibody and T cell responses that are specific for both variola virus and VV protein epitopes. Such a study of survivors of variola virus infection would require comparison to vaccinated individuals and would significantly improve our understanding of the role of prior infection in protecting against or attenuating the response to subsequent exposure to a homologous virus.

Acknowledgments

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