Preexisting Immunity to Pandemic (H1N1) 2009

To the Editor: The influenza A pandemic (H1N1) 2009 virus contains a combination of 8 gene segments (1–3) antigenically similar to North American influenza A virus (H1N1) but different from seasonal human influenza A viruses (H1N1) (3). Despite the initial high number of deaths among patients in Mexico and among patients with specific preexisting conditions, pandemic (H1N1) 2009 virus in general has caused mild symptoms, and the overall death rate remains around 0.45% (www.who.int/csr/don/2009_07_06/en). Low virulence of the virus and preexisting immune status are among the main factors that account for lower death rates in influenza outbreaks. The Centers for Disease Control and Prevention (Atlanta, GA, USA) reported that among persons >60 years old, 33% have preexisting, cross-reactive neutralizing antibodies against the new virus, but seasonal influenza vaccines do not elicit cross-reactive neutralizing antibodies against pandemic (H1N1) 2009 virus in either younger or older populations (1). However, current data cannot be used to evaluate the full immune capacities of human populations because cell-mediated immunity (CMI) has not been characterized in humans infected with pandemic (H1N1) 2009 virus.

We performed a survey (4) for known human immune epitopes present in the various proteins of seasonal influenza A virus strains and known to be efficient in stimulating lymphocytes. We found that multiple major histocompatibility complex (MHC)–restricted epitopes are conserved in nucleoprotein (NP) and matrix protein (MP), and even a few in the more variable hemagglutinin (HA) protein, in A/California/04/2009, A/Texas/04/2009, and A/New York/18/2009.

For MHC class II antigen-restricted epitopes essential for antibody and Th1 responses, HA of pandemic (H1N1) 2009 virus contains HLA-DRA*0101/DRB1*0101-restricted SVIEKMNTQF-TAV (5), as well as HLA-DRA*0101/DRB1*0401-restricted EKMTQF-TAVGKE, TGLRNIPISOQR, and ELLVLLLENERTLDY (5), and HLA-DRB5*0101-restricted DYELREQLSSVSSFERFE (5) epitopes. These antigen-restricted epitopes were present in globally-distributed seasonal H1N1 viruses, including classical A/New Caledonia/20/1999 (H1N1) and A/Solomon Islands/3/2006 (H1N1). Overall, high levels of cross-reactive microneutralization (MN) or hemagglutination inhibition (HI) antibodies may not be detected against pandemic (H1N1) 2009 virus. This lack of detection of MN or HI antibodies is probably because most of these epitopes may not elicit MN/HI detectable antibodies, or these epitopes may be present in earlier seasonal influenza strains but not present in the current trivalent, inactivated influenza vaccine. Even though many do not contribute to neutralizing antibodies, these MHC class II antigen-restricted epitopes may initiate the Th1 response, including activation of infected macrophages and antiviral cytokine production, and help host defenses as well.

For MHC class I antigen-restricted epitopes essential for CD8+ T cell activation and CMI, HA of pandemic (H1N1) 2009 virus contains HLA-A*0201-restricted GLFGAIAGFI (6), which is present also in the HA of A/New Caledonia/20/1999 (H1N1) and A/Solomon Islands/3/2006 (H1N1) viruses. More MHC class I antigen-restricted epitopes in NP and MP of seasonal epidemic influenza viruses (H1N1) and (H3N2) are conserved in pandemic (H1N1) virus. These seasonal influenza viruses were isolated in North America, Europe, Africa, and Asia–Pacific regions. Conserved epitopes in the HA and NP of pandemic (H1N1) 2009 virus are listed in the Table. In addition, ≥15 completely conserved epitopes are in the M protein of pandemic (H1N1) 2009 virus (data not shown).

Studies have demonstrated that both humoral and cell-mediated immune responses may contribute to protection in influenza-vaccinated persons. As for humoral immunity, results have consistently indicated that serum HI or MN antibody titers correlate inversely with morbidity rates after vaccination, which are the most valuable correlates of protection (7). Studies supporting the role of CMI in influenza viral clearance and host survival are well-documented in mouse models, but data are limited for humans. However, emerging evidence has demonstrated that either infection or vaccination can induce T cell-mediated immune responses in humans (8,9). Moreover, higher levels of CD8+ T cells correlate with reduced viral shedding among experimentally infected humans (10). Notably, among vaccinated persons ≥60 years of age, measures of the ex vivo cellular immune response are statistically correlated with protection against influenza illness but serum HI antibody levels are not, suggesting a role for CMI (9). Therefore, it is rational to expect that CMI does provide a protective role, and cross-reactive CMI to pandemic (H1N1) 2009 virus through conserved MHC class I-restricted epitopes may exist in persons previously vaccinated for or exposed to seasonal influenza.

We note that ≥80% of MHC class I epitopes in NP of seasonal and flu vaccine viruses (Table) are also completely conserved in the highly pathogenic avian H5N1 virus (A/Hong Kong/156/1997 and A/Hong Kong/97/1998) (www.ncbi.nlm.nih.gov/genomes/FLU). Several points have to be made regarding the relevance of these epitopes to its high associated mortality rate. First, influenza virus (H5N1) is known to be highly virulent, replicating at a much faster
pace than other influenza A viruses and spreading in vital organs shortly after infection and before epitope-mediated protective immunity can be launched, which may account for its high fatality rate. Second, the epitopes are MHC class I antigen-restricted, which means that only a fraction of the human population will possess the correct MHC class I molecules capable of presenting a specific epitope and eliciting appropriate and protective CMI responses. This lack of correct MHC class I molecules could explain why patients of varied genetic backgrounds may have different prognoses upon infection with pandemic (H1N1) 2009 virus or even influenza virus (H5N1).

In fact, although there are no experiments establishing a solid link, cross-reactive immunity from seasonal influenza virus or vaccination may result in partial cell-mediated or humoral immunity to influenza virus (H5N1). The same type of immunity may have happened in persons exposed to pandemic (H1N1) 2009 virus as well.

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Table. Conserved MHC class I antigen-restricted epitopes present in HA and NP proteins of pandemic (H1N1) 2009 virus*

| MHC antigen | Epitope position | Sequence | Identified in selected virus isolates |
|-------------|------------------|----------|--------------------------------------|
| HA          |                  |          |                                      |
| HLA-A*0201  | 344–353          | GLFGAIAGFI | A/New Caledonia/20/1999 (H1N1) A/Solomon Islands/3/2006 (H1N1) A/Macau/229/2008 (H1N1) A/Managua/254.01/2008/H1N1 |
| NP          |                  |          |                                      |
| HLA-A1      | 44–52            | CTELKLSDY | A/Hong Kong/HKU4/2004 (H3N2) A/Canterbury/200/2004 (H3N2) |
| HLA-A*0101  |                  |          |                                      |
| HLA-A3      | 265–273          | ILRGVSVAHK | A/New Caledonia/20/1999 (H1N1) A/New York/55/2004 (H3N2) A/Managua/254.01/2008/H1N1 A/Taiwan/2645/2006 (H1N1) |
| HLA-B27     | 380–393          | ELRSRYWAI TRSG | A/New Caledonia/20/1999 (H1N1) A/Taiwan/2645/2006 (H1N1) A/Managua/254.01/2008/H1N1 A/Florida/UR06-0383/2007/H1N1 |
| HLA-B27     | 174–184          | RRSGAAGAAVK | A/New Caledonia/20/1999 (H1N1) A/New York/55/2004 (H3N2) A/California/UR07-0067/2008 (H3N2) |
| HLA-B*2705  | 357–370          | KLSTRGVQIASNEN | A/New Caledonia/20/1999 (H1N1) A/New York/55/2004 (H3N2) A/Hong Kong/HKU77/2005 (H3N2) |
| HLA-B*2705  | 383–391          | SRYWAI TRTR | A/New Caledonia/20/1999 A/Taiwan/2645/2006 (H1N1) A/Managua/4537.03/2008/H1N1 A/Florida/UR07-0026/2008/H1N1 |
| HLA-B*2702  | 381–388          | LRSRYWAI | A/New Caledonia/20/1999 A/Florida/UR06-0383/2007/H1N1 A/Canterbury/200/2004 (H3N2) A/Hong Kong/HKU77/2005 (H3N2) |
| HLA-B*4002  | 251–259          | AEIEDLIFL | A/Canterbury/200/2004 (H3N2) A/Hong Kong/HKU77/2005 (H3N2) A/Florida/UR06-0383/2007/H1N1 |
| HLA-B8      | 225–233          | ILKGKFQTA | A/New Caledonia/20/1999 A/California/UR07-0067/2008 (H3N2) |
| HLA-B8      | 380–388          | ELRSRYWAI | A/New Caledonia/20/1999 (H1N1) A/Taiwan/2645/2006 (H1N1) A/Managua/107.01/2008/H1N1 A/Florida/UR07-0026/2008/H1N1 |

*HA, hemagglutinin; NP, nucleoprotein; MHC, major histocompatibility complex; NA, neuraminidase. Epitope binding to and/or activation of specific lymphocytes prepared from human peripheral blood mononuclear cells (PBMC): MHC-tetramer staining; T-cell receptor binding; ELISPOT and intracellular cytokine staining (interferon-γ), and/or 51Chromium release and killing have been demonstrated in published studies. Data on epitope characterization were collected from the Immune Epitope Database (IEDB; www.immuneepitope.org) (4).
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The current outbreak presents the first opportunity to directly observe this process. We used hemagglutination inhibition (HI) and virus neutralization (VN) assays to detect antibodies in 4,043 serum samples from residents (7–84 years of age) of 2 counties in Guangxi Province, People’s Republic of China, collected during July–August 2008. These persons were mostly farmers who lived in rural areas. Serum samples were obtained, transported, and frozen at –80°C as needed human infection with H1 swine influenza virus (H1N1). The current study was designed to directly observe this process.

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