The pandemic influenza virus of 1918–1919 killed an estimated 20 to 50 million people worldwide. With the recent availability of the complete 1918 influenza virus coding sequence, we used reverse genetics to generate an influenza virus bearing all eight gene segments of the pandemic virus to study the properties associated with its extraordinary virulence. In stark contrast to contemporary human influenza H1N1 viruses, the 1918 pandemic virus had the ability to replicate in the absence of trypsin, caused death in mice and embryonated chicken eggs, and displayed a high-growth phenotype in human bronchial epithelial cells. Moreover, the coordinated expression of the 1918 virus genes most certainly confers the unique high-virulence phenotype observed with this pandemic virus.

The influenza pandemic of 1918 was exceptional, resulting in the deaths of up to 50 million people worldwide, including an estimated 675,000 deaths in the United States (1, 2). The pandemic’s most striking feature was the unusually high death rate among healthy adults aged 15 to 34 years, which consequently lowered the average life expectancy in the United States by more than 10 years (3). A similarly high death rate has not occurred in this age group in either prior or subsequent influenza A pandemics or epidemics (4).

Genomic RNA of the 1918 virus was recovered from archived formalin-fixed lung autopsy samples from an Alaskan influenza victim who was embalmed in the absence of trypsin, and from frozen, unfixed lung tissues obtained from archived formalin-fixed lung autopsy samples. The complete coding sequences of all eight viral RNA segments have now been determined, providing insights into the nature and origin of this pathogen (5–11). Plasmid-based reverse genetics has allowed for the generation of recombinant viruses containing 1918 hemagglutinin (HA) with or without the 1918 neuraminidase (NA) rescued in the genetic background of contemporary human H1N1 or H3N2 influenza viruses. The resulting strains were demonstrated to cause mortality in mice only at high infection doses (12, 13); however, the virulence of the complete 1918 virus has not been evaluated.

In the present study, we generated a virus containing the complete coding sequences of the eight viral gene segments from the 1918 virus in an effort to understand the molecular basis of virulence of this pandemic virus. Genes encoding the 1918 influenza virus were reconstructed from deoxycytidinonucleotides and corresponded to the reported coding sequences of the 1918 virus as previously described (5–11). Because the 1918 S’ and 3’ noncoding regions have not been sequenced, the genes were constructed such that they had the noncoding regions corresponding to the closely related influenza A/WSN/33 (H1N1) virus. The 1918 virus and recombinant H1N1 influenza viruses were generated using the previously described reverse genetics system (8, 14). All viruses containing one or more gene segments from the 1918 influenza virus were generated and handled under high-containment [biosafety level 3 enhanced (BSL3)] laboratory conditions in accordance with guidelines of the National Institutes of Health and the Centers for Disease Control and Prevention (15). Viruses were grown in Madin-Darby canine kidney cells (MDCK) and/or the allantoic cavity of 10-day-old embryonated hen’s eggs (table S1). The control viruses included an avian A/duck/Alberta/35/76 H1N1 virus, two contemporary human H1N1 influenza viruses, the wild-type A/New Caledonia/20/99 (N. Cal/99, H1N1) virus and A/Texas/36/91 (Tx/91, H1N1) virus generated by reverse genetics. The other recombinant viruses used were a virus having only the HA from the Tx/91 virus with the remaining seven genes from the 1918 virus (Tx/91 HA:1918); a virus having the NA from 1918 with the remaining seven genes from the Tx/91 virus (1918 NA:Tx/91); and recombinant viruses having two 1918 (1918 HA/NA:Tx/91) or five 1918 genes (1918 HA/NA/M/NP/NS:Tx/91) with the remaining genes derived from the Tx/91 virus. The HA of the 1918 viruses used throughout these studies was derived from A/South Carolina/1/18 strain that was shown to preferentially bind the α2,6 sialic acid (human) cellular receptor (16). The identity of the 1918 and Tx/91 influenza virus genes in the rescued viruses was confirmed by reverse transcription polymerase chain reaction and sequence analysis.

The infectivity of the 1918 virus and the ability to form plaques in the presence and in the absence of the protease trypsin were assayed in MDCK cells by the plaque method. The proteolytic cleavage of the HA molecule is a prerequisite for multicycle replication, and the ability of an influenza virus to replicate in the absence of trypsin has been thought to be an important determinant of influenza virus pathogenicity in mammals (17–20). In contrast to the contemporary human Tx/91 and N. Cal/99 H1N1 viruses, which require an exogenous protease source for their multicycle replication and plaque formation (Table 1), the 1918 virus and a recombinant influenza virus bearing the 1918 HA and NA segments only (1918 HA/NA:Tx/91) formed visible plaques without the addition of trypsin (Table 1). Furthermore, a virus having only the 1918 NA with the remaining genes, including the HA, from Tx/91 virus (1918 NA:Tx/91) also replicated in the absence of trypsin, which suggests that the 1918 NA activity facilitates HA cleavage. However, the 1918 HA and NA gene sequences lack the obvious genetic features that have previously been associated with the ability

<table>
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<tr>
<th>Virus</th>
<th>Infectivity titer (PFU/ml)</th>
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<tr>
<td></td>
<td>Trypsin (+)</td>
</tr>
<tr>
<td>N. Cal/99</td>
<td>3.6 x 10^7</td>
</tr>
<tr>
<td>Tx/91</td>
<td>1.4 x 10^7</td>
</tr>
<tr>
<td>1918 HA/NA/ Tx/91</td>
<td>2.4 x 10^7</td>
</tr>
<tr>
<td>1918 NA/ Tx/91</td>
<td>3.4 x 10^7</td>
</tr>
<tr>
<td>1918 (1)</td>
<td>4.8 x 10^7</td>
</tr>
<tr>
<td>1918 (2)</td>
<td>1.4 x 10^8</td>
</tr>
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*No visible plaques were formed at the lowest dilution (1:10) tested.
to replicate in the absence of trypsin; that is, the 1918 virus has neither a series of basic amino acids at the HA cleavage site (as seen in highly pathogenic avian H5 or H7 influenza viruses) nor mutations (N146R or N146Y) in the NA that lead to the loss of a glycosylation site like those that allow the A/WSN/33 virus NA to sequester plasminogen (6, 7, 17, 18, 20). We will need to consider other mechanisms of NA-mediated cleavage that may be relevant to the replication and virulence of the 1918 virus.

To evaluate the pathogenicity of the 1918 virus in a mammalian species, we intranasally inoculated BALB/c mice with two independently generated 1918 viruses and then determined morbidity (measured by weight loss), virus replication, and 50% lethal dose (LD$_{50}$) titers (21, 22). For comparison, groups of mice were infected with a recombinant influenza virus containing five 1918 genes with the remaining three polymerase genes from Tx/91 (1918 HA/NA/M/NP/NS/Tx/91) virus or a recombinant 1918 virus (Tx/91 HA:1918), in which the HA has been replaced with that of Tx/91 virus. Infection of mice with the 1918 virus resulted in lung virus titers, on day 4 post inoculation (p.i.), that were at least 125 and 39,000 times those of mice infected with the Tx/91 HA:1918 and Tx/91 viruses, respectively (Fig. 1A). The 1918 HA/NA/M/NP/NS/Tx/91 virus also replicated efficiently in the mouse lung, but virus titers were markedly lower than those for mice infected with the 1918 virus, and the lethality of the 1918 virus (LD$_{50}$ = $10^{3.85}$) was at least 100 times that for the 1918 HA/NA/M/NP/NS/Tx/91 virus (LD$_{50}$ = $10^{5.75}$). Strikingly, the mice infected with 10$^6$ PFU (plaque-forming units) of the 1918 virus succumbed to infection as early as 3 days p.i. (Fig. 1B), and they lost up to 13% of their body weight 2 days after infection (Fig. 1C). In contrast to the lethal outcome of the 1918 virus infection, the Tx/91 HA:1918 virus, like the wild-type Tx/91 virus, did not kill mice (LD$_{50}$ = $10^{6.5}$) (Fig. 1B) and displayed only transient weight reduction (Fig. 1C). To determine whether the 1918 virus replicated systemically in the mouse after intranasal infection with 10$^6$ PFU of virus, we harvested brain, heart, liver, and spleen tissues from four mice each on days 4 and 5 p.i. All eight mice infected with the 1918 virus or the Tx/91 control viruses had undetectable levels ($\leq 10^{3.8}$ of the 50% egg infectious dose (EID$_{50}$) determined by serial titration in chicken eggs) of virus in these tissues (fig. S1), which indicated that the 1918 virus did not spread systemically to other organs in the mouse.

Histopathological analysis of lung tissues from individuals who died from primary influenza pneumonia in 1918 frequently showed acute pulmonary edema and/or hemorrhage with acute bronchiolitis, alveolitis, and bronchopneumonia (23). In mice, the most severe lung lesions were observed after infection with the 1918 virus (21). On day 4 p.i., mice infected with the pandemic strain had necrotizing bronchiitis and bronchiolitis and moderate to severe alveolitis (Fig. 2A). The alveolitis varied from peribronchial to diffuse in distribution and was composed of neutrophils and macrophages. Accompanying the inflammation was moderate-to-severe peribronchial and alveolar edema (Fig. 2B) and alveolar hemorrhage. Neutrophils were the predominant inflammatory cells, but alveolar macrophages were also prominent (Fig. 2C). As in autopsy studies performed in 1918, there was no histological evidence of systemic infection such as necrosis or inflammation in the liver, kidney, spleen, heart, and brain tissues of mice infected with the 1918 virus. In general, the 1918 HA/NA/M/NP/NS/Tx/91 virus induced less severe pathology; however, mild-to-moderate peribronchial alveolitis with some mild-to-moderate alveolar edema was observed (Fig. 2D). The 1918 HA gene was essential for severe pulmonary lesion development, because the Tx/91 HA:1918 virus produced very mild pulmonary lesions characterized by minimal, diffuse alveolitis and mild focal lymphocytic-histiocytic bronchitis (Fig. 2E). The Tx/91-inoculated mice lacked noteworthy lesions in the lungs (Fig. 2E).

Table 2. Lethality of the 1918 influenza virus for 10-day-old embryonated chicken eggs. Fifty percent egg infectious dose (EID$_{50}$) and egg lethal dose (ELD$_{50}$) titers were determined as described in the text. For ELD$_{50}$ titers, embryo viability was visually determined by daily candling. EID$_{50}$ and ELD$_{50}$ titers were determined simultaneously and calculated by the method of Reed and Muench (27). The mean death time (MDT) of embryo death was calculated by examining embryo viability daily for 7 days. The MDT is the mean time in days for the minimum lethal dose to kill embryos.

<table>
<thead>
<tr>
<th>Virus</th>
<th>log$<em>{10}$ EID$</em>{50}$/ml</th>
<th>log$<em>{10}$ ELD$</em>{50}$/ml</th>
<th>MDT (day)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N. Cal/99</td>
<td>8.5</td>
<td>1.2</td>
<td>-</td>
</tr>
<tr>
<td>Tx/91</td>
<td>8.7</td>
<td>1.2</td>
<td>-</td>
</tr>
<tr>
<td>Tx/91 HA:1918</td>
<td>9.0</td>
<td>1.2</td>
<td>-</td>
</tr>
<tr>
<td>1918 HA/NA/M/NP/NS/Tx/91</td>
<td>8.7</td>
<td>1.2</td>
<td>-</td>
</tr>
<tr>
<td>1918 (1)</td>
<td>9.0</td>
<td>7.2</td>
<td>4.5</td>
</tr>
<tr>
<td>1918 (2)</td>
<td>9.5</td>
<td>8.2</td>
<td>4.5</td>
</tr>
<tr>
<td>A/duck/Alberta/35/76</td>
<td>9.0</td>
<td>8.5</td>
<td>2</td>
</tr>
</tbody>
</table>

Fig. 1. Pathogenicity of H1N1 viruses. Comparison of lung virus titers (A), lethality (B) and weight loss (C), in BALB/c mice (13 per virus group) intranasally infected with 10$^6$ PFU of Tx/91 (△), Tx/91 HA:1918 (▵), 1918 HA/NA/M/NP/NS/Tx/91 (●), 1918 (1) (○), or 1918 (clone 2) (▲) virus. Four mice from each virus-infected group were killed on day 4 p.i., and titers of individual lungs were determined in eggs and expressed as the mean ± SD. The limit of virus detection was 10$^{1.2}$ EID$_{50}$/ml. The remaining nine mice from each group were observed for weight loss and mortality through a 14-day observation period. *The 1918 virus lung titers are significantly (P < 0.05) different from those of all other virus infection groups as determined by analysis of variance.
other mammalian H1N1 strains (5–11), it was of interest to determine whether the 1918 strain would be lethal for fertile chicken eggs, a pathogenic feature of avian H1N1 viruses. After serial titration in 10-day-old embryonated chicken eggs, the 1918 virus was lethal for chicken embryos: 50% egg lethal dose (ELD$_{50}$) titers were similar to those for an avian H1N1 representative, A/duck/Alberta/35/76 virus (Table 2). By contrast, neither contemporary human H1N1 viruses nor any of the 1918 recombinant viruses containing two, five, or seven genes from the 1918 virus caused mortality of embryos by day 7 p.i., which indicated that the 1918 HA and 1918 polymerase genes were associated with virulence of chicken embryos.

We evaluated growth and release of the virus in polarized Calu-3 cells, a human lung epithelial cell line, grown on membrane inserts (24). Culture medium was collected from the apical and basolateral chambers at different times after inoculation and examined for virus production in the presence or absence of trypsin. With all viruses tested, titers of progeny virus progressively increased during the first 24 hours p.i., and virus was detected almost exclusively in the apical supernatant (Fig. 3). Regardless of the presence or absence of trypsin, infectivity titers of the 1918 virus were significantly higher than virus titers released in the Tx/91 HA:1918 and 1918 HA/NA/M/NP/NS:Tx/91 virus-infected cultures at 12, 16 and 24 hours p.i., which suggests that the 1918 HA and 1918 polymerase genes are essential for maximal replication of the virus in human bronchial epithelial cells. At 16 and 24 hours p.i., 1918 virus release was at least 50 times that observed in Tx/91 virus-infected cultures. The evidence of significantly higher apical release of 1918 progeny virus after an apical infection supports the hypothesis of an increase in the amount of virus present in the infected lung and may provide important insight into the virulence of this virus.

Until now, the exceptional virulence of the 1918 pandemic influenza virus has been a question of historical curiosity. Herein, we demonstrate the successful reconstruction of the 1918 pandemic virus in order to understand more fully the virulence of this virus and possibly of other human influenza pandemic viruses. Because the emergence of another pandemic virus is considered likely, if not inevitable (25), characterization of the 1918 virus may enable us to recognize the potential threat posed by new influenza virus strains, and it will shed light on the prophylactic and therapeutic countermeasures that will be needed to control pandemic viruses. A number of biological properties associated with this unusually virulent influenza virus were found. Comparison of the 1918 virus with recombinant viruses expressing one or more 1918 virus genes was essential for optimal virulence.
and that the constellation of all eight genes together make an exceptionally virulent virus in the model systems examined. In fact, no other human influenza viruses that have been tested show a similar pathogenicity for mice 3 to 4 days after infection. This information provides a partial explanation for what made this virus so lethal. In this regard, it should be noted that the U.S. Food and Drug Administration (FDA)–approved antiviral drugs, oseltamivir and amantadine, have been shown to be effective against viruses carrying the 1918 NA and the 1918 M gene, respectively (19). Furthermore, vaccines containing the 1918 HA and NA were protective in mice (20).

Note added in proof: This research was done by staff taking antiviral prophylaxis and using stringent biosafety precautions (15) to protect the researchers, the environment, and the public. The fundamental purpose of this work was to provide information critical to protect public health and to develop measures effective against future influenza pandemics.

References and Notes
15. Interim CDC-NIH Recommendation (www.cdc.gov/flu/h2n2s13.htm).
21. Materials and methods are available as supporting material on Science Online.

Reports

A Reversible, Unidirectional Molecular Rotary Motor Driven by Chemical Energy

Stephen P. Fletcher, Frédéric Dumur, Michael M. Pollard, Ben L. Feringa*

With the long-term goal of producing nanometer-scale machines, we describe here the unidirectional rotary motion of a synthetical molecular structure fueled by chemical conversions. The basis of the rotation is the movement of a phenyl rotor relative to a naphthyl stator about a single bond axle. The sense of rotation is governed by the choice of chemical reagents that power the motor through four chemically distinct stations. Within the stations, the rotor is held in place by structural features that limit the extent of the rotor’s Brownian motion relative to the stator.

One of the most challenging components required for the fabrication of molecular machines (1–5) is the rotary motor (6, 7): the element that converts energy into controlled rotational motion (8). Natural systems often use adenosine triphosphate (ATP) as an energy source. Rotation in these systems is powered by the energy released upon the hydrolytic conversion of ATP to the diphosphate ADP (6, 9, 10). The system introduced here analogously uses exothermic chemical reactions to power unidirectional rotary motion.

Previous synthetic molecular motor designs have often relied on external light or voltage as an energy source. Recent reports include repetitive, light-driven unidirectional motion about double bonds (11–13), as well as multistep reaction sequences that induce unidirectional (14) and reversible (15) mechanical motion in interlocked molecular rings, partly through photochemistry. A nanometer-scale rotational actuator based on multivalved carbon nanotubes has also been demonstrated (16). Theory further predicts the feasibility of inducing unidirectional rotation about a single bond in a chiral system by applying linearly polarized laser pulses within optimized electric fields (17, 18), and inducing mechanical motion in a double-walled carbon nanotube by applying axially varying electrical voltage (19).

Purely chemical strategies have been scarcer (7, 20, 21). Limited unidirectional (120°) rotation about a single bond in a helically shaped molecule (22, 23) has been achieved with a modified molecular ratchet (24–26). Here, we report a system that uses chemical energy to achieve unidirectional 360° rotation of one half of the molecule relative to the other half (Fig. 1). The rotation is driven by a combination of chemical reactions and random thermal (Brownian) motion. Understanding these processes may be relevant to natural molecular motors, which work on similar principles.

Our system consists of a rotor half and a stator half, connected by a single carbon-carbon bond which acts as the axis of rotation (Fig. 1). Chemical reactions control movement of the rotor through four structurally distinct stations, with the net effect of turning the rotor 360° relative to the stator. Bonding and steric constraints limit the extent of the rotor’s uncontrolled Brownian motion. In two of the stations (Fig. 1, stations A and C) the rotor’s position is restricted by the action of additional chemical bonds, although helix inversion can occur. In stations B and D (Fig. 1), the rotor and stator cannot pass each other due to nonbonding interactions. Movement between the stations is guided by four power strokes, or chemically induced rotational events. The complete cycle involves two bond-breaking steps (step 1 and step 3) and two bond-making steps (step 2 and step 4), each of which provides the driving force for approximately 90° unidirectional rotation to the next station.

Two general mechanisms (27) for the conversion of energy into mechanical motion have